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Can Mg isotopes be used to trace cyanobacteria-mediated magnesium carbonate precipitation in alkaline lakes?

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BGD

8, 6473–6517, 2011

Can Mg isotopes be used to trace cyanobacteria-mediated

L. S. Shirokova et al.

Title Page

Abstract

Introduction

Conclusions

References

Tables

Figures

⏪

⏩

◀

▶

Back

Close

Full Screen / Esc

Printer-friendly Version

Interactive Discussion



Abstract

The fractionation of Mg isotopes was determined during the cyanobacterial mediated precipitation of hydrous magnesium carbonate precipitation in both natural environments and in the laboratory. Natural samples were obtained from Lake Salda (SE Turkey), one of the few modern environments on the Earth's surface where hydrous Mg-carbonates are the dominant precipitating minerals. This precipitation was associated with cyanobacterial stromatolites which were abundant in this aquatic ecosystem. Mg isotope analyses were performed on samples of incoming streams, groundwaters, lake waters, stromatolites, and hydromagnesite-rich sediments. Laboratory Mg carbonate precipitation experiments were conducted in the presence of purified *Synechococcus* sp cyanobacteria that were isolated from the lake water and stromatolites. The hydrous magnesium carbonates nesquehonite ($\text{MgCO}_3 \cdot 3\text{H}_2\text{O}$) and dypingite ($\text{Mg}_5(\text{CO}_3)_4(\text{OH})_2 \cdot 5(\text{H}_2\text{O})$) were precipitated in these batch reactor experiments from aqueous solutions containing either synthetic $\text{NaHCO}_3/\text{MgCl}_2$ mixtures or natural Lake Salda water, in the presence and absence of live photosynthesizing *Synechococcus* sp. Bulk precipitation rates were not affected by the presence of bacteria when air was bubbled through the system. In the stirred non-bubbled reactors, conditions similar to natural settings, bacterial photosynthesis provoked nesquehonite precipitation, whilst no precipitation occurred in bacteria-free systems in the absence of air bubbling, despite the fluids achieving a similar or higher degree of supersaturation. The extent of Mg isotope fractionation ($\Delta^{26}\text{Mg}_{\text{solid-solution}}$) between the mineral and solution in the abiotic experiments was found to be identical, within uncertainty, to that measured in cyanobacteria-bearing experiments, and ranges from -1.4 to -0.7% . This similarity refutes the use of Mg isotopes to validate microbial mediated precipitation of hydrous Mg carbonates.

BGD

8, 6473–6517, 2011

Can Mg isotopes be used to trace cyanobacteria-mediated

L. S. Shirokova et al.

Title Page

Abstract

Introduction

Conclusions

References

Tables

Figures

⏪

⏩

◀

▶

Back

Close

Full Screen / Esc

Printer-friendly Version

Interactive Discussion

1 Introduction

Traditional stable isotopes (C, O, N, S, B) have been used extensively to elucidate biogeochemical processes in aquatic lacustrine environments (e.g. Pentecost and Spiro, 1990; Peterson and Fry, 1987). The stable isotopic composition of “non-traditional” metals such as Ca, Cu, Fe, Li, Mg, Zn are just starting to be used to understanding the processes operating in lake systems. Of these isotope systems, Mg is of particular interest because: (1) it is a major component of the lake water; (2) it is an important component of chlorophyll in aquatic microorganisms (Black et al., 2006, 2007); (3) hydrous Mg-minerals are known to form in lake sediments (Castanier et al., 1993; Power et al., 2009), and (4) stable Mg isotopes fractionate more significantly compared to other alkaline earth metals such as Ca and Sr (Galy et al., 2002; Tipper et al., 2006; Higgins and Schrag, 2010; Li et al., 2011; Schauble, 2011).

One of the most important processes controlling the biogeochemical cycling of Mg in continental waters is carbonate biomineralization (Lowenstam and Weiner, 1989; Dove, 2010). Cyanobacteria-induced mineralization has occurred in both ancient and modern environments since the Precambrian (Kempe and Kazmierczak, 1990; Knoll et al., 1993; Brady et al., 2009; Planavsky et al., 2009; Raven and Giordano, 2009; Riding, 2000; Ries, 2010). Most modern freshwater cyanobacteria-dominated carbonate formation is observed in alkaline aquatic environments with high Ca to Mg ratios (Scholl and Taft, 1964; Müller et al., 1972; Otsuki and Wetzel, 1974; Kelts and Hsü, 1978; Pentecost, 1978; Stabel, 1986; Thompson and Ferris, 1990; Pedone and Folk, 1996; Ferris et al., 1997; Thompson et al., 1997; Kazmierczak and Kempe, 2006; Dupraz et al., 2009; Power et al., 2011), and produces various calcium carbonate minerals. In contrast, the formation of Mg-rich carbonate minerals by cyanobacteria occurs only in specific Earth surface environments, such as Lake Salda in Turkey, which is fed by ultra-mafic rock weathering products (e.g. Braithwaite and Zedef, 1994), alkaline lakes such as those in British Columbia (Renaut, 1990; Power et al., 2007, 2009), and in some saline lake sediments (Renaut and Long, 1989; Renaut and Douglas, 1990; Queralt et

BGD

8, 6473–6517, 2011

Can Mg isotopes be used to trace cyanobacteria-mediated

L. S. Shirokova et al.

Title Page

Abstract

Introduction

Conclusions

References

Tables

Figures

⏪

⏩

◀

▶

Back

Close

Full Screen / Esc

Printer-friendly Version

Interactive Discussion

al., 1997). Interest in these $\text{Mg}(\text{HCO}_3)_2$ -dominated lakes has been recently stimulated by discovery of Mg carbonates on the surface of the Mars, which may provide evidence for the presence of Mg-rich waters (Calvin et al., 1994; Russell et al., 1999; Edwards et al., 2005; Palomba et al., 2009).

5 The reaction rates and mechanisms of Ca-carbonate precipitation associated with cyanobacterial activity, both in liquid suspension (Hartley et al., 1995, Obst and Dittrich, 2006; Obst et al., 2009; Dittrich and Sibling, 2010; Kranz et al., 2010) and in biofilms (Jorgensen et al., 1983; Cox et al., 1989; Hartley et al., 1996) are relatively well understood. In contrast, the main biological and physico-chemical factors control-
10 ling hydrous Mg carbonate precipitation and Mg isotope fractionation in natural waters are still poorly constrained. In this regard, the alkaline Lake Salda (SW Turkey) represents an excellent natural laboratory where contemporary Mg carbonate precipitation can be studied. Previous studies provided a comprehensive understanding of the geology, lithology, biology, and mineral precipitation processes occurring in the lake basin (e.g. Braithwaite and Zedef, 1994, 1996), and suggested that most of the hydro-
15 magnesite ($\text{Mg}_5(\text{CO}_3)_4(\text{OH})_2\cdot 4\text{H}_2\text{O}$) microbialites (stromatolites) developed along the lake coast were formed by cyanobacterial and algal activity (Braithwaite and Zedef, 1994). In this study we sampled Lake Salda waters, sediments, and stromatolites and performed laboratory experiments to characterize the range and mechanisms of Mg
20 isotope fractionation occurring in this lacustrine environment. These results represent the first quantitative experimental and field calibration of Mg isotope fractionation between the aqueous solution and biotically and abiotically formed hydrous magnesium carbonates under conditions similar to the lake water. As such, these results provide the fundamental basis for using magnesium isotopes as a proxy for tracing biomineral-
25 ization processes in past and contemporary continental aquatic environments.

BGD

8, 6473–6517, 2011

Can Mg isotopes be used to trace cyanobacteria-mediated

L. S. Shirokova et al.

Title Page

Abstract

Introduction

Conclusions

References

Tables

Figures

⏪

⏩

◀

▶

Back

Close

Full Screen / Esc

Printer-friendly Version

Interactive Discussion



2 Materials and methods

2.1 Site description

Lake Salda (also known as Salda Gölü in Turkish) is located in SW Turkey and has a surface area close to 45 km² and an average depth of 80 m, with a maximum reported depth of 200 m (see Fig. 1a). Its limnology, geology, and geochemistry has been extensively studied (Schmid, 1987; Braithwaite and Zedef, 1994, 1996; Russell et al., 1999; Zedef et al., 2000; Kazanchi et al., 2004). This lake is a natural analogue for mineral carbonation (e.g. Oelkers et al., 2008), as meteoric waters feeding the lake dissolve adjacent ultramafic rocks and precipitate hydromagnesite in shallow littoral zones. This is similar to the process recently documented in the British Columbia playas (Power et al., 2009). The lake has no outlet and the water level varies annually depending on precipitation and evaporation.

Contemporary hydromagnesite stromatolites are developed in the coastal zone (littoral) of the SW part of the lake, known as Kocaadalar Burnu, where they form three 20 to 100 m² mounds situated ~50 m offshore and rise ~10 m from the lake bottom, reaching within 3–4 m of the lake surface. Similar to Braithwaite and Zedef (1994, 1996), we define all of the modern, actively grown microbialite structures as stromatolites. Underwater diving examination of the deepest part of these mounds showed no evidence of stromatolites below 6–10 m water depth. 1–1.5 m² stromatolites were found in other parts of the lake within 10–20 m of the shoreline (see Fig. 1b). Finally, at the littoral of the north side of the lake, most of the submerged stones were covered by actively forming, non-solidified stromatolites (see Fig. 1c) that had the same surface texture as the larger mounds. Detailed underwater examination of the stromatolite surfaces demonstrated that they are alive and actively growing. The mineral surface was covered by a layer of green algae, diatoms, and cyanobacteria with oxygen bubbles adjacent to the surface of the microbial mats (see Fig. 1d).

