

Biogeosciences Discussions is the access reviewed discussion forum of *Biogeosciences*

**Copper in
foraminiferal calcite**

L. J. de Nooijer et al.

Copper incorporation in foraminiferal calcite: results from culturing experiments

L. J. de Nooijer, G. J. Reichart, A. Dueñas-Bohórquez, M. Wolthers, S. R. Ernst, P. R. D. Mason, and G. J. van der Zwaan

Dept. of Earth Sciences, Utrecht University, Budapestlaan 4, 3584 CD Utrecht, The Netherlands

Received: 5 March 2007 – Accepted: 6 March 2007 – Published: 2 April 2007

Correspondence to: L. J. de Nooijer (nooijer@geo.uu.nl)

Title Page

Abstract

Introduction

Conclusions

References

Tables

Figures

◀

▶

◀

▶

Back

Close

Full Screen / Esc

Printer-friendly Version

Interactive Discussion

EGU

Abstract

A partition coefficient for copper (D_{Cu}) in foraminiferal calcite has been determined by culturing individuals of two benthic species under controlled laboratory conditions. The partition coefficient of a trace element (TE) is an empirically determined relation between the TE/Ca ratio in seawater and the TE/Ca ratio in foraminiferal calcite and has been established for many divalent cations. Despite its potential to act as a tracer of human-induced, heavy metal pollution, data is not yet available for copper. Since partition coefficients are usually a function of multiple factors (seawater temperature, pH, salinity, metabolic activity of the organism, etc.), we chose to analyze calcite from specimens cultured under controlled laboratory conditions. They were subjected to different concentrations of Cu^{2+} (0.1–20 $\mu\text{mol/l}$) and constant temperature (10 and 20°C), seawater salinity and pH. We monitored the growth of new calcite in specimens of the temperate, shallow-water foraminifer *Ammonia tepida* and in the tropical, symbiont-bearing *Heterostegina depressa*. Newly formed chambers were analyzed for Cu/Ca ratios by laser ablation-ICP-MS. The calculated partition coefficient (0.1–0.4) was constant to within experimental error over a large range of $(Cu/Ca)_{\text{seawater}}$ ratios and was remarkably similar for both species. Neither did the presence or absence of symbionts affect the D_{Cu} , nor did we find a significant effect of temperature or salinity on Cu-uptake.

1 Introduction

Trace elements incorporated in foraminiferal calcite tests are widely used in paleoceanography: Mg/Ca ratios are used to reconstruct sea surface (Nürnberg et al., 1996) and deep-sea temperatures (Rathburn and DeDecker, 1997), Cd and Ba are used to estimate past seawater nutrient levels and alkalinity, respectively (Boyle, 1988; Rosenthal et al., 1997; Lea and Boyle, 1991). These proxies rely on empirically derived partition coefficients (D_{TE}) and the dependence of these coefficients on environmental variables. Temperature, salinity and the pH of sea water all potentially affect the D_{TE} in

BGD

4, 961–991, 2007

Copper in foraminiferal calcite

L. J. de Nooijer et al.

Title Page

Abstract

Introduction

Conclusions

References

Tables

Figures

◀

▶

◀

▶

Back

Close

Full Screen / Esc

Printer-friendly Version

Interactive Discussion

EGU

foraminiferal calcite (e.g. Nürnberg et al., 1996; Segev and Erez, 2006).

Although field experiments are useful to determine first order proxy relationships, reliable proxy calibrations should include the contribution of so-called vital effects and separate the effects of other possible contributing factors. The best way to unravel the contribution of separate variables is through culturing experiments, in which one variable is varied and all the others are kept constant. In the case of some divalent cations (e.g. Mg^{2+} and Sr^{2+} : Nürnberg et al., 1996; Lea et al., 1999), culturing experiments also allow calibration of proxies out of the range of naturally occurring environmental conditions. This is important for trace elements that are associated with anthropogenic pollution with significantly raised concentrations above natural background levels.

Anthropogenic heavy metal pollution is often characterized by, amongst others, high Cu-concentrations (Borrego et al., 2004; Sáinz and Ruiz, 2006). Foraminifera have been used in several ways to investigate environmental pollution as high levels of Cu and other heavy metals potentially deform foraminiferal chamber alignment and influence foraminiferal community structure (Ellison et al., 1986; Samir and El-Din, 2001; Hallock et al., 2003; Armynot du Châtelet et al., 2004; Ruiz et al., 2004; Ferraro et al., 2006). However, a number of studies state that test deformations under high heavy metal concentrations occur less often than under medium pollution loads (Alve and Olsgard, 1999; Geslin et al., 2002; Le Cadre and Debenay, 2006). This suggests that reconstructions based on test deformations alone are not accurate.

Cu-concentrations can also be high in the proximity of hydrothermal vents (Iizasa, 1993; Douville et al., 2002; Kadar et al., 2005). Although the low pH close to acidic vent fluids dissolves foraminiferal calcite, records of benthic foraminiferal assemblages described further away from these vents may be used to reconstruct past hydrothermal activity and the impact of high concentrations of heavy metals in its vicinity (Molina-Cruz and Ayala-López, 1988; Jonasson et al., 1995; Panieri et al., 2005).

In previous studies, Cu has been one of the most difficult elements to analyze in foraminiferal calcite (Boyle, 1981). However, recent advances in analytical methodology for trace element determination in foraminiferal calcite (Reichert et al., 2003) now

BGD

4, 961–991, 2007

**Copper in
foraminiferal calcite**

L. J. de Nooijer et al.

Title Page

Abstract

Introduction

Conclusions

References

Tables

Figures

◀

▶

◀

▶

Back

Close

Full Screen / Esc

Printer-friendly Version

Interactive Discussion

EGU

enable the calibration of foraminiferal Cu to seawater chemistry for the first time, using cultured benthic foraminifera. Two different intertidal to neritic species (one temperate and one tropical) were cultured to establish possible interspecific differences in the partition coefficient of Cu in foraminiferal calcite.

2 Methods

2.1 Collecting and culturing foraminifera

Two similar culturing experiments were conducted in series. For the first experiment, sediment was collected at an intertidal flat in the Dutch Wadden Sea and was kept in the laboratory in the dark at 15°C. Large (>150 µm), living individuals of *Ammonia* cf. molecular type T6 (Hayward et al., 2004: further referred to as *A. tepida*) were transferred to our in-house designed flow-through culture vessels (Fig. 1). Vessels consist of a 24-well culture tray, sandwiched between two Plexiglas plates and cells were connected by silicon tubes, attached with luers in the upper Plexiglas lid (Fig. 1). A small filter was placed between each cell and tube to prevent specimens from moving between cells. Trays were connected individually to a 2-liter reservoir with chemically altered sea water (see below) and a peristaltic pump was used to circulate sea water through the cells with a speed of 9 ml/h: in this way, six groups of 12 cells were formed, each connected to its own sea water reservoir. Four foraminiferal specimens were placed in each cell. Sea water was enriched with Cu from a stock solution at concentrations of 0, 0.10, 0.20, 0.50, 10 and 20 µmol/l. Calcein (C0875, Sigma-Aldrich, St Louis, USA) was added to the Cu-enriched sea water at a concentration of 5 mg/l. Calcein is incorporated into biogenic calcite, while existing calcite (i.e. earlier formed chambers) is not affected. Since (incorporated) calcein is fluorescent, foraminiferal chambers that have been built during the time when individuals were incubated can be recognized (Bernhard et al., 2004).

Cells with specimens of *Ammonia tepida*, contained a thin layer (<0.5 mm) of artificial

BGD

4, 961–991, 2007

Copper in foraminiferal calcite

L. J. de Nooijer et al.

Title Page

Abstract

Introduction

Conclusions

References

Tables

Figures

◀

▶

◀

▶

Back

Close

Full Screen / Esc

Printer-friendly Version

Interactive Discussion

EGU

sediment (Silica, 52–63 μm). Natural seawater from the eastern Mediterranean Sea was adjusted with MilliQ water to a salinity of 17 to mimic average Wadden Sea salinity. Salinity levels were regularly checked during the experiment with a WTW LF330 conductivity meter. All 6*12 cells were kept at a constant temperature of 10°C for two months: before and after experiments, reservoirs were sub-sampled and sea water was analyzed by ICP-MS for Cu, Mg and Ca. At the start of the incubation period, the individuals were fed ~0.5 mg of autoclaved (20 min at 121°C) *Dunaliella* sp. During experiments, the set-up was subjected to the daily sunlight cycle (app. 14 h light/10 h dark).

For experiment 2, trays were replaced and lids rigorously cleaned with HCl, rinsed with MiliQ and re-used to incubate individuals of *Heterostegina depressa* in seawater with similar Cu-enrichments used in the first experiments. *H. depressa* is an epibenthic, tropical and symbiont-bearing foraminifer, that was kept in our laboratory under high light intensities (15W tropical reef lamp; Arcadia, FO15) and after transferring them into the culturing set-up, similar light conditions were maintained in a daily rhythm (14 h light/10 h dark). No sediment was added to the cells, seawater salinity was kept at 35, with a constant temperature of 20°C. Because of their large size, only two specimens were placed in each cell.