BGD

8, 6473–6517, 2011

Can Mg isotopes be used to trace cyanobacteria-mediated

L. S. Shirokova et al.

Title Page

Abstract

Introduction

Conclusions

References

Tables

Figures

⏪

⏩

◀

▶

Back

Close

Full Screen / Esc

Printer-friendly Version

Interactive Discussion

2.2 Lake water and mineral sampling

Water samples were taken from inflowing streams and the lake littoral zone in February 2008, February 2010, and September 2010. A profile of the lake water column (down to 70 m depth) was also collected from the middle of the lake in September 2010 using an Aquatic Research Co. horizontal polycarbonate water sampler, enabling ultraclean sampling in the field (Shirokova et al., 2010). The water samples were immediately filtered through sterile, single-use Sartorius Minisart 0.45 µm acetate cellulose filters. The first 100 ml of the filtrate was systematically discarded. Dissolved oxygen, pH, and temperature were measured on-site with an uncertainty of 5 %, 0.02 units, and 0.5 °C, respectively. Concentrations of dissolved organic carbon (DOC), Cl, SO₄, alkalinity, cations and trace element (TE) were subsequently measured in the laboratory using methods routinely applied for analysis of lake and river water samples (Pokrovsky et al., 2010, 2011; Shirokova et al., 2010; Vasyukova et al., 2010). From 2 to 5 liters of surface water were also filtered on-site using a sterile Nalgene disposable filter unit and 0.22 µm polycarbonate filters. These samples were stored in sterile polypropylene containers and subsequently used for the bacterial culturing. Once filtered, all water samples were stored at 5 °C.

Hydromagnesite samples were collected from the massive stromatolite mounds of Kocaadalar Burnu, other air-exposed stromatolite islands, and parts of stromatolites collected from 4–5 m depth. We also sampled hydromagnesite sand from the beach and carbonate mineral coatings on submerged branches and grasses. A map of the lake with position of sampling points is given in Electronic Supplementary Material ESM-1.

2.3 Culture and characterization of cyanobacteria

A culture of *Synechococcus* sp. cyanobacteria was isolated from the surface of coastal stromatolites sampled in February 2008 from the depth of 1 m at 50 m from the Lake Salda shoreline. Cyanobacterial strains *Chroococcus turgidus*, *Plankothrix*, *Anabaena*,

BGD

8, 6473–6517, 2011

Can Mg isotopes be used to trace cyanobacteria-mediated

L. S. Shirokova et al.

Title Page

Abstract

Introduction

Conclusions

References

Tables

Figures

⏪

⏩

◀

▶

Back

Close

Full Screen / Esc

Printer-friendly Version

Interactive Discussion



2.4 Laboratory precipitation experiments

Laboratory experiments were aimed at precipitating hydrous Mg carbonates in the presence of the cyanobacterial cultures extracted from the Lake Salda stromatolites under controlled conditions. Experiments were performed in 1000 ml sterile borosilicate glass reactors. Two distinct fluid phases were used in the experiments. One is a low-phosphate Cyanobacteria BG-11 freshwater solution medium (Sigma-Aldrich C3061, Rippka et al., 1979); this fluid had the same composition as the BG-11 growth medium other than for its phosphate concentration, which was 50 μM or 10 % that of the traditional BG-11. The second was the cell supernatant (sterile filtered BG-11 media after 14–30 days of culture growth during the stationary phase), into which 0.025–0.030 M MgCl_2 and ~ 0.05 M NaHCO_3 were added. In addition, sterile Lake Salda water, (0.014 M Mg, 0.03 M DIC) amended with low-phosphate BG-11 nutrient components, was used for additional bacterial growth experiments. A summary of these reactive fluid compositions is provided in Table 1.

Several distinct types of biotic experiments were performed at 25 ± 2 °C. Experiments S-Bio-2, S-Bio-5, and S-Bio-7 were performed in reactors that were continuously stirred with a magnetic stirring bar and bubbled with sterile humid air with an average flow rate of 1.5 ± 0.3 L/min. Experiments S-Bio-1, S-Bio-3, S-Bio-4, and S-Bio-6 were performed in reactors that were stirred, but without air bubbling. Experiments S-Bio-8, S-Bio-9, and S-Bio-10 were run without shaking and bubbling. Each of these experiments was performed under continuous fluorescent light of $30 \mu\text{mol photon m}^{-2} \text{s}^{-1}$.

Abiotic, cell-free control experiments were also performed at a variety of conditions. Experiment S-Abio-1 was performed with stirring and air bubbling in the presence of a sterile supernatant of the *Synechococcus* sp. cyanobacteria containing $50 \pm 10 \text{ mg l}^{-1}$ of dissolved organic carbon (DOC) in the form of cell exometabolites in which MgCl_2 and NaHCO_3 were added in concentrations similar to those of the biotic experiments. The supernatant solution used in these abiotic experiments was generated after centrifugation and filtration through a $0.22 \mu\text{m}$ sterile filter of the *Synechococcus* sp. culture

BGD

8, 6473–6517, 2011

Can Mg isotopes be used to trace cyanobacteria-mediated

L. S. Shirokova et al.

Title Page

Abstract

Introduction

Conclusions

References

Tables

Figures

⏪

⏩

◀

▶

Back

Close

Full Screen / Esc

Printer-friendly Version

Interactive Discussion



Can Mg isotopes be used to trace cyanobacteria-mediated

L. S. Shirokova et al.

Title Page

Abstract

Introduction

Conclusions

References

Tables

Figures

⏪

⏩

◀

▶

Back

Close

Full Screen / Esc

Printer-friendly Version

Interactive Discussion

collected ~20 days after attainment of the stationary phase. Abiotic control experiments S-Abio-2, S-Abio-5, and S-Abio-6 were performed without air bubbling in sterile Lake Salda water amended with low-phosphate BG-11 culture media. Experiment S-Abio-4 was performed without air bubbling in a solution having elevated concentrations of Mg and alkalinity (0.05 M MgCl₂ + 0.1 M NaHCO₃). All abiotic experiments were performed in the presence of 0.01 M NaN₃ to prevent potential microbial growth.

One additional experiment, S-f, was run to assess Mg consumption and isotopic fractionation by *Synechococcus* sp. without carbonate precipitation. This experiment was performed in the absence of dissolved MgCl₂ and NaHCO₃ and *Synechococcus* sp. was grown in BG-11 medium. The fluid phase and biomass in this experiment was sampled after 5 days and 3 months of growth.

2.5 Sampling and analyses

30–50 ml aliquots of the homogeneous suspension (containing the fluid, precipitated mineral phase, and cells if present) were sampled periodically from the reactors in a sterile laminar hood box. The optical density and pH were measured in liquid subsamples, whilst the solution supernatants were initially filtered, using 0.22 μm filters, and used for alkalinity, DOC, and Mg concentration measurements. Trace elements were measured without preconcentration by ICP-MS (e.g., Pokrovsky et al., 2010, 2011). Alkalinity was determined by HCl titration using an automatic Schott TitroLine alpha TA10^{plus} titrator with an uncertainty of ±2 % and a detection limit of 5 × 10⁻⁵ M. The DOC content was determined using a Shimadzu TOC-6000 SCN Carbon Total Analyzer with an uncertainty of 3 % and a detection limit of 40 μM. Magnesium concentrations were measured by flame atomic absorption spectroscopy using a Perkin Elmer AAnalyst 400 with an uncertainty of ±2 % and a detection limit of 0.2 μM. pH was measured using a Mettler Toledo combined electrode, with a precision of ±0.01. The uncertainty of biomass concentration determination via optical density is estimated to be ±10 %.

Can Mg isotopes be used to trace cyanobacteria-mediated

L. S. Shirokova et al.

Title Page

Abstract

Introduction

Conclusions

References

Tables

Figures

⏪

⏩

◀

▶

Back

Close

Full Screen / Esc

Printer-friendly Version

Interactive Discussion



Prior to sample characterization by Scanning Electron Microscopy (SEM), organic matter was removed from the solid phases by treating them with 10 % H₂O₂ for 2–3 days at the same pH as the experimental fluids. The residual solids were then thoroughly rinsed with de-ionized water and freeze dried at –55 °C. The mineral phases were then characterized using a Jeol JSM840a SEM, and an INEL CPS 120 X-ray diffractometer using Co_{Kα}, with a scan speed of 0.02 ° s^{–1}. Untreated solids were kept for chemical analysis as described below.

Transmission Electron Microscopy (TEM) analysis was performed using the TEM-SCAN facilities of the University of Toulouse, with a JEOL JEM 2100F unit equipped with a field emission gun (FEG) source and a PGT EDX detector. TEM samples for analyses were prepared by immersing 200 mesh copper grids coated with a carbon film for 10 s in solutions containing live bacteria taken from the experiments that either produced or did not produce mineral precipitates. To minimize the potential effect of crystallization from salts present in reactor solutions on TEM/EDX measurements, 20 ml aliquots were centrifuged at 4000 rpm for 10 min and washed twice with sterile MilliQ water. Grids were dried and covered with a 20 nm carbon layer prior to TEM analysis.

2.6 Magnesium isotope analyses

The Mg isotope compositions of liquid and solid samples collected in the field and produced during laboratory experiments were analyzed according to the procedure described by Mavromatis et al. (2011). Magnesium isotopic ratios were measured using a Thermo-Finnigan “Neptune” Multi Collector Inductively Coupled Plasma Mass Spectrometer (MC-ICP-MS) at the GET (Toulouse, France). Instrumental mass fractionation effects were corrected via sample-standard bracketing, and all results are presented in delta notation with respect to the DSM-3 international reference material (Galy et al., 2001):

$$\delta^x \text{Mg} = \left(\frac{\left(\frac{x \text{Mg}}{24 \text{Mg}} \right)_{\text{sample}}}{\left(\frac{x \text{Mg}}{24 \text{Mg}} \right)_{\text{DSM3}}} - 1 \right) \cdot 1000 \quad (1)$$

where x refers to the Mg mass of interest. The reproducibility of these $\delta^{26}\text{Mg}$ analyses was typically $<0.08\text{‰}$ and was confirmed by replicate analyses of three international Mg reference standards (DSM-3, CAM-1 and OUMg). The isotopic offset between the Mg in the fluid and that incorporated into the solid phase is defined as:

$$\Delta^{26}\text{Mg}_{\text{solid-liquid}} \equiv \delta^{26}\text{Mg}_{\text{solid}} - \delta^{26}\text{Mg}_{\text{liquid}} \quad (2)$$

This value was determined for all samples where both the fluid and solid phases were collected in the present study.

3 Results

3.1 Chemical and Mg isotopic composition of Lake Salda water and minerals

3.2 Hydrochemistry

The major and trace element compositions of all collected samples are listed in the Electronic Supplementary Material (ESM) 1. The Mg concentration, alkalinity, and pH values measured in Lake Salda waters and inflowing streams are in general in agreement with values reported by Braithwaite and Zedef (1994, 1996) and Kazanci et al. (2004). In February 2008 and February 2010, the surface water temperature was around $8\text{--}10\text{°C}$ whereas in September 2010, it varied from 27.5°C at the surface to 13°C at 70 m depth. The most significant stromatolite growth occurs during the summer (Braithwaite and Zedef, 1996) and as such, detailed chemical and isotopic analysis of the lake water composition was performed on water samples collected during September 2010. At this time, pH decreased from 9.20 ± 0.05 at the surface to 9.03 ± 0.03

Can Mg isotopes be used to trace cyanobacteria-mediated

L. S. Shirokova et al.

Title Page

Abstract

Introduction

Conclusions

References

Tables

Figures

⏪

⏩

◀

▶

Back

Close

Full Screen / Esc

Printer-friendly Version

Interactive Discussion



at the bottom of the water column; DOC concentrations ranged from 4.5 to 3.5 mg l⁻¹, whilst Alkalinity and Mg, Ca, and Cl concentrations remained constant in the water column and were equal to 0.032 ± 0.001 mol l⁻¹, 390 ± 5 mg l⁻¹, 4.0 ± 0.1 mg l⁻¹ and 195 ± 5 mg l⁻¹, respectively. The Mg concentration of incoming streams and groundwaters was ~3 times lower than the lake water, and were thereby strongly undersaturated with respect to nesquehonite.

3.2.1 Stromatolite mineralogy

The examined stromatolites were dominated by hydromagnesite, as shown by XRD analysis of multiple spots of a 20 cm-thick representative sample and littoral sediments. The carbonate sand of the littoral zone of the lake (the white deposits shown in Fig. 1a) was also composed of hydromagnesite as it originated from the wave abrasion of growing stromatolites (Braithwaite and Zedef, 1996). The external stromatolite surface had a typical honeycomb-like structure (see Fig. 2a, b), likely formed by the heterotrophic degradation of cyanobacteria cell which are mineralized within their ExoPolySaccharide (EPS) layers (e.g. Dupraz et al., 2004). Platelets of hydromagnesite were clearly seen in the inner parts of stromatolites, which were not exposed directly to lake water (see Fig. 2d). Similar needle-like habits occur on hydromagnesite covering submerged solids such as tree branches (see Fig. 2c). These crystal forms were commonly observed and they are abundant in all investigated hydromagnesite samples. The imprints of *Pyrogira* green algae were often recognized by the SEM on stromatolite surfaces (not shown) as these algae efficiently colonized active stromatolite surfaces forming visible air bubbles (see Fig. 1d).

3.2.2 Mg isotopes

The δ²⁶Mg composition of the water column was homogeneous at the time of sampling (September 2010), and it did not vary between different sites in the littoral zone, or due to the presence of stromatolites (see Table 2). The mean δ²⁶Mg composition of

BGD

8, 6473–6517, 2011

Can Mg isotopes be used to trace cyanobacteria-mediated

L. S. Shirokova et al.

Title Page

Abstract

Introduction

Conclusions

References

Tables

Figures

⏪

⏩

◀

▶

Back

Close

Full Screen / Esc

Printer-friendly Version

Interactive Discussion



the water column in September 2010 was $0.12 \pm 0.04\text{‰}$, but was found to be slightly lighter at the other sampling times; the mean $\delta^{26}\text{Mg}$ composition of the water found to be -0.005 and -0.02‰ in February 2008 and February 2010 respectively. The incoming spring and ground waters had lighter $\delta^{26}\text{Mg}$ compositions ranging between -1.4 and -0.8‰ . The internal and external parts of stromatolites and the littoral sand exhibited $\delta^{26}\text{Mg}$ composition of $-0.99 \pm 0.07\text{‰}$.

3.3 Experimental modeling of hydrous Mg carbonate precipitation in laboratory

The measured chemical composition of the reactive fluids and the mineralogy of precipitated solid phases are listed in the Table ESM-2 of the Electronic Supporting Material and are described in detail below.

3.3.1 Solid phases

X-ray diffraction analysis demonstrated the precipitation of nesquehonite ($\text{MgCO}_3 \cdot 3\text{H}_2\text{O}$), dypingite ($\text{Mg}_5(\text{CO}_3)_4(\text{OH})_2 \cdot 5\text{H}_2\text{O}$), and hydromagnesite ($\text{Mg}_5(\text{CO}_3)_4(\text{OH})_2 \cdot 4\text{H}_2\text{O}$) at distinct times during the experiments. Nesquehonite precipitation was limited to the first 12–23 days of experiments S-Bio-2 and S-Bio-5, but it was present after 48 days in experiment S-Abio-4, which was performed in fluids two times more enriched in Mg and DIC. Two biotic experiments (S-Bio-1 and S-Bio-2) yielded brucite at the end of the experiment with nesquehonite followed by dypingite at the beginning. For the majority of the abiotic (S-Abio-1, S-Abio-3) and some biotic experiments (S-Bio-2, S-Bio-3 and S-Bio-5), dypingite or mixtures of dypingite and hydromagnesite, were the main mineral phases present at the end of experiment. Experiments performed on Lake Salda water without air bubbling often yielded hydromagnesite. SEM images (see Fig. 2e–j) revealed that nesquehonite exhibited a needle-like habit (see Fig. 2e), whilst the dypingite was present as 2 to 10 μm diameter aggregates that grow with time to 5–15 μm rosettes (see Fig. 2f). The hydromagnesite was characterized by 0.5–1 μm size rounded elongated crystals

BGD

8, 6473–6517, 2011

Can Mg isotopes be used to trace cyanobacteria-mediated

L. S. Shirokova et al.

Title Page

Abstract

Introduction

Conclusions

References

Tables

Figures

⏪

⏩

◀

▶

Back

Close

Full Screen / Esc

Printer-friendly Version

Interactive Discussion

forming large associates (see Fig. 2g). Overall, the sequence of precipitation and recrystallization appears to be: nesquehonite (20–120 μm long, $10 \pm 5 \mu\text{m}$ wide needles,) \rightarrow dypingite (5–15 μm rosettes) \rightarrow hydromagnesite 20–40 μm (aggregates).

It is worth noting that the hydromagnesite crystal associates formed in fluids containing *Gloeocapsa* sp. cyanobacteria during experiment S-Bio-8 (see Fig. 2h) were surprisingly similar to the external surface of Lake Salda stromatolites (see Fig. 2a, b), whereas the rosette-like dypingite and hydromagnesite aggregates obtained in experiments with *Synechococcus* sp. cyanobacteria during experiment Exp S-Bio-10 (Fig. 2f) and Exp S-Bio-3 (Fig. 2i) were similar to the natural hydromagnesite coatings of submerged surfaces in Lake Salda (see Fig. 2c). Note the shape of hydromagnesite crystals formed in experiment S-Bio-3 (Fig. 2g) was similar to that of coccoid *Synechococcus* sp. cyanobacteria (see Fig. 3), suggesting some embedding of the cells. Such an embedding of cells and cell associates was evident in experiment Exp S-Bio-11 which was performed with *Gloeocapsa* sp. cyanobacteria cultured 40 days in Mg, HCO_3^- -enriched BG-11 media (see Fig. 2j) and was likely responsible for the typical honeycomb structure of Mg hydrous carbonates formed in the presence of cyanobacteria (see Fig. 2h) as well as in natural stromatolites (see Fig. 2a, b).

The TEM examination of bacteria grown during 2 weeks in low-phosphate BG-11 nutrient media containing initially 0.025 M MgCl_2 + 0.05 M NaHCO_3 evidenced the presence of nanometer-size spherulite crystals assembled in network-like 1–2 μm sized associates usually adjacent to the surfaces of live cells (see Fig. 3a, d, e). In addition, mineral coating of whole cell surfaces (see Fig. 3c) or cell sheaths (see Fig. 3b) was frequently observed. Cells grown in Mg^{2+} , HCO_3^- low nutrient media remained mineral precipitate-free (see Fig. 3f).

3.3.2 Chemical composition of the fluid phase

The temporal evolution of pH and Mg concentration in all experiments as well as the temporal evolution of alkalinity and biomass concentrations during representative experiments are illustrated in Fig. 4. The Mg concentration and alkalinity of the reactive

BGD

8, 6473–6517, 2011

Can Mg isotopes be used to trace cyanobacteria-mediated

L. S. Shirokova et al.

Title Page

Abstract

Introduction

Conclusions

References

Tables

Figures

⏪

⏩

◀

▶

Back

Close

Full Screen / Esc

Printer-friendly Version

Interactive Discussion

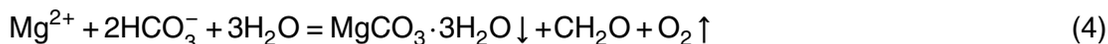


fluids tended to decrease and the pH tended to increase with time during all biotic experiments, notably those with air bubbling (see Fig. 4a, c, e). Abiotic stirred experiments without air bubbling did not produce a measurable decrease in Mg concentration or alkalinity, although a slight pH increase was observed (Fig. 4b, d, f). The stirred biotic experiments both with and without air bubbling produced very similar variations in pH, Mg concentration, and alkalinity (see Fig. 4), though the biomass production was a factor of two higher in experiments with air bubbling (see Fig. 5). The addition of nutrient BG-11 components to Lake Salda water had a significant effect on biomass production and Mg hydrous carbonate precipitation, as evident from the decrease in Mg concentration (see Fig. 6).