3 Temperature and salinity of culture media

For culturing *Ammonia tepida*, we diluted 35 PSU seawater with MiliQ water to mimic intertidal ambient conditions with seawater of 17 PSU. The dilution decreased both $[\text{Ca}^{2+}]$ and $[\text{CO}_3^{2-}]$ and the alkalinity by approximately 50%, resulting in a pronounced reduction of the carbonate saturation state (Ω). The temperature maintained during these experiments was kept at 10°C, compared to 20°C for the *Heterostegina depressa* experiments, allowing gas exchange with the atmosphere (ambient pCO_2) in both cases. The combined effect of these changes is a reduction in saturation state from about 5.5 for the *H. depressa* experiment to about 1.0 for the *A. tepida* experiment (calculations

BGD

4, 961–991, 2007

Copper in foraminiferal calcite

L. J. de Nooijer et al.

Title Page

Abstract

Introduction

Conclusions

References

Tables

Figures

◀

▶

◀

▶

Back

Close

Full Screen / Esc

Printer-friendly Version

Interactive Discussion

EGU

were performed in CO₂sys; Lewis and Wallace, 1998). The lower seawater saturation state for the *A. tepida* cultures was most likely responsible for the fact that newly formed chambers were thinner than the pre-experiment chambers (see Results).

4 Laser ablation ICP-MS

5 Newly formed chambers were ablated using an Excimer laser (Lambda Physik) with GeoLas 200Q optics inside a helium atmosphere flushed ablation chamber. Pulse repetition rate was set at 6 Hz, with an energy density at the sample surface of 10 J/cm². Ablation craters were 60 μm in diameter and ablated material was analyzed with respect to time (and hence depth) using a quadrupole ICP-MS instrument (Micromass Platform ICP).

10 Simultaneous monitoring of Al allowed us to discard the parts of the ablation profiles contaminated by clay minerals from further calculations of elemental concentrations. Since the analytical error increases with shorter ablation time we cleaned all specimens by an incubation of 24 h in 5% NaOCl (Gaffey and Brönniman, 1993) before analysis, maximizing the amount of data that could be used for calculating (Cu/Ca)_{calcite} ratios.

5 Calibration strategy

The low calcite saturation state used in the experiment with *Ammonia tepida* resulted in formation of new chambers with thin walls. A similar correlation between test wall thickness and carbonate saturation state has been observed earlier for tests of cultured planktonic foraminifera (Bijma et al., 2002). Unfortunately, these thin chambers break easily during ablation when high laser energies are used. Therefore, we ablated *Ammonia tepida* with a laser energy of 1 J/cm², ten times less than the 10 J/cm² used to ablate newly formed chambers of *Heterostegina depressa*.

25 Analyses were calibrated against NIST glasses 610 and 612, using concentration data of Pearce et al. (1997). Calibrating calcites against glasses is possible because

Title Page

Abstract

Introduction

Conclusions

References

Tables

Figures

◀

▶

◀

▶

Back

Close

Full Screen / Esc

Printer-friendly Version

Interactive Discussion

of the relatively matrix independent ablation by the Excimer laser (Mason and Kraan, 2002). However, a fluence of $<2\text{ J/cm}^2$ was close to the ablation threshold for glass and calibration was performed instead against matrix matched in-house standards (i.e. pressed calcite powder tablets). Calcium was used as an internal standard because (1) the concentration is constant at 40 wt % in calcite and (2) it allows direct comparisons with trace metal to Ca ratios from wet-chemical studies. A collision and reaction cell was used to give improved results by reducing spectral interferences on the minor isotopes of Ca (^{42}Ca , ^{43}Ca and ^{44}Ca : Mason and Kraan, 2002). Good agreement was observed when using both ^{63}Cu and ^{65}Cu isotopes to calculate Cu-concentrations. Relative analytical precision for copper analyses was 15% on average, based on variability during the ablation calculated by GLITTER (New Wave Research, Fremont, CA, USA). This error includes both analytical uncertainties and internal, natural variability in test chemistry.

6 Seawater Cu-concentration

The concentration of Cu did not vary considerably in most of our experiments during the experimental period across the range of concentrations used (Table 1).

In the first experiment, all measured Cu-concentrations were lower than the target concentration and most total Cu-concentrations increased during the experiment, resulting in increased seawater Cu/Ca ratios (on average 17%). In experiment 2, most Cu-concentrations and all Cu/Ca ratios were higher at the start than at the end of the experiment. Identical procedures and techniques were used before and after subsampling the culture media, making it unlikely that sampling artifacts affected our measurements. Therefore, we used average solution Cu/Ca ratios to calculate the partition coefficient of Cu in foraminiferal calcite and incorporated differences between start and end concentrations for uncertainty calculations. Error bars plotted in the different graphs are based on these calculations and largely stem from these changes, which are an order of magnitude larger than the analytical uncertainties.

BGD

4, 961–991, 2007

Copper in foraminiferal calcite

L. J. de Nooijer et al.

Title Page

Abstract

Introduction

Conclusions

References

Tables

Figures

◀

▶

◀

▶

Back

Close

Full Screen / Esc

Printer-friendly Version

Interactive Discussion

EGU

7 Cu speciation in seawater

In the absence of organic matter, Cu in sea water forms mainly $\text{Cu}(\text{OH})_2$ and CuCO_3 , while small amounts of Cu^{2+} and CuOH^- are also present (Zirino and Yamamoto, 1972). In natural sea water, however, usually more than 99.9% of the Cu is bound to organic compounds (Eriksen et al., 2001), mainly in the colloidal state (Mackey and Zirino, 1994). Foraminifera take up organic particles and sea water by endocytosis, likely ingesting both free Cu and Cu-ligand complexes. The internal routes that organic compounds follow are virtually uninvestigated in foraminifera and therefore, we do not know which Cu species are present at the site of calcification.

8 Modelling Cu speciation

The calcein added in our experiments is a ligand that can bind TE's and could thus cause concentrations of free Cu to drop. Traditionally, total calcium and TE concentrations in solution are used to calculate partition coefficients for TE's in calcite. Ideally, activities or effective concentrations, of relevant metals are used to allow application of partition coefficients in solutions of different compositions (Morse and Bender, 1990).

To correct for Cu binding to calcein, we calculated speciation of all abundant cations (Cu, Ca and Mg) in our solution. Speciation calculations were performed in PHREEQC 2.8.03 (Parkhurst and Appelo, 1999) with the IlnI database and thermodynamic data for calcein listed in Table 2. For calcein complexation with calcium and magnesium, Reactions (11) and (14) are reported in the literature. These reactions will lead to competition between copper, calcium and magnesium in binding to an amine group on H_2L^{4-} , thus decreasing calcein-bound Cu. It is, therefore, likely that the behavior of Ca and Mg towards calcein is similar to Cu and similar competition between the three metals occurs in binding according to Reactions (7) and (9) via carboxyl groups (Lu and Allen, 2002). Composition of the solution in the model was either the Cu-enriched sea water with a salinity of 35 or 17 for the experiment with *Ammonia tepida*, and were

BGD

4, 961–991, 2007

Copper in foraminiferal calcite

L. J. de Nooijer et al.

Title Page

Abstract

Introduction

Conclusions

References

Tables

Figures

◀

▶

◀

▶

Back

Close

Full Screen / Esc

Printer-friendly Version

Interactive Discussion

EGU

both open to atmospheric CO₂.

Cu and Ca-concentrations used to calculate the partition coefficients were those corrected for Cu and Ca complexed with calcein (Table 3).

9 Results

5 9.1 New calcite and survival rates

Specimens that grew new calcite, were recognized by fluorescent, outer chambers (Fig. 2).

10 None of the individuals of *Heterostegina depressa* incubated at the target Cu-concentration of 20 μmol/l, survived the experimental period. At 10 μmol/l of added Cu, however, several survived of which 1 individual grew new calcite. At lower concentrations, generally more chambers were formed (Table 4). None of the added chambers (n=88) showed abnormal alignments or deformations.

15 For *Ammonia tepida*, the number of successful laser-ablation analyses was significantly lower (3) than the number of added chambers (34). The limited size (<100 μm) of newly added chambers did not allow multiple analyses of a single chamber. After an unsuccessful attempt to analyze a targeted chamber it was not possible to repeat this measurement as the largest part of the carbonate was consumed (Fig. 3).

9.2 Partition coefficient of Cu – *Ammonia tepida*

20 Two ablation profiles were obtained from two specimens of *Ammonia tepida* that grew new chambers at a low (0.20 μmol/l) Cu-concentration (Fig. 4). Measurements indicate that the partition coefficient lies between 0.1 and 0.4. In the right panel of Fig. 4, the same two measurements are depicted at the left end of the graph. Ratios for calcite formed at higher (Cu/Ca)_{seawater}, indicated a partition coefficient between 0.1 and 0.4.