The mass of precipitated hydrous magnesium carbonate is plotted as a function of the measured biomass present in the reactor fluid in Fig. 7. The line drawn in this figure, consistent with all the results of all experiments containing biomass can be described by:

$$\text{Mg}_{\text{precipitated}}(\text{mmoles}) = (7.1 \pm 0.9) \times \text{Biomass}_{\text{produced}}(g_{\text{wet}}), r^2 = 0.93 - 0.84 \quad (3)$$

Converting this relationship into molar scale and taking into account that the ratio of wet to dry biomass of *Synechococcus* sp. is 8 ± 2 and that the proportion of carbon in dry biomass is 50% yield the molar inorganic Mg to organic C ratio in reaction product of 1.4 ± 0.2 . This value is compatible with the theoretical $\text{Mg}/\text{C}_{\text{org}}$ ratio of 1 for nesquehonite as first precipitating phase during cyanobacterial photosynthesis:



The speciation and saturation state of the reactive fluids with respect to potentially precipitating mineral phases for all experiments was calculated using PHREEQC software together with its MINTEQA2 database (Parkhurst and Appelo, 1999). The speciation of aqueous Mg during the experiments was dominated by aqueous Mg^{2+} , but also contained significant concentrations of $\text{MgCO}_3^-_{(\text{aq})}$ and MgHCO_3^0 . The evolution of the saturation state of the reactive fluids during the biotic and abiotic experiments with and

BGD

8, 6473–6517, 2011

Can Mg isotopes be used to trace cyanobacteria-mediated

L. S. Shirokova et al.

Title Page

Abstract

Introduction

Conclusions

References

Tables

Figures

⏪

⏩

◀

▶

Back

Close

Full Screen / Esc

Printer-friendly Version

Interactive Discussion



without air bubbling is illustrated in Fig. 8a and b, respectively. The initial fluid supersaturation state with respect to nesquehonite, the first precipitating phase, ranged from 0.2 to 0.6, at the beginning of the experiments, then maximized to 0.8–1.6, before massive mineral precipitation, after which it decreased to 0.1–0.3. The saturation state of these fluids with respect to dypingite was not calculated owing to lack of relevant thermodynamic data. Experiments with lower HCO_3^- concentrations and thus a lower degree of supersaturation (≤ 0.3) failed to produce sufficient mineral precipitation for analysis. The abiotic experiments resulted in mineral precipitation only when air bubbling was applied; the stirred no-bubbling experiments failed to precipitate detectable quantities of hydrous Mg carbonate, despite the fact that $\Omega_{\text{nesquehonite}}$ was similar or even higher than that in the biotic experiments (see Fig. 8b).

Apparent precipitation rates (r_i) were calculated from the first derivative of the fluid phase Mg concentration with respect to time, from the onset of precipitation to the attainment of constant fluid Mg concentrations using:

$$r_i = \frac{dc_{\text{Mg}}}{dt} \quad (5)$$

where c_{Mg} stands for the concentration of Mg in the reactive fluid and t designates time. Experiments performed in the presence of sterile humid air bubbling attained steady-state Mg concentrations over shorter time periods (12 ± 3 days) and exhibited approximately twice higher apparent precipitation rates compared to bubbling-free experiments. Under similar environmental conditions (Mg, alkalinity, bubbling regime), the precipitation rate measured in biotic experiment S-Bio-2 was significantly higher than that measured in its abiotic counterpart (experiment S-Abio-1).

3.3.3 Magnesium isotopic composition

Mg-isotope analyses were performed on selected samples of experiments S-Bio-1, S-Bio-2, S-Abio-1, S-Abio-4, and S-Abio-5 where nesquehonite and dypingite were the main precipitated mineral phases. The Mg isotope compositions for all of the analyzed

BGD

8, 6473–6517, 2011

Can Mg isotopes be used to trace cyanobacteria-mediated

L. S. Shirokova et al.

Title Page

Abstract

Introduction

Conclusions

References

Tables

Figures

⏪

⏩

◀

▶

Back

Close

Full Screen / Esc

Printer-friendly Version

Interactive Discussion



samples are given in Table 3, and the evolution of $\delta^{26}\text{Mg}_{\text{solid-solution}}$ during the experiments is plotted in Fig. 9. All of the experiments exhibited mass-dependent fractionation between the liquid and the solid phase. The precipitated hydrous Mg carbonates had $\delta^{26}\text{Mg}$ compositions that were 0.5–1.4‰ lighter than their liquid counterparts, whilst the precipitated brucite was only 0.1–0.2‰ lighter than the aqueous solution. The isotopic composition of *Synechococcus* sp. cells from the mineral-free experiment was found to be 0.15‰ heavier than the growth medium.

Based on analysis of solution chemical and isotopical composition, there was no evidence for any relationship between the extent of Mg fractionation and the fluid pH, $\text{IAP}_{\text{hydromagnesite}}$, Mg, alkalinity and biomass concentrations. Generally, during the course of experiments in which a single mineral hydrous Mg carbonate phase was precipitated (e.g. S-Bio-5, S-Abio-1), the $\Delta^{26}\text{Mg}_{\text{solid-liquid}}$ values were found to remain constant (see Fig. 9a, b).

4 Discussion

4.1 Comparison of Lake Salda sediments and laboratory precipitates

Although stromatolites from Lake Salda have been extensively studied in the past, the present study confirmed the dominance of hydromagnesite ($\text{Mg}_5(\text{CO}_3)_4(\text{OH})_2\cdot 4\text{H}_2\text{O}$) as the main mineral of these microbialites. The external surface of stromatolites had an extremely porous, void-like, honeycomb-like structure reflecting the presence of bacterial associates or cyanobacterial colonies. The persistence of hydromagnesite both in the lake littoral sediments and in live microbialites confirmed its long-term stability in the lake water. This observation agrees with reports of hydromagnesite's dominance in alkaline playas of British Columbia (Power et al., 2009) and the persistence of hydromagnesite in other alkaline lake sediments (Renaut and Long, 1989; Queralt et al., 1997).

BGD

8, 6473–6517, 2011

Can Mg isotopes be used to trace cyanobacteria-mediated

L. S. Shirokova et al.

Title Page

Abstract

Introduction

Conclusions

References

Tables

Figures

⏪

⏩

◀

▶

Back

Close

Full Screen / Esc

Printer-friendly Version

Interactive Discussion

The bottom waters of the lake and the incoming streams were undersaturated with respect to hydromagnesite and nesquehonite, which suggests the importance of both (i) microbial photosynthetic activity and (ii) surface temperature in creating the fluid supersaturation necessary for mineral precipitation.

In contrast to the natural samples, experiments produced mainly nesquehonite and dypingite in both biotic and abiotic systems. Although the transformation of nesquehonite into hydromagnesite is relatively fast (on the order of weeks) it has only been studied at elevated temperatures (Davies and Bubela, 1973; Hopkinson et al., 2008). These same authors also suggested that dypingite ($\text{Mg}_5(\text{CO}_3)_4(\text{OH})_25(\text{H}_2\text{O})$) may represent an intermediate metastable phase during the transition of nesquehonite to hydromagnesite. This is in agreement with the results of our long-term experiments (S-Bio-8, S-Bio-9 and S-Bio-10) where the transformation of dypingite to hydromagnesite occurred. This long-term transformation can explain the presence of less stable intermediate phases (e.g. nesquehonite and dypingite) in experiments run at 25 °C for only 14 to 42 days. Similarly, Power et al. (2007), reported dypingite formation in biotic mesocosm experiments, and nesquehonite formation in abiotic control experiments performed in the presence of a microbial consortia isolated from Atlin Playas, British Columbia, Canada, at a pH of ~9.5

A striking similarity was observed between hydrous Mg carbonates forming at the surface of live stromatolites in Lake Salda (see Fig. 2b) and those precipitating in long-term laboratory experiments in the presence of *Gloeocapsa* sp. (see Fig. 2h), as well as in numerous field observations (e.g. Fig. 9c and Fig. 11 in Dupraz et al., 2004). Dupraz et al. (2009) observed that discontinuous EPS calcification generates a micropeloidal structure resulting from the presence of coccoid clusters or filamentous bacteria remnants. Furthermore, these authors reported that no precipitation is observed in or on the sheaths of cyanobacteria, and only a negligible precipitation is directly associated with the inner layers of the active filamentous cyanobacteria mats. Instead, precipitation occurs at the uppermost mat layer, which is composed of EPS, empty filamentous bacteria, and coccoids (*Gloeocapsa* spp.). Results of the present study corroborates

Can Mg isotopes be used to trace cyanobacteria-mediated

L. S. Shirokova et al.

Title Page

Abstract

Introduction

Conclusions

References

Tables

Figures

⏪

⏩

◀

▶

Back

Close

Full Screen / Esc

Printer-friendly Version

Interactive Discussion



the mechanism of honeycomb - like structure formation via hydrous Mg carbonate precipitation due to the presence of EPS and cell capsules.

The amount of precipitated hydrous Mg carbonate and biomass production were highly correlated ($r^2 \sim 0.9$) in our laboratory experiments which was consistent with theoretical mineral yield during the biomass production (see Fig. 7). Moreover, the mineral yield due to microbial photosynthesis and, thus, mineral precipitation rates, were very similar among different experiments despite the fact that the experiments were performed in various distinct fluids, durations, biomass concentration, and air bubbling regimes. Given the large variety of investigated conditions, we suggest that the linear relationship shown in Fig. 7 (Eq. 3) can be used for the quantitative prediction ($\pm 25\%$) of the amount of hydrous Mg carbonate precipitation in the presence of photosynthesizing cyanobacteria.

4.2 The mechanisms of Mg carbonate precipitation in the presence of cyanobacteria

Carbonate mineral formation in the presence of photosynthetic bacteria has been attributed to the alkaline environment produced due to release of hydroxyl ions as a result of photosynthesis (Thompson and Ferris, 1990; Douglas and Beveridge, 1998). pH was observed to increase in all experiments, although this increase was almost negligible in abiotic experiments performed without air bubbling. In the abiotic experiments, this pH increase originated from the degassing of the initial reactive fluid which contained $3\text{--}5 \times 10^{-2} \text{ mol kg}^{-1} \text{ NaHCO}_3$ at pH $\sim 8.2\text{--}9.2$. Owing to this high aqueous bicarbonate content, these initial fluids had a $p\text{CO}_2$ of $\sim (0.3\text{--}10) \times 10^{-2.0} \text{ atm}$, which is supersaturated with respect to the atmosphere. The bubbling of sterile humid air liberated CO_2 from the reactive fluid leading to both an increase in pH and degree of supersaturation with respect to Mg carbonate minerals. In the biotic experiments, this $p\text{CO}_2$ decrease was accompanied by an additional pH increase due to photosynthetic uptake of HCO_3^- ions and OH^- release.