BGD

4, 961–991, 2007

Copper in foraminiferal calcite

L. J. de Nooijer et al.

Title Page

Abstract

Introduction

Conclusions

References

Tables

Figures

◀

▶

◀

▶

Back

Close

Full Screen / Esc

Printer-friendly Version

Interactive Discussion

EGU

9.3 Partition coefficient of Cu – *Heterostegina depressa*

Although individuals of *Heterostegina* did not survive the highest Cu-levels, we obtained two ratios from specimens that added new chambers at a target concentration of $10 \mu\text{mol/l}$. From incubations with lower Cu-concentrations, more specimens were available that grew new chambers that could be analyzed for Cu-concentration (Fig. 5).

10 Discussion

Within the experimental and analytical error both species show an identical $(\text{Cu}/\text{Ca})_{\text{seawater}}$ to $(\text{Cu}/\text{Ca})_{\text{calcite}}$ relation, resulting in a partition coefficient (D_{Cu}) of 0.25 ± 0.15 . No significant difference was observed in copper incorporation between *Ammonia tepida* and *Heterostegina depressa*, despite large differences in ecology and habitat. Some inter-specimen variation in $\text{Cu}/\text{Ca}_{\text{calcite}}$ was observed in *H. depressa* grown at low Cu-concentrations, in which rather large uncertainties in culture water Cu-concentration (Fig. 5) resulted from changes in $[\text{Cu}^{2+}]$ over time. Moreover, the alternative calibration method used for the thin-walled chambers of *A. tepida* (i.e. using a pressed calcite pellet and a lower ablation energy) increased the analytical uncertainty in the laser ablation-ICP-MS analyses (Fig. 5). Despite these errors, the calculated D_{Cu} was not significantly dependent on either temperature or salinity.

10.1 Experimental uncertainties

The seawater Cu-concentrations in most culture media increased during the incubation period (Table 1), however, there is no systematical difference in $\text{Cu}/\text{Ca}_{\text{seawater}}$ ratios before and after the two experiments. Therefore, an analytical error during ICP-MS or during subsampling of the culture vessels is unlikely to play an important role in this offset. Sorption of Cu onto organic matter in the culture media or onto the calcite of the foraminifera would have lowered the Cu-concentrations in the media, which happened in some cases, and therefore could have contributed to the differences in Cu/Ca

BGD

4, 961–991, 2007

Copper in foraminiferal calcite

L. J. de Nooijer et al.

Title Page

Abstract

Introduction

Conclusions

References

Tables

Figures

◀

▶

◀

▶

Back

Close

Full Screen / Esc

Printer-friendly Version

Interactive Discussion

EGU

ratios. Alternatively, release of Cu by the culture vessels or trays could have increased seawater Cu-concentrations, although this is unlikely since the equipment used was new or cleaned prior to the experiment. Moreover, the control media (no Cu added) did not contain considerable amounts of Cu, indicating that contamination by the materials used played a minor role. The foraminifera themselves are unlikely to have contained high concentrations of Cu at the start of the experiments since concentrations of this metal are low in natural seawater. Cu-pollution of seawater during subsampling for seawater Cu-analyses may have caused an increase in Cu-concentrations, although care was taken to avoid such contamination. The difference in seawater Cu-concentrations before and after the incubation period are likely to be the result of a combination of the processes mentioned above.

The number of new chambers formed by the cultured foraminifera was generally low, especially at high Cu-concentrations (Table 4). The limited production of new calcite may indicate that the conditions during the experiments were not optimal for the foraminifera. Consequently, the reaction of the foraminifera to these conditions may have influenced their calcification and thus the derived D_{Cu} . Improving culturing set-ups and optimizing laboratory conditions is important and comparison of our results with those from optimized, future experiments is necessary to establish a more precise D_{Cu} for foraminiferal calcite.

10.2 Cu in the calcite lattice

Crystalline $CuCO_3$ does not exist, because the most common coordination of Cu-carbonate complexes are distorted tetragonal pyramids or distorted octahedrons (Wells, 1984). These shapes do not allow precipitation of pure $CuCO_3$ crystals and rather $Cu_2CO_3(OH)_2$ (malachite) will form. However, sorption studies have shown that at the calcite-water interface these so-called Jahn-Teller distortions can be overcome and a solid solution $Cu_xCa_{(1-x)}CO_3$ forms (Schlosseler et al., 1999). It has been proposed that copper in calcite is present in clusters, based on studying the transformation of vaterite to calcite (Nassrallah-Aboukaïs et al., 1996, 1998). Recent XAFS work, how-

BGD

4, 961–991, 2007

Copper in foraminiferal calcite

L. J. de Nooijer et al.

Title Page

Abstract

Introduction

Conclusions

References

Tables

Figures

◀

▶

◀

▶

Back

Close

Full Screen / Esc

Printer-friendly Version

Interactive Discussion

EGU

ever, has shown that this mechanism is not applicable to calcite surfaces (Elzinga and Reeder, 2002). This rather unusual complexation behavior would suggest that sorption and subsequent incorporation into the crystal lattice for copper is limited to part of the crystal surface only. This, in turn, would result in a lower partition coefficient for copper than expected based on its ionic radius only.

Contrary to these results, it has been shown that during inorganic coprecipitation experiments Cu is incorporated in calcite with a distribution coefficient (K_{Cu}) of 23 and constant under a range of Cu-concentrations (Kitano et al., 1980). In the initial stage of calcification, the K_{Cu} can be even higher (40) probably due to the strong affinity of $Cu(OH)_2$ for calcite surfaces (Franklin and Morse, 1982; Pickering, 1983; Papadopoulos and Rowell, 1989). This indicates that the Jahn-Teller distortions are easily overcome during calcification.

Generally, divalent cations with an ionic radius close to Ca ($=1.0 \text{ \AA}$) have a partition coefficient in calcite close to 1. Cd has an ionic radius of 0.95 (Shannon, 1976) and is incorporated in both planktonic and benthic species with a D between 1 and 4 (Boyle, 1981, 1988; Havach et al., 2001), independent of temperature (Marchitto, 2004). Sr (ionic radius = 1.31 \AA) is incorporated in foraminiferal calcite with a D of 0.11–0.19, measured in several planktonic genera (Bender et al., 1975) and 0.05–0.25 in *Cibicides* (Elderfield et al., 1996). Coretop studies on *Cibicides* and *Uvigerina* show that Ba (1.47 \AA) is incorporated with a partition coefficient of 0.3–0.4 at 3°C (Lea and Boyle, 1989). In the planktonic genera *Globorotalia* and *Globoquadrina*, Ba is incorporated with a D of 0.19 (Lea and Boyle, 1991). Cu has an ionic radius close to Mg (0.73 and 0.72 \AA , respectively), but the partition coefficient of Mg is much lower ($0.1-1 \times 10^{-3}$; Bender et al., 1975; Delaney et al., 1985) than the measured 0.1–0.4 for Cu (Fig. 7). The large difference between the foraminifer-mediated Cu-incorporation and the inorganic incorporation of Cu in calcite indicates that much energy is spent on removal of Cu at the site of calcification.

BGD

4, 961–991, 2007

Copper in foraminiferal calcite

L. J. de Nooijer et al.

Title Page

Abstract

Introduction

Conclusions

References

Tables

Figures

◀

▶

◀

▶

Back

Close

Full Screen / Esc

Printer-friendly Version

Interactive Discussion

EGU

10.3 Biological control on D_{Cu}

Since magnesium inhibits calcite growth (Berner, 1975; Mucci and Morse, 1983) and high levels of Mg are likely to be present in foraminiferal calcifying reservoirs, it is necessary for foraminifera to remove Mg before calcification. It has been suggested that foraminifera actively pump Mg from their calcifying reservoir in order to stimulate $CaCO_3$ precipitation. Usually, Cu is present only in very low concentrations in seawater and therefore no need exists to actively remove Cu from calcifying reservoirs, despite its ability to modify the crystalline structure of calcite. Although under high Cu-concentrations it may be beneficial to remove Cu from calcifying reservoirs, apparently the foraminifera does not do so, as the D_{Cu} is similar for high and low Cu-concentrations. Alternatively, the concentrations used in our experiments may still be too low to seriously impede $CaCO_3$ precipitation.

Another reason for active removal of trace elements from calcifying reservoirs is that these elements are necessary for cellular processes. Since Cu is known to play only minor roles in eukaryotic metabolic processes (Bruland et al., 1991; Sunda and Huntsman, 1995; Chang and Reinfelder, 2000), it is unlikely that the D_{Cu} is affected by cellular needs.

Organic compounds may increase Mg contents in foraminiferal calcite (Bentov and Erez, 2006). High concentrations of Mg in the primary organic membrane (Hemleben et al., 1986) may explain the observed intra-test variability of Mg/Ca (e.g. Toyofuku and Kitazato, 2005). Cu also has a strong affinity for organic compounds (see below), so that the D_{Cu} may be partly determined by the presence of organic compounds in the calcite.

Bresler and Yanko (1995) showed that some benthic, epiphytic foraminifera have tryptofan-containing proteins that can bind Cu^{2+} and prevent intracellular Cu-concentrations from becoming harmful. When a significant part of the Cu^{2+} would have been immobilized by these Cu-binding proteins this would also have lowered the Cu activity in the solution and thus D_{Cu} . Since we have not observed such a de-

BGD

4, 961–991, 2007

Copper in foraminiferal calcite

L. J. de Nooijer et al.

Title Page

Abstract

Introduction

Conclusions

References

Tables

Figures

◀

▶

◀

▶

Back

Close

Full Screen / Esc

Printer-friendly Version

Interactive Discussion

EGU

crease, it is unlikely that such molecules play a major role in decreasing intracellular Cu-concentrations.

Seawater pH is a potentially important modulator of trace metal uptake (Lea et al., 1999; Zeebe and Sanyal, 2002). To investigate the potential effect we compared species with and without symbionts. In the symbiont-bearing *H. depressa* the photosynthetic activity of the symbionts changes the local carbonate chemistry because CO₂ is taken up and pH lowered during light conditions. However, the lack of any systematic offset in Cu/Ca between the *H. depressa* and *A. tepida* suggests no significant effect of pH on Cu incorporation.

10.4 Test deformation and mortality

A number of studies over the last 20 years have attempted to correlate the number of deformed tests to environmental pollution (Alve, 1991; Elberling et al., 2003; Armynot du Châtelet et al., 2004). The empirical correlation between number of deformed tests and for instance heavy metal or hydrocarbon concentration levels was interpreted to signify a causal relationship. Although in several of our experiments Cu-concentrations were well above levels found at even the most polluted sites not a single deformed chamber alignment was observed. Although the limited number of observations does not allow a statistical evaluation, our results strongly suggest that high levels of Cu do not cause test deformities. This is also shown by Alve and Olsgard (1999), who found no test deformities in foraminifera living in seawater with high Cu-concentrations. Therefore, we think that relative abundances of deformed tests in fossil samples are not suitable to reconstruct past copper concentrations. Most likely other environmental factors, co-varying with environmental trace metal levels must have been responsible for the observed increase in test deformities. The complete absence of deformations in our experiments is in high contrast to the low but still detectable levels of natural occurring test deformities under environmental pristine conditions. This suggests that the protected environment of the culture trays may actually have shielded our foraminifera by providing them with a constant temperature, salinity and seawater chemistry in gen-

BGD

4, 961–991, 2007

Copper in foraminiferal calcite

L. J. de Nooijer et al.

Title Page

Abstract

Introduction

Conclusions

References

Tables

Figures

◀

▶

◀

▶

Back

Close

Full Screen / Esc

Printer-friendly Version

Interactive Discussion

EGU

eral.

In seawater with the highest concentration of Cu ($20 \mu\text{mol/l}$), none of the *Heterostegina*'s survived and only one specimen grew new chambers when cultured at $10 \mu\text{mol/l}$. Since the growth or survival of *Ammonia tepida* did not appear to be hampered by high concentrations of Cu, we hypothesize that either the symbionts of the tropical foraminifera are vulnerable to high Cu-concentrations (Brandt et al., 1986), or that individuals of *A. tepida* are adapted to cope with (occasional) high levels of heavy metals.

10.5 Application of Cu/Ca ratios in foraminiferal calcite

In order to quantify pollution levels, heavy metal concentrations are often analyzed using strong acid extractions and subsequent ICP-MS analyses of bulk sediment. Since heavily polluted sites are frequently characterized by high concentrations of (labile) organic matter, polluted sediments are often anoxic with high levels of sulphate reduction and associated production of free sulfide. These high sulfide-levels result in immobilization of heavy metals such as Zn, Cu, Cd and Pb, which are precipitated as the highly insoluble minerals PbS, CuS and ZnS, or as co-precipitates in pyrite. Because these metals are no longer bioavailable they do not reflect toxicity of the overlying water to, for instance, benthic biota. Actual analyses of the overlying water itself or organisms living in these waters would give a much more applicable concentration to assess pollution. This becomes even more important when at a later stage organic loads decrease and/or the oxygen level increases, (e.g. after improved wastewater treatment). Under these conditions lower sedimentary trace metal levels could result in higher actual toxicity as these metals are remobilized by progressive reoxidation of the sediment and escape to the overlying water. Monitoring foraminiferal test Cu/Ca ratios could be used to establish the bioavailable fraction of Cu and potentially also could record relatively short episodes with elevated bottom water Cu-levels.

Sludge dump sites and associated elevated concentration levels of heavy metals are mostly limited to coastal and estuarine environments. These settings experience considerably varying seasonal and even daily temperatures and salinity levels. A signif-

BGD

4, 961–991, 2007

Copper in foraminiferal calcite

L. J. de Nooijer et al.

Title Page

Abstract

Introduction

Conclusions

References

Tables

Figures

◀

▶

◀

▶

Back

Close

Full Screen / Esc

Printer-friendly Version

Interactive Discussion

EGU

5 icant impact of either temperature or salinity on partition coefficients would, therefore, render foraminiferal trace metal records useless for any reliable reconstruction and/or monitoring of such dump sites. Since the obtained D_{Cu} is not markedly dependent on either temperature or salinity, foraminiferal Cu/Ca ratios may be a powerful proxy for the quantitative reconstruction of past heavy metal pollution, even in highly variable environments.

11 Conclusions

10 Copper is incorporated into foraminiferal calcite with a partition coefficient of 0.25 ± 0.15 with respect to sea water Cu/Ca values. No effects on the D_{Cu} of species-specific control or of temperature and salinity could be observed with the experimental setup used here. Additional experiments are needed to better constrain D_{Cu} and unravel the effects of other likely important environmental factors such as temperature, salinity and seawater carbonate chemistry.

15 *Acknowledgements.* The authors thank M. Janse from Burger's Zoo and W. Renema from Naturalis for providing specimens of *Heterostegina depressa*. P. Kleingeld is acknowledged for his help with developing the culturing set-up. The laser ablation measurements were greatly aided by G. Nobbe. E. van Vilsteren and B. van Os provided help with the ICP-MS-analyses and discussion on sea water chemistry.

References

- 20 Alve, E.: Benthic foraminifera in sediment across cores reflecting heavy metal pollution inm Sør fjord, western Norway, *J. Foraminifer. Res.*, 21(1), 1–19, 1991.
- Alve, E. and Olsgard, F.: Benthic foraminiferal colonization in experiments with copper-contaminated sediment, *J. Foraminifer. Res.*, 29(3), 186–195, 1999.
- 25 Armynot du Châtelet, E., Debenay, J.-P., and Soulard, R.: Foraminiferal proxies for pollution monitoring in moderately polluted harbors, *Environ. Pollut.*, 127, 27–40, 2004.

BGD

4, 961–991, 2007

Copper in foraminiferal calcite

L. J. de Nooijer et al.

Title Page

Abstract

Introduction

Conclusions

References

Tables

Figures

◀

▶

◀

▶

Back

Close

Full Screen / Esc

Printer-friendly Version

Interactive Discussion

EGU

- Bender, M. L., Lorens, R. B., and Williams, D. F.: Sodium, magnesium, and strontium in the tests of planktonic foraminifera, *Micropaleontol.*, 21, 448–459, 1975.
- Bentov, S. and Erez, J.: Impact of biomineralization processes on the Mg content of foraminiferal shells: A biological perspective, *Geochem. Geophys. Geosyst.*, 7(1), Q01P08, 2006.
- Bernhard, J. M., Blanks, J. K., Hintz, C. J., and Chandler, G. T.: Use of fluorescent calcite marker calcein to label foraminiferal tests, *J. Foraminif. Res.*, 34(2), 96–101, 2004.
- Berner, R. A.: The role of magnesium in the crystal growth of calcite and aragonite from seawater, *Geochim. Cosmochim. Acta*, 39(4), 489–504, 1975.
- Bijma, J., Honisch, B., and Zeebe, R. E.: Impact of the ocean carbonate chemistry on living foraminiferal shell weight: Comment on “carbonate ion concentration in glacial-age deep waters of the Caribbean Sea” by W. S. Broecker and E. Clark., *Geochem. Geophys. Geosyst.*, 3(11), 1064, 2002.
- Borrego, J., López-González, N., and Carro, B.: Geochemical signature as paleoenvironmental markers in Holocene sediments of the Tinto River estuary (Southwestern Spain), *Est., Coast. Shelf Sci.*, 61, 631–641, 2004.
- Boyle, E. A.: Cadmium, zinc, copper, and barium in foraminifera tests, *Earth Planet. Sci. Lett.*, 53, 11–35, 1981.
- Boyle, E. A.: Cadmium: chemical tracer of deepwater paleoceanography, *Paleoceanography*, 3, 471–489, 1988.
- Brandt, L. E., Sunda, W. G., and Guillard, R. R. L.: Reduction of marine phytoplankton reproduction rates by copper and cadmium, *J. Exp. Mar. Biol. Ecol.*, 96, 225–250, 1986.
- Bresler, V. and Yanko, V.: Chemical ecology : A new approach to the study of living benthic epiphytic foraminifera, *J. Foraminif. Res.*, 25(3), 267–279, 1995.
- Bruland, K. W., Donat, J. R., and Hutchins, D. A.: Interactive influences of bioactive trace metals on biological production in oceanic waters, *Limnol. Oceanogr.*, 36(1), 1555–1577, 1991.
- Chang, S. I. and Reinfelder, J. R.: Bioaccumulation, subcellular distribution, and trophic transfer of copper in a coastal marine diatom, *Environ. Sci. Technol.*, 34, 4931–4935, 2000.
- Delaney, M. L., Bé, A. W. H., and Boyle, E. A.: Li, Sr, Mg, and Na in foraminiferal calcite shells from laboratory culture, sediment traps, and sediment cores, *Geochim. Cosmochim. Acta*, 49(6), 1327–1341, 1985.
- Douville, E., Charlou, J. L., Oelkers, E. H., Bienvvenu, P., Jove Colon, C. F., Donval, J. P., Fouquet, Y., Prieur, D., and Appriou, P.: The Rainbow vent fluids (36°14' N, MAR): the influence