BGD

8, 6473–6517, 2011

Can Mg isotopes be used to trace cyanobacteria-mediated

L. S. Shirokova et al.

Title Page

Abstract

Introduction

Conclusions

References

Tables

Figures

⏪

⏩

◀

▶

Back

Close

Full Screen / Esc

Printer-friendly Version

Interactive Discussion

Can Mg isotopes be used to trace cyanobacteria-mediated

L. S. Shirokova et al.

Title Page

Abstract

Introduction

Conclusions

References

Tables

Figures

⏪

⏩

◀

▶

Back

Close

Full Screen / Esc

Printer-friendly Version

Interactive Discussion



It is widely accepted that actively metabolizing microbial cells can avoid calcium carbonate mineral encrustation and cell entombment (Thompson et al., 1997; Aloisi et al., 2006; Bontognali et al., 2008; Martinez et al., 2010) via cell electric potential regulation, S-layer formation, and extracellular EPS generation (e.g., Krumbein et al., 1977; Chafetz and Buczynski, 1992; Arp et al., 1999a, b; Martinez et al., 2008). Unlike biocalcification (Dupraz et al., 2009; Martinez et al., 2010; Bundeleva et al., 2011), hydrous Mg carbonate formation does not appear to trigger a cell protection mechanism. The presence of hydrous Mg carbonate nanoclusters at the surface of live cells, detectable in the TEM images (see Fig. 3a, d, e), and the complete cell encrustation by precipitated mineral seen by the SEM (Fig. 2h, j) suggests a significant role of the cell surface in governing hydrous Mg carbonate formation in the presence of cyanobacteria.

An additional difference between calcite versus hydrous Mg carbonate nucleation at the vicinity of photosynthesizing cells may be the different size of forming crystals. Unlike calcite or dolomite submicron nano-globules (e.g. Aloisi et al., 2006; Bontognali et al., 2008; Spadafora et al., 2010), the first precipitated hydrous Mg carbonate, nesquehonite, are 50 to 200 μm length crystals (Fig. 2l) and as such they cannot encrust the small *Synechococcus*. Similarly, rosette-like dypingite crystal aggregates (Fig. 2f, i), formed at the second stage of experiment are also too large to cover completely cyanobacterial cells at the beginning of incrustation. Only at the stage of hydromagnesite formation at the end of experiment (Fig. 2h, j) does massive embedding and encrusting becomes possible due to (i) the small size of hydromagnesite platelets, and (ii) the presence of dead cells which are easier to mineralize (e.g., Chafetz and Buczynski, 1992; Martinez et al., 2010).

4.3 Mg isotope fractionation

The extent of Mg isotope fractionation between cells and aqueous solutions ($\delta^{26}\text{Mg}_{\text{solid-solution}}$) due to cell surface adsorption and intracellular uptake during *Synechococcus* sp. Growth was equal to 0.152 ‰. This value is similar to the fractionation factor observed during *Gloeocapsa* sp. cyanobacteria growth (Mavromatis et al., 2011).

Black et al. (2006) reported that chlorophyll-*a*, isolated from cyanobacterium *S. elongates*, preferentially incorporates light Mg isotopes with a fractionation factor of -0.71 to -0.53% depending on the growth stage. However, the chlorophyll represents small fraction of total cell Mg, compared to the cytoplasm. As such, there may be significant variation of the Mg isotopic ratio in different cell fractions, with cytoplasm being “heavier” than the chlorophyll. Unlike other essential metals, Mg is not exchanged during the cell division, but once taken up by the cells, remains inside them (Silver and Walderhaug, 1992). This explains the light Mg isotope composition of mineral-free biomass, collected after 5 days and 3 months (see Table 3).

During the abiotic experiment S-Abio-1 in the presence of air bubbling, the precipitated dypingite exhibited a $\delta^{26}\text{Mg}_{\text{solid-solution}}$ value between -1.4% and -1.25% , whereas abiotic experiment S-Abio-4 without bubbling produced only nesquehonite with $\delta^{26}\text{Mg}_{\text{solid-solution}} = -0.47\%$. In contrast, the biotic experiment with bubbling (experiment S-Bio-5) yielded nesquehonite slightly lighter than its abiotic counterpart and depleted in ^{26}Mg compared to the fluid phase. Finally, biotic experiments produced brucite with $\delta^{26}\text{Mg}_{\text{solid-solution}} = -0.2\%$ to -0.1% . Overall, we observed a systematic enrichment of minerals by isotopically light Mg in the order brucite < nesquehonite < dypingite with dypingite fractionation factors similar to that reported for hydrous Mg carbonates formation in the presence *Gloeocapsa* sp. cyanobacteria (Mavromatis et al., 2011). The preferential incorporation of light Mg isotopes in the precipitating solids is consistent with previous Mg isotope analyses on biogenic skeletal carbonates (Chang et al., 2004; Buhl et al., 2007; Hippler et al., 2009) and abiotically precipitated low Mg-calcite (Galy et al., 2002; Immenhauser et al., 2010).

The $\Delta^{26}\text{Mg}_{\text{solid-liquid}}$ values were almost constant as a function of elapsed time (see Fig. 9a) and the fraction of Mg remaining in solution (see Fig. 9b). The nearly constant $\Delta^{26}\text{Mg}_{\text{solid-liquid}}$ values for each mineral phase were consistent with a closed-system equilibrium fractionation model. This suggests a continuous isotopic exchange/equilibration between the precipitated hydrous Mg-carbonates and the reactive fluid (Criss, 1999). Taking into account earlier results for dypingite (Mavromatis

BGD

8, 6473–6517, 2011

Can Mg isotopes be used to trace cyanobacteria-mediated

L. S. Shirokova et al.

Title Page

Abstract

Introduction

Conclusions

References

Tables

Figures

⏪

⏩

◀

▶

Back

Close

Full Screen / Esc

Printer-friendly Version

Interactive Discussion

et al., 2011) and for other hydrous Mg carbonates (this work), it can be assumed that hydrous Mg-carbonate minerals are in a continuous isotopic equilibrium with the fluid phase in aquatic environments. This conclusion is consistent with our observations that the precipitated magnesium phase transforms several times during the temporal evolution of each experiment. The exchange does not seem to be appreciably affected by physicochemical factors such as pH, $p\text{CO}_2$, Mg and DIC concentration or biological activity

The isotopic shift between solid and solution observed in the laboratory is in close agreement with that found in the Salda lake ecosystem (see Fig. 10). As Salda lake is a closed basin, assuming the Mg concentration of the lake water is constant, the mass of Mg arriving to the lake must equal that precipitated in the form of magnesium carbonates. Consequently, mass balance requires that the precipitated hydromagnesite exhibit the same Mg isotopic offset as that between the incoming streams and groundwaters versus the Salda lake water. This is exactly what was observed. A $\delta^{26}\text{Mg}$ offset of 1.0–1.4‰ was observed between the Salda lake water and the incoming streams and groundwaters; both naturally forming and laboratory synthesized magnesium carbonates were 1.0–1.4‰ lighter than their corresponding fluid phase. It is important to note that the presence of biofilms and other, heterotrophic bacteria in natural stromatolites apparently, had an insignificant effect on the overall isotopic fractionation factor compared to the laboratory cyanobacteria monocultures.

A very significant observation in this study is that the Mg isotope fractionation found in abiotic precipitation experiments was nearly identical to that found in biotic experiments and in the natural system. This similarity refutes the use of Mg isotopes to validate microbial mediated precipitation of hydrous Mg carbonates. Similar, if our observation of continuous Mg isotope exchange between hydrous Mg-carbonates and aqueous solutions is general, the Mg-isotopic signature of hydrous Mg-carbonates cannot be used as a paleoproxy tool. Furthermore, assuming that the observed closed isotopic equilibrium indicates a dissolution/re-precipitation process, a similar isotope exchange mechanism for oxygen and carbon may also occur.

Can Mg isotopes be used to trace cyanobacteria-mediated

L. S. Shirokova et al.

Title Page

Abstract

Introduction

Conclusions

References

Tables

Figures



Back

Close

Full Screen / Esc

Printer-friendly Version

Interactive Discussion



5 Conclusions

The hydrochemistry of Lake Salda and field observations suggested a strong microbial control on hydrous Mg carbonate precipitation in the upper layers of the lake. Laboratory experiments with cyanobacteria culture *Chroococales* sp. isolated from live stromatolite resulted in the precipitation of the hydrous magnesium carbonates nesquehonite and dypingite under various laboratory conditions. We discovered a universal dependence between the amount of Mg carbonate precipitated and bacterial biomass production, consistent with theoretical ratio of mineral precipitation during photosynthesis. This relationship can be used to reconstruct paleoproductivity based on the amount of accumulated Mg carbonates. All laboratory experiments yielded similar bulk hydrous magnesium carbonate precipitation rates, although the pH of the reactive fluid was 0.5–1.0 units higher in biotic compared to abiotic experiments. Taken together these observations suggest that the presence of cyanobacteria is required to increase solution pH and thus the fluid supersaturation state as well as to provide nucleation sites at the cell surface. No cell protection mechanisms was evident to avoid cells encrusting and entombment by precipitating hydrous Mg carbonates. The difference between the identity of mineral precipitates and solution ^{26}Mg concentration obtained from abiotic experiments were similar within uncertainty of those obtained in the presence of *Synechococcus* sp. and in previous experiments performed in the presence of *Gloeocapsa* sp. cyanobacteria. Considering that similar $\Delta^{26}\text{Mg}_{\text{solid-liquid}}$ values were observed between inflow waters and hydromagnesitic stromatolites of Lake Salda, it can be inferred that a continuous exchange of Mg-isotopes occurred between the solid and liquid phase. This conclusion suggests that the Mg isotopic composition of hydrous Mg carbonates cannot be used as a paleoproxy tool.

Supplementary material related to this article is available online at:
[http://www.biogeosciences-discuss.net/8/6473/2011/
bgd-8-6473-2011-supplement.pdf](http://www.biogeosciences-discuss.net/8/6473/2011/bgd-8-6473-2011-supplement.pdf).