BGD

4, 961–991, 2007

**Copper in
foraminiferal calcite**

L. J. de Nooijer et al.

Title Page

Abstract

Introduction

Conclusions

References

Tables

Figures

◀

▶

◀

▶

Back

Close

Full Screen / Esc

Printer-friendly Version

Interactive Discussion

EGU

- of ultramafic rocks and phase separation on trace metal content in Mid-Atlantic Ridge hydrothermal fluids, *Chem. Geol.*, 184, 37–48, 2002.
- Elberling, B., Knudsen, K. L., Kristensen, P. H., and Asmund, G.: Applying foraminiferal stratigraphy as a biomarker for heavy metal contamination and mining impact in a fiord in west Greenland, *Mar. Environ. Res.*, 55, 235–256, 2003.
- Elderfield, H., Bertram, C. J., and Erez, J.: A biomineralization model for the incorporation of trace elements into foraminiferal calcite, *Earth Planet. Sci. Lett.*, 142, 409–423, 1996.
- Ellison, R. L., Broome, R., and Oglivie, R.: Foraminiferal response to trace metal contamination in the Patapsco River and Baltimore harbour, Maryland, *Mar. Pollut. Bull.*, 17(9), 419–423, 1986.
- Elzinga, E. J. and Reeder, R. J.: X-ray absorption spectroscopy study of Cu^{2+} and Zn^{2+} adsorption complexes at the calcite surface: Implications for site-specific metal incorporation preferences during calcite crystal growth, *Geochim. Cosmochim. Acta*, 66(22), 3943–3954, 2002.
- Eriksen, R. S., Mackey, D. J., Van Dam, R., and Nowak, B.: Copper speciation and toxicity in Macquarie Harbour, Tasmania: an investigation using a copper ion selective electrode, *Mare Chem.*, 74, 99–113, 2001.
- Ferraro, L., Sprovieri, M., Alberico, I., Lirer, F., Prevedello, L., and Marsella, E.: Benthic foraminifera and heavy metals distribution: A case study from the Naples harbour (Tyrrhenean Sea, Southern Italy), *Environ. Pollut.*, 142, 274–287, 2006.
- Franklin, M. L. and Morse, J. W.: The interaction of copper with the surface of calcite, *Ocean Sci. Eng.*, 7, 147–174, 1982.
- Gaffey, S. J. and Brönnimann, C. E.: Effects of bleaching on organic and mineral phases in biogenic carbonates, *Journal of Sediment. Petrol.*, 63, 752–754, 1993.
- Geslin, E., Debenay, J.-P., Duleba, W., and Bonetti, C.: Morphological abnormalities of foraminiferal tests in Brazilian environments: comparison between polluted and non-polluted areas, *Mar. Micropaleontol.*, 45, 151–168, 2002.
- Hallock, P., Lidz, B. H., Cockey-Burkhard, E. M., and Donnelly, K. B.: Foraminifera as bioindicators in coral reef assessment and monitoring: the FORAM index, *Environ. Monit. and Assess.*, 83(1–3), 221–238, 2003.
- Havach, S. M., Thomas Chandler, G., Wilson-Finelli, A., and Shaw, T. J.: Experimental determination of trace element partition coefficients in cultured benthic foraminifera, *Geochim. Cosmochim. Acta*, 65(8), 1277–1283, 2001.

BGD

4, 961–991, 2007

**Copper in
foraminiferal calcite**L. J. de Nooijer et al.

[Title Page](#)[Abstract](#)[Introduction](#)[Conclusions](#)[References](#)[Tables](#)[Figures](#)[◀](#)[▶](#)[◀](#)[▶](#)[Back](#)[Close](#)[Full Screen / Esc](#)[Printer-friendly Version](#)[Interactive Discussion](#)**EGU**

- Hayward, B. W., Holzmann, M., Grenfell, H. R., Pawlowski, J., and Triggs, C. M.: Morphological distinction of molecular types in *Ammonia* – towards a taxonomic revision of the world's most commonly misidentified foraminifera, *Mar. Micropaleontol.*, 50, 237–271, 2004.
- Hemleben, C. H., Anderson, O. R., Berthold, W., and Spindler, M.: Calcification and chamber formation in Foraminifera- a brief overview, in: Biomineralization in lower plants and animals, Systematics Association Special Volume 30, edited by: Leadbeater, B. S. C. and Riding, R., Clarendon Press, Oxford (for the Systematics Association), p 237–249, 1986.
- lizasa, K.: Assessment of the hydrothermal contribution to seafloor sediments in the Myojinsho submarine caldera, Shichito-Iwojima ridge, Izu-Ogasawara arc, Japan, *Mar. Geol.*, 114(1–2), 119–132, 1993.
- Jonasson, K. E., Schröder-Adams, C. J., and Patterson, R. T.: Benthic foraminiferal distribution at Middle Valley, Juan de Fuca Ridge, a northeast Pacific hydrothermal venting site, *Mar. Micropaleontol.*, 25, 151–167, 1995.
- Kádár, E., Costa, V., Martins, I., Serrao Santos, R., and Powell, J. J.: Enrichment in trace metals (Al, Mn, Co, Cu, Mo, Cd, Fe, Zn, Pb and Hg) of macro-invertebrate habitats at hydrothermal vents along the Mid-Atlantic Ridge, *Hydrobiologia*, 548, 191–205, 2005.
- Kitano, Y., Okumura, M., and Idogaki, M.: Abnormal behaviors of copper (II) and zinc ions in parent solution at the early stage of calcite formation, *Geochem. J.*, 14, 167–175, 1980.
- Koide, M., Lee, D. S., and Goldberg, E. D.: Metal and transuranic records in mussel shells, byssal threads and tissues, *Estuarine, Coastal and Shelf Science*, 15, 679–695, 1982.
- Lea, D. W. and Boyle, E. A.: Ba content of benthic foraminifera controlled by bottom water composition, *Nature*, 338, 751–753, 1989.
- Lea, D. W. and Boyle, E. A.: Ba in planktonic foraminifera, *Geochim. Cosmochim. Acta*, 55, 3321–3331, 1991.
- Lea, D. W., Mashiotta, T. A., and Spero, H. J.: Controls on magnesium and strontium uptake in planktonic foraminifera determined by live culturing, *Geochim. Cosmochim. Acta*, 63(16), 2369–2379, 1999.
- Le Cadre, V. and Debenay, J.-P.: Morphological and cytological responses of *Ammonia* (foraminifera) to copper contamination: Implications for the use of foraminifera as bioindicators of pollution, *Environ. Pollut.*, 143, 304–317, 2006.
- Lewis, E. and Wallace, D. W. R.: Program Developed for CO₂ System Calculations, ORNL/CDIAC-105, Carbon Dioxide Information Analysis Center, Oak Ridge National Laboratory, U.S. Department of Energy, Oak Ridge, Tennessee, 1998.

BGD

4, 961–991, 2007

**Copper in
foraminiferal calcite**

L. J. de Nooijer et al.

Title Page

Abstract

Introduction

Conclusions

References

Tables

Figures

◀

▶

◀

▶

Back

Close

Full Screen / Esc

Printer-friendly Version

Interactive Discussion

EGU

- Lu, Y. F. and Allen, H. E.: Characterization of copper complexation with natural dissolved organic matter (DOM) – link to acidic moieties of DOM and competition by Ca and Mg, *Water Res.*, 36, 5083–5101, 2002.
- Mackey, D. J. and Zirino, A.: Comments on trace metal speciation in seawater or do “onions” grow in the sea?, *Anal. Chim. Acta*, 284, 635–647, 1994.
- Mason, P. R. D. and Kraan, W. J.: Attenuation of spectral interferences of during laser ablation inductively coupled plasma mass spectrometry (LA-ICP-MS) using an rf only collision and reaction cell, *J. Anal. At. Spectrom.*, 17, 858–867, 2002.
- Marchitto, T. M.: Lack of a significant temperature influence on the incorporation of Cd into benthic foraminiferal tests, *Geochem. Geophys. Geosyst.*, 5(10), Q10D11, 2004.
- Molina-Cruz, A. and Ayala-López, A.: Influence of the hydrothermal vents on the distribution of benthic foraminifera from the Guaymas Basin, Mexico, *Geo-Mar. Lett.*, 8, 49–56, 1988.
- Morse, J. W. and Bender, M. L.: Partition coefficients in calcite: examination of factors influencing the validity of experimental results and their application to natural systems, *Chem. Geol.*, 82, 265–277, 1990.
- Mucci, A.: The solubility of calcite and aragonite in seawater at various salinities, temperatures, and one atmosphere total pressure, *Am. J. Sci.*, 283, 780–799, 1983.
- Mucci, A. and Morse, J. W.: The incorporation of Mg^{2+} and Sr^{2+} into calcite overgrowths: influences of growth rate and solution composition, *Geochim. Cosmochim. Acta*, 47(2), 217–233, 1983.
- Nassrallah-Aboukaïs, N., Boughriet, A., Fischer, J. C., Wartel, M., Langelin, H. R., and Aboukaïs, A.: Electron paramagnetic resonance (EPR) study of Cu^{2+} and Mn^{2+} ions interacting as probes with calcium carbonate during the transformation of vaterite into cubic calcite, *J. Chem. Soc. Lond. Faraday Trans.*, 92, 3211–3216, 1996.
- Nassrallah-Aboukaïs, N., Boughriet, A., Laureyns, J., Aboukaïs, A., Fischer, J. C., Langelin, H. R., and Wartel, M.: Transformations of vaterite into cubic calcite in the presence of $Cu(II)$ species, *Chem. Mater.*, 10, 238–243, 1998.
- Nürnberg, D., Bijma, J., and Hemleben, C.: Assessing the reliability of magnesium in foraminiferal calcite as a proxy for water mass temperatures, *Geochim. Cosmochim. Acta*, 60(5), 803–814, 1996.
- Parkhurst, D. L. and Appelo, C. A. J.: User’s guide to PHREEQC (Version 2) – A computer program for speciation, batch-reaction, one-dimensional transport, and inverse geochemical calculations: U.S. Geological Survey Water-Resources Investigations Report, 99-4259,

BGD

4, 961–991, 2007

**Copper in
foraminiferal calcite**

L. J. de Nooijer et al.

Title Page

Abstract

Introduction

Conclusions

References

Tables

Figures

◀

▶

◀

▶

Back

Close

Full Screen / Esc

Printer-friendly Version

Interactive Discussion

EGU

1999.

Panieri, G., Bamberi, F., Marani, M., and Barbieri, R.: Benthic foraminifera from a recent, shallow-water hydrothermal environment in the Aeolian Arc (Thyrrhenian Sea), *Mar. Geol.*, 218, 207–229, 2005.

5 Papadopoulos, P. and Rowell, D. L.: The reactions of copper and zinc with calcium carbonate surfaces, *J. Soil Sci.*, 40, 39–48, 1989.

Pearce, N. J. G., Perkins, W. T., Westgate, J. A., Gorton, M. P., Jackson, S. E., Neil, C. R., and Chenery, S. P.: A compilation of new and published major and trace element data for NIST SRM 610 and NIST SRM 612 glass reference materials, *Geost. Newslett.*, 21(1), 115–144, 1997.

10 Pickering, W. F.: Extraction of copper, lead, zinc and cadmium ions sorbed on calcium carbonate, *Water, Air, Soil Pollut.*, 20, 299–309, 1983.

Rathburn, A. E. and DeDeckker, P.: Magnesium and strontium compositions of recent benthic foraminifera from the Coral Sea, Australia and Prydz Bay, Antarctica, *Mar. Micropaleontol.*, 32, 231–248, 1997.

15 Reichart, G. J., Jorissen, F. J., Anschutz, P., and Mason, P. R. D.: Single foraminiferal test chemistry, *Geology*, 31(4), 335–358, 2003.

Rosenthal, Y., Boyle, E. A., and Slowley, N.: Temperature control on the incorporation of magnesium, strontium, fluorine, and cadmium into benthic foraminiferal shells from Little Bahama Bank: prospects for thermocline paleoceanography, *Geochim. Cosmochim. Acta*, 61(17), 3633–3643, 1997.

20 Ruiz, F., González-Regalado, M. L., Borrego, J., Abad, M., and Pendón, J. G.: Ostracoda and foraminifera as short-term tracers of environmental changes in very polluted areas: the Odiel estuary (SW Spain), *Environ. Pollut.*, 129, 49–61, 2004.

25 Sáinz, A. and Ruiz, F.: Influence of the very polluted inputs of the Tinto-Odiel system on the adjacent littoral sediments of southwestern Spain: A statistical approach, *Chemosphere*, 62, 1612–1622, 2006.

Samir, A. M. and El-Din, A. B.: Benthic foraminiferal assemblages and morphological abnormalities as pollution proxies in two Egyptian basins, *Mar. Micropaleontol.*, 41, 193–227, 2001.

30 Segev, E. and Erez, J.: Effect of Mg/Ca ratio in seawater on shell composition in shallow benthic foraminifera, *Geochem. Geophys. Geosyst.*, 7(2), Q02P09, 2006.

Shannon, R. D.: Revised effective ionic radii and systematic studies of interatomic distances in halides and chalcogenides, *Acta Crystallogr. A*, 32, 751–767, 1976.

BGD

4, 961–991, 2007

Copper in foraminiferal calcite

L. J. de Nooijer et al.

Title Page

Abstract

Introduction

Conclusions

References

Tables

Figures

◀

▶

◀

▶

Back

Close

Full Screen / Esc

Printer-friendly Version

Interactive Discussion

EGU

- Sunda, W. G. and Huntsman, S. A.: Regulation of copper concentration in the oceanic nutricline by phytoplankton uptake and regeneration cycles, *Limnol. Oceanogr.*, 40(1), 132–137, 1995.
- Toyofuku, T., Kitazato, H., Kawahata, H., Tsuchiya, M., and Nohara, M.: Evaluation of Mg/Ca thermometry in foraminifera: Comparison of experimental results and measurements in nature, *Paleoceanography*, 15(4), 456–464, 2000.
- 5 Toyofuku, T. and Kitazato, H.: Micromapping of Mg/Ca values in cultured specimens of the high-magnesium benthic foraminifera, *Geochem. Geophys. Geosyst.*, 6(11), Q11P05, 2005.
- Ueno, K., Imamura, T., and Cheng, K. L.: *Handbook of organic analytical reagents*, 2nd ed. CRC Publishers, 1992.
- 10 Wells, A. F.: *Structural inorganic chemistry*, 5th ed, Clarendon, 1984.
- Zeebe, R. E. and Sanyal, A.: Comparison of potential strategies of planktonic foraminifera for house building: Mg²⁺ or H⁺ removal?, *Geochim. Cosmochim. Acta*, 66(7), 1159–1169, 2002.
- Zeebe, R. E. and Wolf-Gladrow, D.: *CO₂ in seawater: Equilibrium, kinetics and isotopes*, Elsevier Oceanographic Series, Amsterdam, 2001.
- 15 Zirino, A. and Yamamoto, S.: A pH-dependent model for the chemical speciation of copper, zinc, cadmium, and lead in seawater, *Limnol. Oceanogr.*, 17(5), 661–671, 1972.

BGD

4, 961–991, 2007

**Copper in
foraminiferal calcite**

L. J. de Nooijer et al.

Title Page

Abstract

Introduction

Conclusions

References

Tables

Figures

◀

▶

◀

▶

Back

Close

Full Screen / Esc

Printer-friendly Version

Interactive Discussion

EGU

Table 1. Target concentrations of Cu in sea water and measured $[Cu]_{\text{seawater}}$ and $(Cu/Ca)_{\text{seawater}}$ at start and end of both experiments. n.a. = not available.

Experiment	Target [Cu] in $\mu\text{mol/l}$	Measured Cu		Cu/Ca	
		at the start of ex- periment in $\mu\text{mol/l} + 1\text{SD}$	at the end of ex- periment in $\mu\text{mol/l} + 1\text{SD}$	at start of experi- ment $\times 10^{-6}$ $+ 1\text{SD} \times 10^{-6}$	at end of experi- ment $\times 10^{-6}$ $+ 1\text{SD} \times 10^{-6}$
1. <i>Ammonia tepida</i>	0	0.0843	0.100±0.00584	6.32±5.28	2.91±0.221
	0.10	0.0960±0.0146	n.a.	2.95±0.384	n.a.
	0.20	0.197±0.00173	0.256±0.00206	6.00±0.0326	7.34±0.121
	0.50	0.332±0.00505	n.a.	10.1±0.0143	n.a.
	10	12.6±0.293	14.0±0.867	378±4.62	408±6.94
	20	17.8±0.489	20.0±3.05	547±13.5	473±20.8
2. <i>Heterostegina depressa</i>	0	0.125±0.00554	0.208±0.0119	1.69±0.0236	3.62±0.343
	0.10	0.744±0.00217	0.211±0.0408	2.32±0.0670	3.74±0.500
	0.20	0.242±0.0228	0.368	4.06±0.179	6.71
	0.50	1.28±0.0384	1.25±0.0522	20.2±0.0120	23.1±0.711
	10	11.2±0.00672	10.4±0.277	172±0.844	212±5.88
	20	n.a.	n.a.	n.a.	n.a.