BGD

8, 6473–6517, 2011

Can Mg isotopes be used to trace cyanobacteria-mediated

L. S. Shirokova et al.

Title Page

Abstract

Introduction

Conclusions

References

Tables

Figures

⏪

⏩

◀

▶

Back

Close

Full Screen / Esc

Printer-friendly Version

Interactive Discussion

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Can Mg isotopes be used to trace cyanobacteria-mediated

L. S. Shirokova et al.

Title Page

Abstract

Introduction

Conclusions

References

Tables

Figures



Back

Close

Full Screen / Esc

Printer-friendly Version

Interactive Discussion



Can Mg isotopes be used to trace cyanobacteria-mediated

L. S. Shirokova et al.

Title Page

Abstract

Introduction

Conclusions

References

Tables

Figures

⏪

⏩

◀

▶

Back

Close

Full Screen / Esc

Printer-friendly Version

Interactive Discussion



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Can Mg isotopes be used to trace cyanobacteria-mediated

L. S. Shirokova et al.

Title Page

Abstract

Introduction

Conclusions

References

Tables

Figures

⏪

⏩

◀

▶

Back

Close

Full Screen / Esc

Printer-friendly Version

Interactive Discussion



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Can Mg isotopes be used to trace cyanobacteria-mediated

L. S. Shirokova et al.

Title Page

Abstract

Introduction

Conclusions

References

Tables

Figures

◀

▶

◀

▶

Back

Close

Full Screen / Esc

Printer-friendly Version

Interactive Discussion



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L. S. Shirokova et al.

Title Page

Abstract

Introduction

Conclusions

References

Tables

Figures

◀

▶

◀

▶

Back

Close

Full Screen / Esc

Printer-friendly Version

Interactive Discussion

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L. S. Shirokova et al.

Title Page

Abstract

Introduction

Conclusions

References

Tables

Figures

⏪

⏩

◀

▶

Back

Close

Full Screen / Esc

Printer-friendly Version

Interactive Discussion



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Can Mg isotopes be used to trace cyanobacteria-mediatedL. S. Shirokova et al.

Title Page

Abstract

Introduction

Conclusions

References

Tables

Figures

◀

▶

◀

▶

Back

Close

Full Screen / Esc

Printer-friendly Version

Interactive Discussion



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Can Mg isotopes be used to trace cyanobacteria-mediated

L. S. Shirokova et al.

Title Page

Abstract

Introduction

Conclusions

References

Tables

Figures

⏪

⏩

◀

▶

Back

Close

Full Screen / Esc

Printer-friendly Version

Interactive Discussion

Can Mg isotopes be used to trace cyanobacteria-mediated

L. S. Shirokova et al.

Title Page

Abstract

Introduction

Conclusions

References

Tables

Figures



Back

Close

Full Screen / Esc

Printer-friendly Version

Interactive Discussion



Table 1. Experimental conditions of all experiments performed in this study. All abiotic experiments were conducted in the presence of 0.01 M NaN_3 .

Experiment	Medium	Duration (days)	conditions	Biomass range, σ_{wet} /L	pH range	Mg range (mM)	DOC, mg L^{-1}	Rate (Mg slope) mmol day^{-1}	Solid phase
S-BIO-1	BG-11	43	Stirring, no bubbling	0.1–3.1	8.7–10.6	25.6–2.2	17–62	–0.86	Dypingite+brucite
S-BIO-2	BG-11	30	Stirring, bubbling	0.2–3.7	8.2–10.5	30.8–4.1	60	–1.22	Nesquehonite, Dypingite, brucite
S-BIO-3	BG-11 on Salda lake water	34	Stirring, no bubbling	0.2–2.2	9.2–10.8	15–2.5	7–18	–0.38**	Dypingite, hydromagnesite
S-BIO-4	BG-11 on Salda lake water	36	Stirring, no bubbling	0.1–3.0	9.3–10.8	14–3	9–17	–0.435**	Dypingite, hydromagnesite
S-BIO-5	BG-11	30	Stirring, bubbling	0.4–2.8	8.2–10.4	31–7	23–29	–0.71	Nesquehonite, dypingite
S-BIO-6	Salda lake water w/o BG-11	34	Stirring, no bubbling	0.2–0.9	9.2–9.6	9.5–12		No precipitation	
S-BIO-7	0.05 M MgCl_2 + 0.005 M NaHCO_3	31	Stirring, bubbling	0.05–3.0	8.1–10.0	52–40		No precipitation	
S-BIO-8*	BG-11: 0.025 M MgCl_2 + 0.036 M NaHCO_3	100	No stirring, no bubbling	0.05–4.0	8.4–10.4	25–5	20–120	N.D.	Hydromagnesite
S-BIO-9	BG-11: 0.025 M MgCl_2 + 0.036 M NaHCO_3	100	No stirring, bubbling	0.05–3.5	8.4–10.7	25–5	10–65	N.D.	Hydromagnesite
S-BIO-10	BG-11: 0.025 M MgCl_2 + 0.036 M NaHCO_3	100	No stirring, no bubbling	0.05–3.5	8.5–10.8	25–5	10–40	N.D.	Hydromagnesite
S-BIO-11*	BG-11: 0.025 M MgCl_2 + 0.05 M NaHCO_3	40	Stirring, no bubbling	0.2–2.2	8.7–10.6	25–4	6.2–90	N.D.	Dypingite, brucite
S-ABIO-1	Supernatant	25	Stirring, bubbling		8.15–9.3	23–9	90–30	–0.50	
S-ABIO-2	BG-11 on Salda lake water	35	Stirring, no bubbling	0	9.2–9.39	13–12	7.4	No precipitation	
S-ABIO-3	Supernatant	25	Stirring, no bubbling	0	8.1–9.25	37–13	30	–1.39	Dypingite
S-ABIO-4	0.05 M MgCl_2 + 0.1 M NaHCO_3	48	Stirring, no bubbling	0	8.33–8.90	52–27		–0.67	Nesquehonite, dypingite
S-ABIO-5	BG-11 on Salda lake water	35	Stirring, no bubbling	0	8.2–9.46	21–22		No precipitation	
S-ABIO-6	BG-11 on Salda lake water	45	Stirring, no bubbling	0	9.3–9.4	13.1–13.8		No precipitation	

* *Gloeocapsa* sp. Culture;

** The rates are significantly lower due to lower initial Mg concentration.

Can Mg isotopes be used to trace cyanobacteria-mediated

L. S. Shirokova et al.

Discussion Paper | Discussion Paper | Discussion Paper | Discussion Paper | Discussion Paper

Title Page

Abstract Introduction

Conclusions References

Tables Figures

⏪ ⏩

◀ ▶

Back Close

Full Screen / Esc

Printer-friendly Version

Interactive Discussion

Table 2. Mg isotopic composition of natural samples.

Sample description	LIQUIDS				SOLIDS				$\Delta^{26}\text{Mg}_{\text{solid-liquid}}$	Mineralogy
	$\delta^{26}\text{Mg}$	2σ	$\delta^{26}\text{Mg}$	2σ	$\delta^{26}\text{Mg}$	2σ	$\delta^{26}\text{Mg}$	2σ		
T3 Salda Lake coast and stromatolites	0.1337	0.0162	0.2569	0.0721	-0.365	0.048	-0.696	0.036	-0.952	Hydromagnesite
Coastal water and carbonate sediments	-0.0637	0.0772	-0.1360	0.1509	-0.504	0.066	-0.993	0.154	-0.857	Hydromagnesite
Stromatolite – Interior part					-0.532	0.041	-1.038	0.032	-1.118*	Hydromagnesite
Stromatolite – Exterior part					-0.479	0.004	-0.938	0.023	-1.018*	Hydromagnesite
Salda-Water February 2010	-0.0053	0.0077	-0.0208	0.0046						
Live stromatolites + bacteria					-0.395	0.026	-0.754	0.018		
Lake depth profile:										
T07-0	0.075	0.017	0.150	0.010						
T07-20	0.035	0.044	0.094	0.049						
T07-40	0.100	0.031	0.167	0.031						
T07-60	0.040	0.034	0.117	0.080						
T07-70	0.079	0.011	0.137	0.063						
T08, Stromatolite islands	0.050	0.024	0.084	0.027						
T11, 1.5 m depth, fragm. stromatolites	0.045	0.024	0.080	0.042						
T09, Incoming spring	-0.733	0.042	-1.402	0.053						
T10, Spring under mountain, Yuoruk MZL	-0.535	0.101	-1.010	0.107						
T1, Coastal lake water, Feb 2008	-0.019	0.046	-0.005	0.070						
T2 Incoming spring, Feb 2008	-0.384	0.049	-0.759	0.028						
Epsomite deposits at the wood			0.1000		-0.183	0.023	-0.340	0.047	-0.440	Epsomite

* Assuming $\delta^{26}\text{Mg}$ in Salda lake water of 0.08‰.



Can Mg isotopes be used to trace cyanobacteria-mediated

L. S. Shirokova et al.

Table 3. Mg isotopic composition of laboratory samples.

Expt.	Sample No.	Time (Days)	LIQUIDS			SOLIDS			$\Delta^{26}\text{Mg}$		Mineralogy	
			$\delta^{25}\text{Mg}$	2σ	$\delta^{26}\text{Mg}$	2σ	$\delta^{25}\text{Mg}$	2σ	$\delta^{26}\text{Mg}$	2σ		solid-liquid
S-Bio-1	1	0	-0.270	0.034	-0.524	0.050						
	8	13	0.048	0.060	-0.017	0.056						
	13	37	-0.091	0.055	-0.462	0.026	-0.050	0.071	-0.567	0.005	-0.105	Brucite
S-Bio-2	6	7	-0.080	0.059	-0.168	0.102	-0.807	0.032	-1.573	0.016	-1.406	Nesquehonite
	10	15	-0.409	0.072	-0.757	0.121	-0.565	0.040	-1.067	0.036	-0.310	Dypingite
	14	26	-0.233	0.050	-0.447	0.026	-0.366	0.010	-0.699	0.041	-0.252	dypingite+brucite
S-ABIO-1	4	6	-0.074	0.041	-0.103	0.033	-0.738	0.016	-1.428	0.005	-1.325	Dypingite
	8	13	0.113	0.036	0.213	0.011	-0.590	0.032	-1.191	0.020	-1.404	Dypingite
	13	25	0.090	0.028	0.195	0.031	-0.555	0.023	-1.061	0.030	-1.256	Dypingite
S-BIO-5	4	10	-0.068	0.068	-0.165	0.102	-0.467	0.017	-0.965	0.055	-0.800	nesquehonite
	9	20	-0.025	0.030	-0.101	0.051	-0.368	0.002	-0.769	0.129	-0.668	nesquehonite
	11	26	0.184	0.030	0.361	0.073	-0.332	0.051	-0.705	0.087	-1.066	nesquehonite
S-ABIO-4	10	40	-0.206	0.026	-0.382	0.063	-0.418	0.045	-0.849	0.094	-0.468	nesquehonite
S-f-5 culture	5 days 3 months	5	-0.474	0.046	-0.973	0.052	-0.385	0.052	-0.821	0.067	0.152	
								-0.582	0.005	-1.184	0.019	-0.211

Title Page

Abstract

Introduction

Conclusions

References

Tables

Figures

⏪

⏩

◀

▶

Back

Close

Full Screen / Esc

Printer-friendly Version

Interactive Discussion



Fig. 1. (A): View of the Salda lake coast. The white and light blue layers in the littoral zone originate from hydromagnesite sand formed due to modern and paleostromatolites wave abrasion. (B): modern stromatolite formations in the littoral zone. (C): stromatolite coatings of the peridotite rock debris taken from a depth of 1–1.5 m in the littoral zone. (D): the surface of active stromatolite covered by *Pyrogira* algae and diatoms with oxygen bubbles illustrating active on-going photosynthesis.