Title Page

Abstract

Introduction

Conclusions

References

Tables

Figures

◀

▶

◀

▶

Back

Close

Full Screen / Esc

Printer-friendly Version

Interactive Discussion

Title Page

Abstract

Introduction

Conclusions

References

Tables

Figures

◀

▶

◀

▶

Back

Close

Full Screen / Esc

Printer-friendly Version

Interactive Discussion

Table 2. Thermodynamic data for calcein (Ueno et al., 1992). L = calceine. ^a reactions leads to negligible metal binding; ^b values extrapolated by assuming chemical behavior similar to Cu²⁺ (see text for discussion).

	Reaction	Functional groups	Log <i>K</i>
(1)	$L^{6-} + H^+ = HL^{5-}$	–COOH	11.7
(2)	$HL^{5-} + H^+ = H_2L^{4-}$	–COOH	10.8
(3)	$H_2L^{4-} + H^+ = H_3L^{3-}$	–COOH	5.5
(4)	$H_3L^{3-} + H^+ = H_4L^{2-}$	–OH	4.2
(5)	$H_4L^{2-} + H^+ = H_5L^{-}$	≡NH ⁺	2.9
(6)	$H_5L^{-} + H^+ = H_6L$	≡NH ⁺	2.1
(7)	$2Cu^{2+} + L^{6-} = Cu_2L^{2-}$	–COOH	28.9
(8)	$Cu^{2+} + H_2L^{4-} = CuH_2L^{2-}$	≡NH ⁺	8.3
(9)	$Cu^{2+} + H_4L^{2-} + H_3L^{3-} = Cu(H_4L)(H_3L)^{3-}$	–COOH + –OH	10.4 ^a
(10)	$2Ca^{2+} + L^{6-} = Ca_2L^{2-}$	–COOH	27.2 ^b
(11)	$Ca^{2+} + H_2L^{4-} = CaH_2L^{2-}$	≡NH ⁺	6.63
(12)	$Ca^{2+} + H_4L^{2-} + H_3L^{3-} = Ca(H_4L)(H_3L)^{3-}$	–COOH + –OH	8.73 ^{ab}
(13)	$2Mg^{2+} + L^{6-} = Mg_2L^{2-}$	–COOH	28.5 ^b
(14)	$Mg^{2+} + H_2L^{4-} = MgH_2L^{2-}$	≡NH ⁺	7.9
(15)	$Mg^{2+} + H_4L^{2-} + H_3L^{3-} = Mg(H_4L)(H_3L)^{3-}$	–COOH + –OH	10.0 ^{ab}

Table 3. Added and free copper concentrations and activities in the experiments. Suffix t = total copper concentration added to the experiment; suffix c = corrected ratios (total Cu or Ca minus calceine-complexed Cu or Ca).

Experiment	[Cu] _t (μmol/l)	[Cu] _f (μmol/l)	[Cu] _t /[Ca] _t	[Cu] _c (μmol/l)	[Cu] _c (μmol/l)	[Cu] _c /[Ca] _c
1. <i>Ammonia tepida</i>	0.156	0.133	2.6 × 10 ⁻⁵	0.156	0.133	2.6 × 10 ⁻⁵
	0.168	0.146	2.9 × 10 ⁻⁵	0.168	0.146	2.9 × 10 ⁻⁵
	0.384	0.333	6.6 × 10 ⁻⁵	0.384	0.333	6.6 × 10 ⁻⁵
	1.05	0.911	18 × 10 ⁻⁵	1.05	0.911	18 × 10 ⁻⁵
	18.6	16.1	320 × 10 ⁻⁵	18.6	16.1	320 × 10 ⁻⁵
	38.0	32.9	660 × 10 ⁻⁵	38.0	32.9	660 × 10 ⁻⁵
2. <i>Heterostegina depressa</i>	0.167	0.140	1.6 × 10 ⁻⁵	0.167	16	1.6 × 10 ⁻⁵
	0.190	0.163	1.8 × 10 ⁻⁵	0.190	18	1.8 × 10 ⁻⁵
	0.255	0.218	2.4 × 10 ⁻⁵	0.255	24	2.4 × 10 ⁻⁵
	1.36	1.16	13 × 10 ⁻⁵	1.36	130	13 × 10 ⁻⁵
	12.1	10.3	110 × 10 ⁻⁵	12.1	1100	110 × 10 ⁻⁵

Title Page

Abstract

Introduction

Conclusions

References

Tables

Figures

◀

▶

◀

▶

Back

Close

Full Screen / Esc

Printer-friendly Version

Interactive Discussion

Table 4. Number of individuals at the start of the experiments, number of specimens that formed new calcite and total number of added chambers.

Experiment	Target [Cu ²⁺] in μmol/l	Number of specimens at start of experiment	Number of individuals that grew new chambers	Number of new chambers added
1. <i>Ammonia tepida</i>	0	48	7	7
	0.10	48	11	12
	0.20	48	6	6
	0.50	48	3	3
	10	48	5	5
	20	48	1	1
2. <i>Heterostegina depressa</i>	0	24	5	8
	0.10	24	3	6
	0.20	24	0	0
	0.50	24	19	37
	10	24	1	3
	20	24	0	0

Title Page

Abstract

Introduction

Conclusions

References

Tables

Figures

◀

▶

◀

▶

Back

Close

Full Screen / Esc

Printer-friendly Version

Interactive Discussion

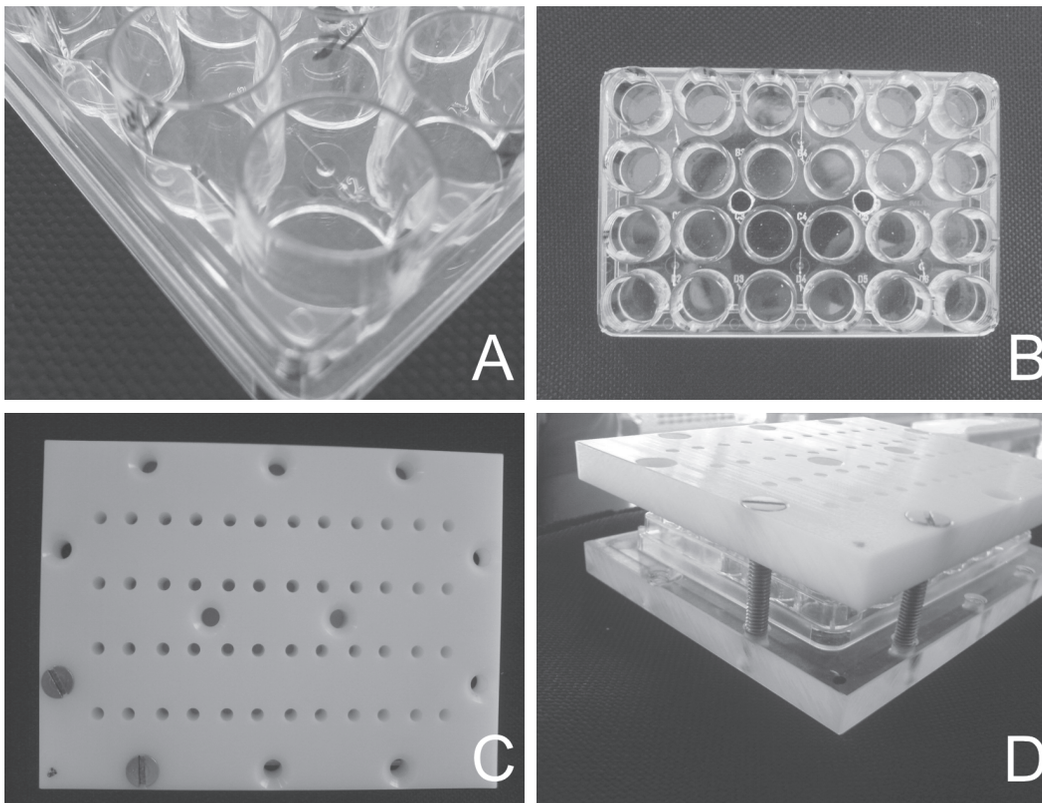


Fig. 1. Design of the experimental set-up. **(A)**: close-up of a culture tray. **(B)**: overview of a culture tray. **(C)** and **(D)**: culture tray between plexiglass lids, with in- and outflow openings in upper lid.

BGD

4, 961–991, 2007

**Copper in
foraminiferal calcite**

L. J. de Nooijer et al.

Title Page

Abstract

Introduction

Conclusions

References

Tables

Figures

◀

▶

◀

▶

Back

Close

Full Screen / Esc

Printer-friendly Version

Interactive Discussion

EGU

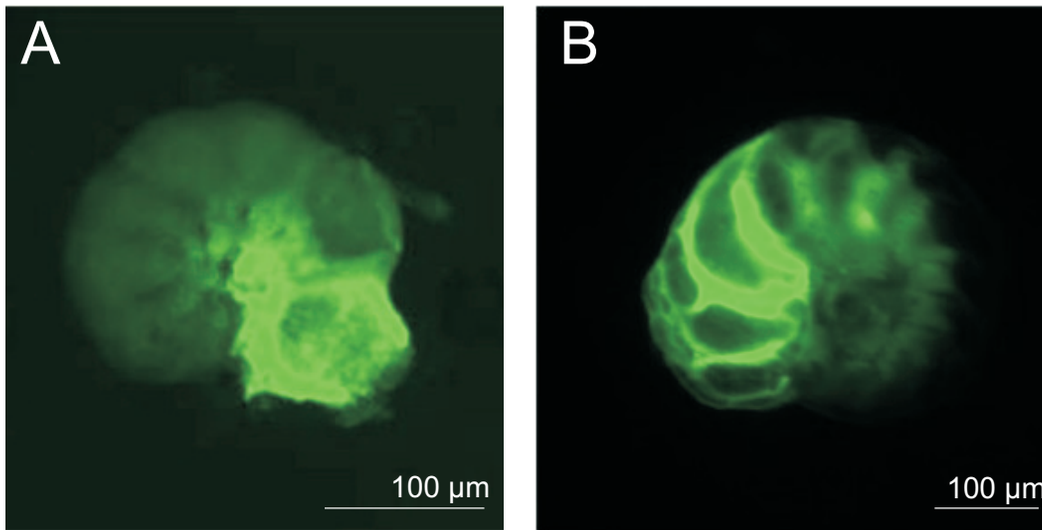


Fig. 2. New chambers added by *Ammonia tepida* (A) and *Heterostegina depressa* (B), visible by fluorescence of incorporated calcein.

BGD

4, 961–991, 2007

**Copper in
foraminiferal calcite**

L. J. de Nooijer et al.

Title Page

Abstract

Introduction

Conclusions

References

Tables

Figures

◀

▶

◀

▶

Back

Close

Full Screen / Esc

Printer-friendly Version

Interactive Discussion

EGU

Copper in
foraminiferal calcite

L. J. de Nooijer et al.

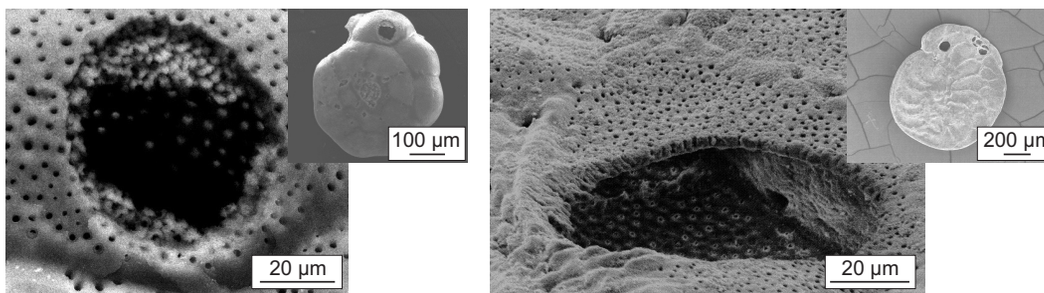


Fig. 3. Scanning electron microscope image of laser ablation craters in *Ammonia* (left) and *Heterostegina* (right). Insets depict the whole specimen. Scale bar is indicated in the lower right corner only for the magnified image.

Title Page

Abstract

Introduction

Conclusions

References

Tables

Figures

◀

▶

◀

▶

Back

Close

Full Screen / Esc

Printer-friendly Version

Interactive Discussion

Copper in
foraminiferal calcite

L. J. de Nooijer et al.

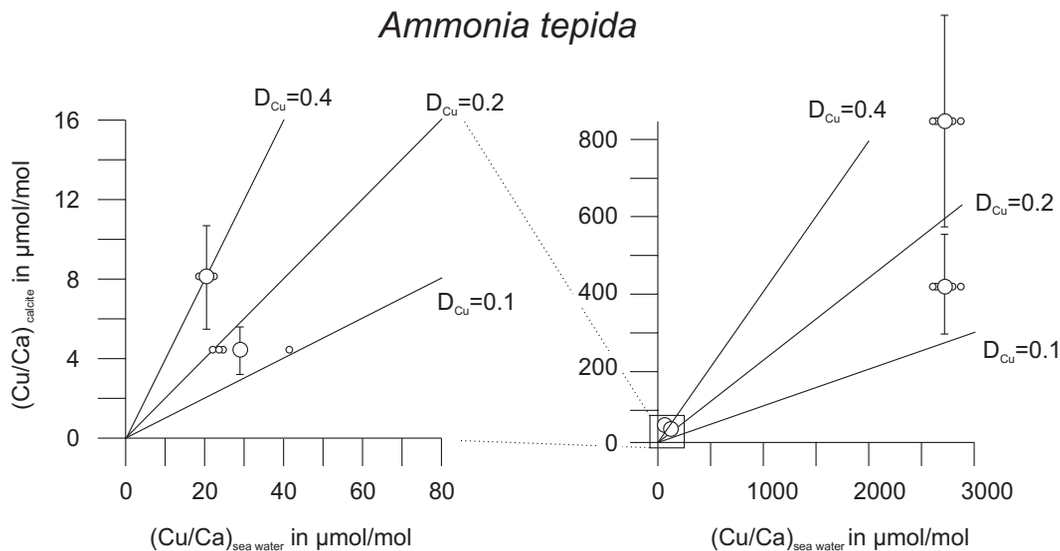


Fig. 4. Cu/Ca ratios in *Ammonia tepida* test carbonate versus Cu/Ca in sea water. Left graph is an enlargement of right one: small circles indicate the separate $[\text{Cu}]_{\text{seawater}}$ measurements (replicate samples before and after the incubation period), large circles represent the average $\text{Cu/Ca}_{\text{seawater}}$ values. The two measurements in the right panel are from chambers of the same specimen. Lines represent partition coefficients of 0.1, 0.2 and 0.4.

Title Page

Abstract

Introduction

Conclusions

References

Tables

Figures

I◀

▶I

◀

▶

Back

Close

Full Screen / Esc

Printer-friendly Version

Interactive Discussion

Copper in
foraminiferal calcite

L. J. de Nooijer et al.

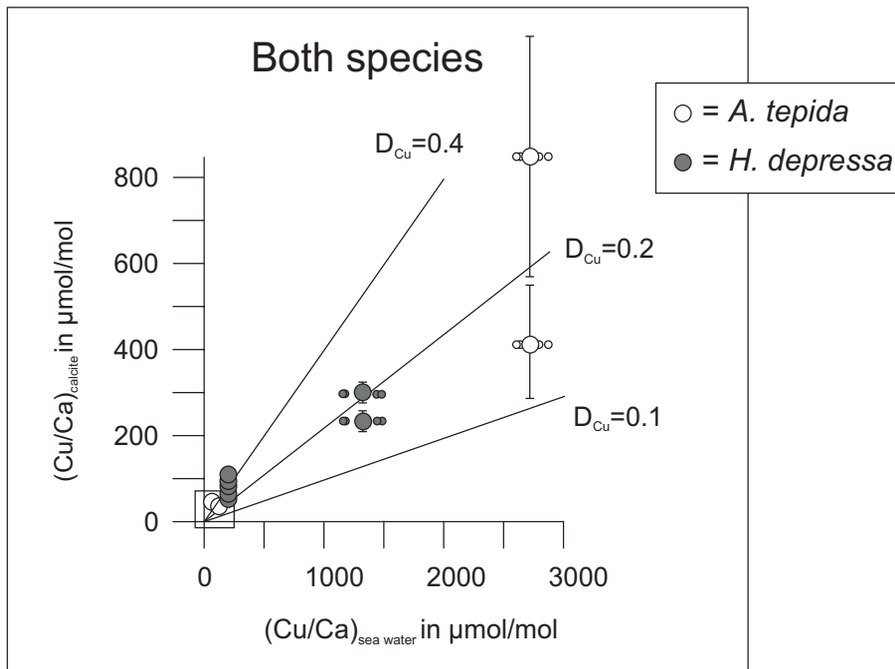


Fig. 5. Plot of Cu/Ca ratios in foraminiferal calcite of the added chamber versus the Cu/Ca of the sea water in which they were incubated. The two *Heterostegina*-measurements in the center of the graph represent two chambers of the same specimen. Lines indicate partition coefficients of 0.1, 0.2 and 0.4.

Title Page

Abstract

Introduction

Conclusions

References

Tables

Figures

I◀

▶I

◀

▶

Back

Close

Full Screen / Esc

Printer-friendly Version

Interactive Discussion