Can Mg isotopes be used to trace cyanobacteria-mediated

L. S. Shirokova et al.

Title Page

Abstract

Introduction

Conclusions

References

Tables

Figures

◀

▶

◀

▶

Back

Close

Full Screen / Esc

Printer-friendly Version

Interactive Discussion

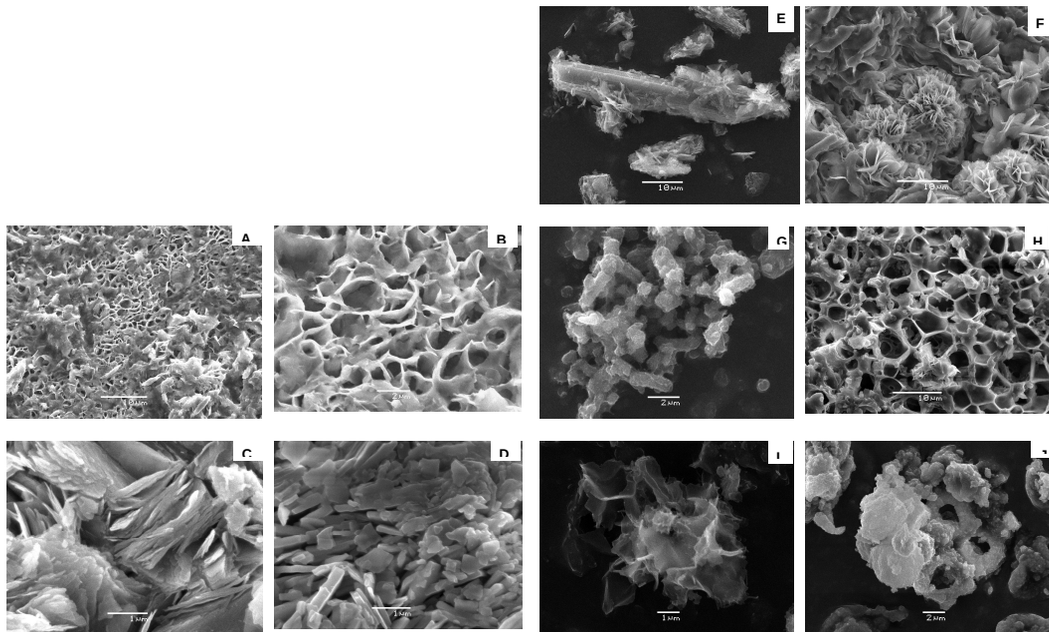


Fig. 2. SEM images of natural stromatolites and experimentally precipitated minerals: **(A–D)**, natural samples; **(E–J)**, laboratory precipitates. **(A)**: Sample T3-1 from the coastal zone, **(B)** close-up showing the external surface of this stromatolite **(C)**: 2–10 mm thick mineral coatings taken from grasses and tree branches in the littoral zone water. **(D)**: interior part of the microbialite. EDS analyses demonstrated the presence of only C, Mg, and O in all the natural samples shown in **A–D**. **(E–J)**: nesquehonite and dypingite precipitates recovered from experiment S-Abio-4, **(F)** dypingite recovered from experiment S-Bio-10, **(G)**, hydromagnesite recovered from experiment S-Bio-3 **(H)**, hydromagnesite recovered from experiment S-Bio-8, **(I)** dypingite recovered from experiment S-Bio-3, and **(J)** dypingite and hydromagnesite recovered from experiment S-Bio-11.

Can Mg isotopes be used to trace cyanobacteria-mediated

L. S. Shirokova et al.

Title Page

Abstract

Introduction

Conclusions

References

Tables

Figures

◀

▶

◀

▶

Back

Close

Full Screen / Esc

Printer-friendly Version

Interactive Discussion



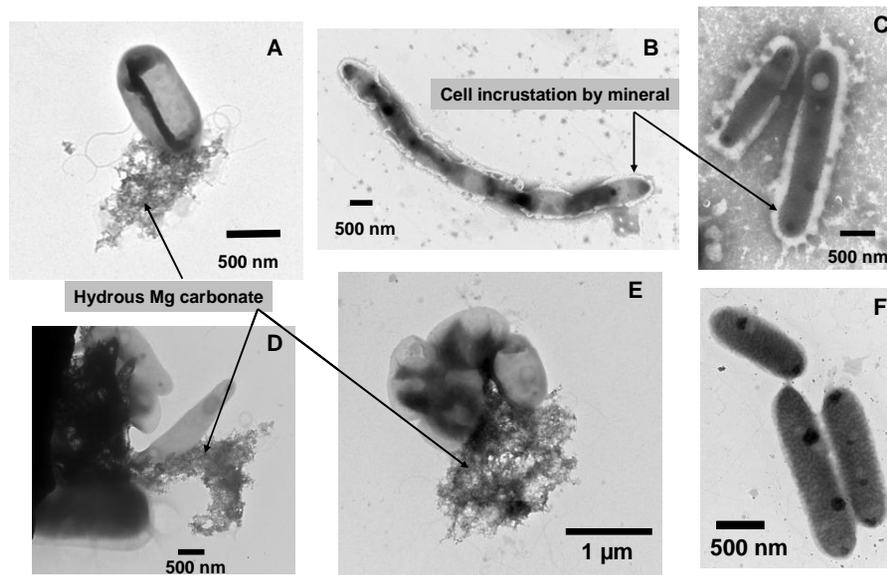


Fig. 3. TEM images of hydrous precipitated Mg carbonate after 14 days in 0.025 M MgCl_2 + 0.05 M NaHCO_3 stirred solution without air bubbling (**A–E**) and *Synechococcus* sp. after its growth during 14 days in BG-11 media without mineral precipitation (**F**).

Can Mg isotopes be used to trace cyanobacteria-mediated

L. S. Shirokova et al.

Title Page

Abstract Introduction

Conclusions References

Tables Figures

⏪ ⏩

◀ ▶

Back Close

Full Screen / Esc

Printer-friendly Version

Interactive Discussion

Can Mg isotopes be used to trace cyanobacteria-mediated

L. S. Shirokova et al.

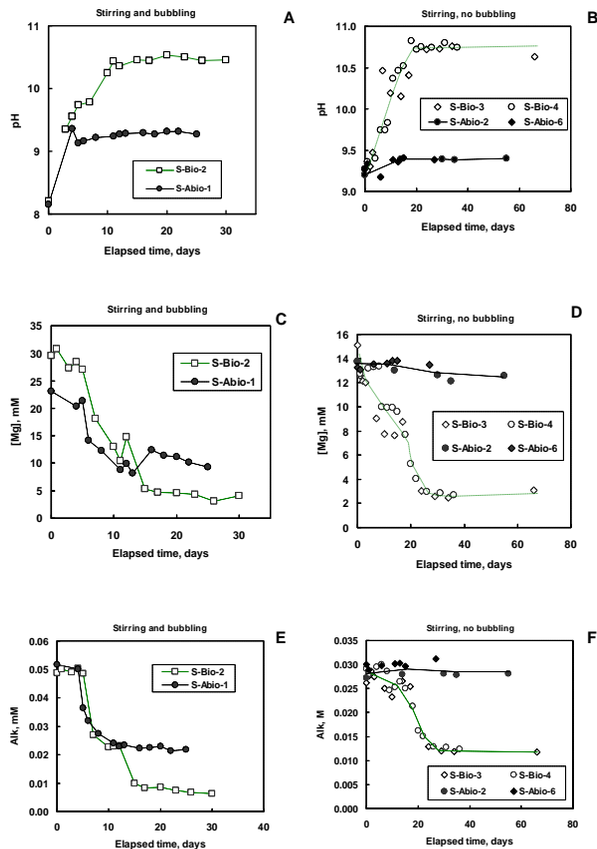


Fig. 4. Reactive fluid evolution during all experiments performed in this study. Temporal evolution of pH (**A**, **B**), Mg concentration (**C**, **D**), and alkalinity (**E**, **F**) during experiments performed in continuously stirred reactors with air bubbling (left panel, **A**, **C**, **E**) and without air bubbling (right panel, **B**, **D**, **F**). The symbol size is encompasses the uncertainty of the analyses. The lines connecting the data points are for the aid of the reader.

Title Page

Abstract

Introduction

Conclusions

References

Tables

Figures

◀

▶

◀

▶

Back

Close

Full Screen / Esc

Printer-friendly Version

Interactive Discussion

Can Mg isotopes be used to trace cyanobacteria-mediated

L. S. Shirokova et al.

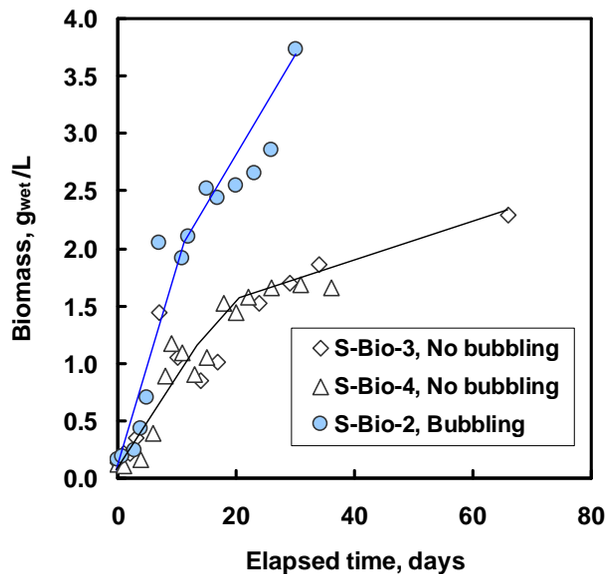


Fig. 5. The biomass evolution during experiments performed in stirred reactors with air bubbling (experiment S-Bio-2, circles) and without air bubbling (experiment S-Bio-3, and S-Bio-4, diamonds and triangles). The lines connecting the data points are for the aid of the reader.

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

Can Mg isotopes be used to trace cyanobacteria-mediated

L. S. Shirokova et al.

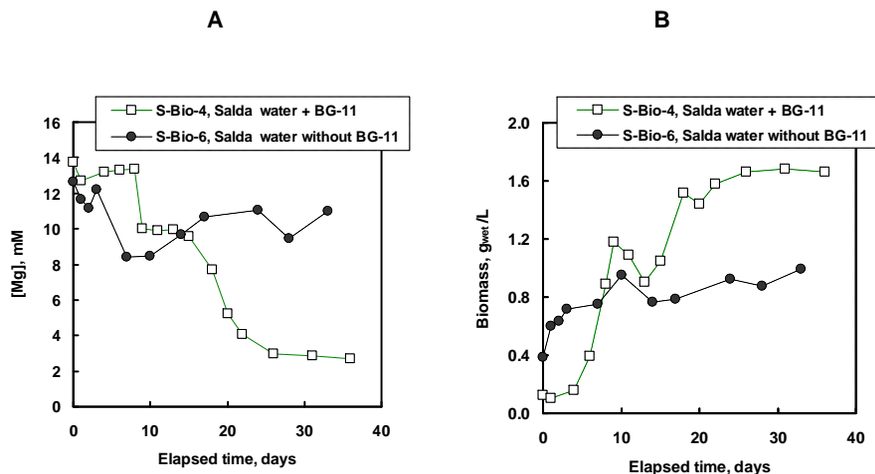


Fig. 6. Magnesium and biomass concentration evolution during experiments in stirred reactors with Salda lake water amended with BG-11 (S-Bio-4, open squares) and without BG-11 components (S-Bio-6, solid circles). The lines connecting the data points are for the aid of the reader.

Title Page

Abstract Introduction

Conclusions References

Tables Figures

⏪ ⏩

◀ ▶

Back Close

Full Screen / Esc

Printer-friendly Version

Interactive Discussion



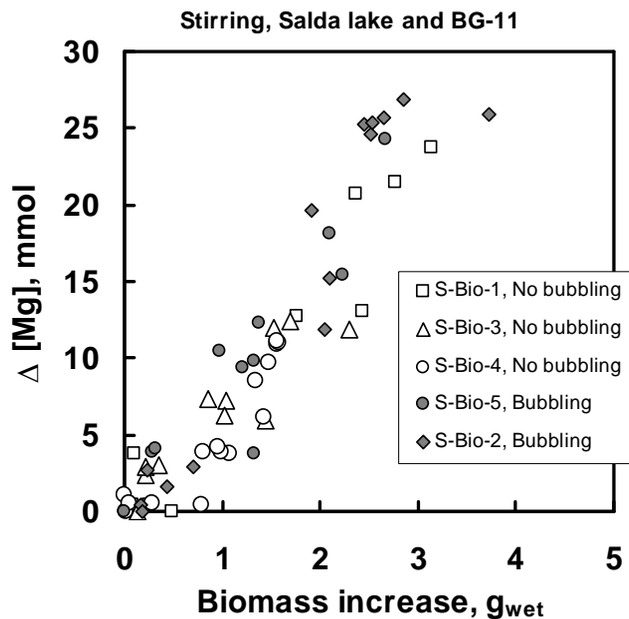
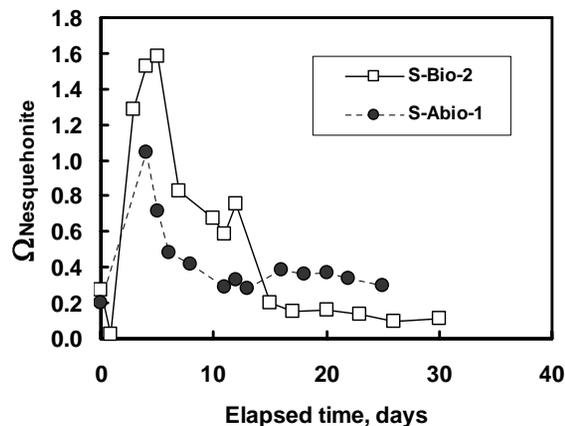


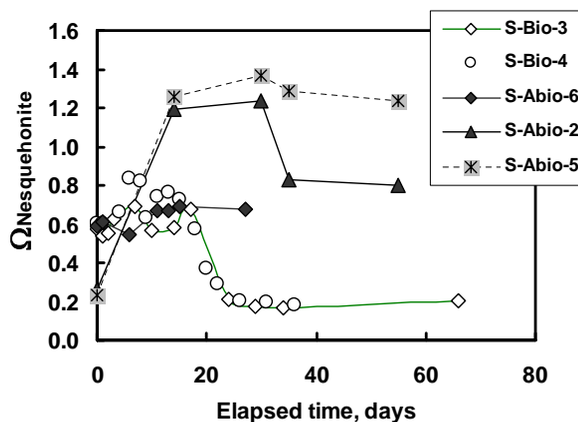
Fig. 7. Relationship between the amount of precipitated magnesium (mmol) and the increase of the biomass (g_{wet}) in different biotic experiments ($r^2 = 0.84 - 0.93$).

Can Mg isotopes be used to trace cyanobacteria-mediated

L. S. Shirokova et al.



(a)



(b)

Fig. 8. Temporal evolution of degree of saturation of nesquehonite ($\Omega_{\text{nesquehonite}}$) of the reactive fluids in stirred experiments. **A:** During biotic experiment S-Bio-2 and abiotic experiment S-Abio-1 with air bubbling; **B:** During biotic and abiotic experiments without air bubbling. The lines connecting the data points are for the aid of the reader.

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

Can Mg isotopes be used to trace cyanobacteria-mediated

L. S. Shirokova et al.

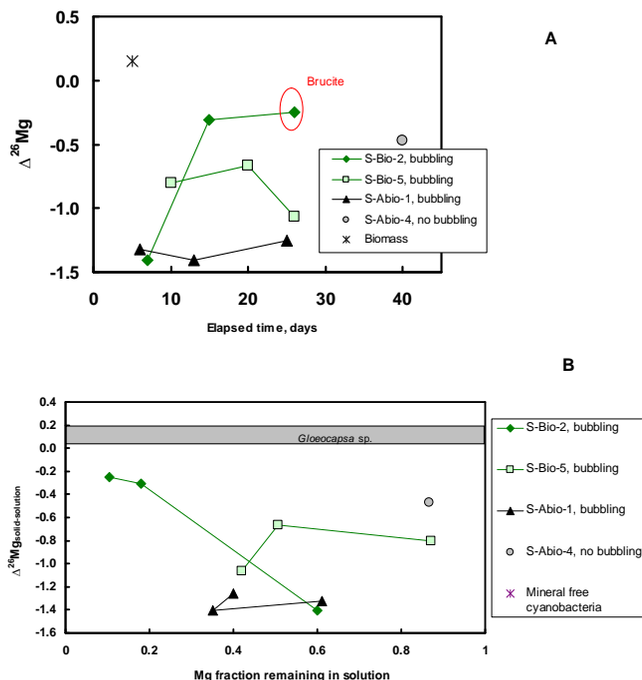


Fig. 9. Temporal evolution of $\delta^{26}\text{Mg}$ in the liquid and solid samples collected during experiments Bio-A and Abio-A plotted as a function of elapsed time (**A**) and fraction of Mg remaining in solution (**B**). The size of the symbols incorporates the uncertainty (2σ) on $\delta^{26}\text{Mg}$. Lines connecting data points (criterion is elapsed time) are for the aid of the viewer.

Title Page

Abstract

Introduction

Conclusions

References

Tables

Figures

⏪

⏩

◀

▶

Back

Close

Full Screen / Esc

Printer-friendly Version

Interactive Discussion

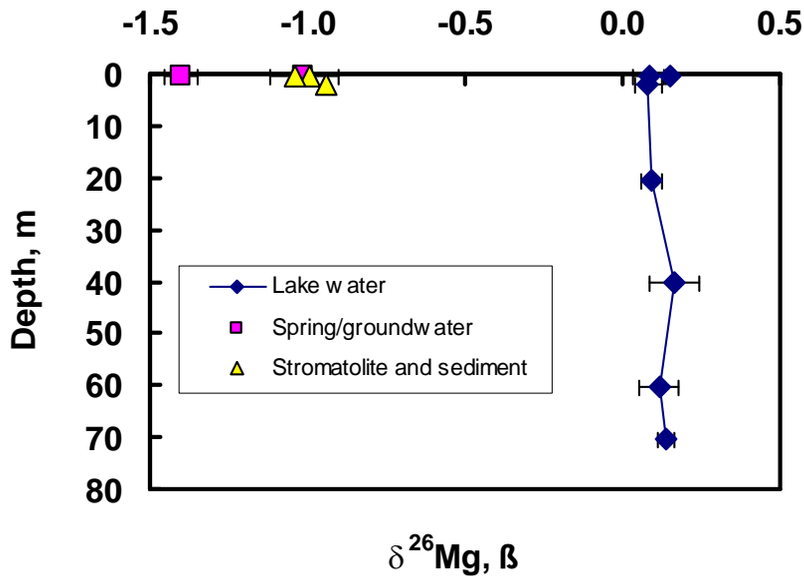


Fig. 10. Mg isotopic composition in Salda lake water (blue diamonds), inflowing springs (pink squares), stromatolites and sediments (yellow triangles).

Can Mg isotopes be used to trace cyanobacteria-mediated

L. S. Shirokova et al.

Title Page

Abstract Introduction

Conclusions References

Tables Figures

⏪ ⏩

◀ ▶

Back Close

Full Screen / Esc

Printer-friendly Version

Interactive Discussion

