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HAL Id: hal-01138257
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Submitted on 7 Dec 2015

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Processes controlling the Si-isotopic composition in the Southern Ocean and application for paleoceanography

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Received: 8 September 2011 – Published in Biogeosciences Discuss.: 17 October 2011
Received: 25 April 2012 – Accepted: 16 May 2012 – Published: 6 July 2012

Abstract. Southern Ocean biogeochemical processes have an impact on global marine primary production and global elemental cycling, e.g. by likely controlling glacial-interglacial pCO2 variation. In this context, the natural silicon isotopic composition (δ30Si) of sedimentary biogenic silica has been used to reconstruct past Si-consumption:supply ratios in the surface waters. We present a new dataset in the Southern Ocean from a IPY-GEOTrACES transect (Bonus-GoodHope) which includes for the first time summer δ30Si signatures of suspended biogenic silica (i) for the whole water column at three stations and (ii) in the mixed layer at seven stations from the subtropical zone up to the Weddell Gyre. In general, the isotopic composition of biogenic opal exported to depth was comparable to the opal leaving the mixed layer and did not seem to be affected by any diagenetic processes during settling, even if an effect of biogenic silica dissolution cannot be ruled out in the northern part of the Weddell Gyre. In general, the isotopic composition of biogenic opal exported to depth was comparable to the opal leaving the mixed layer and did not seem to be affected by any diagenetic processes during settling, even if an effect of biogenic silica dissolution cannot be ruled out in the northern part of the Weddell Gyre. We develop a mechanistic understanding of the processes involved in the modern Si-isotopic balance, by implementing a mixed layer model. We observe that the accumulated biogenic silica (sensu Rayleigh distillation) should satisfactorily describe the δ30Si composition of biogenic silica exported out of the mixed layer, within the limit of the current analytical precision on the δ30Si. The failures of previous models (Rayleigh and steady state) become apparent especially at the end of the productive period in the mixed layer, when biogenic silica production and export are low. This results from (1) a higher biogenic silica dissolution:production ratio imposing a lower net fractionation factor and (2) a higher Si-supply:Si-uptake ratio supplying light Si-isotopes into the mixed layer. The latter effect is especially expressed when the summer mixed layer becomes strongly Si-depleted, together with a large vertical silicic acid gradient, e.g. in the Polar Front Zone and at the Polar Front.

1 Introduction

In the Southern Ocean, the deep nutrient-rich waters ascend into the surface layer south of the Polar Front (PF), and are returned to the subsurface northward before the available pools of nitrogen and phosphorus are fully used by phytoplankton. In contrast, silicic acid (Si(OH)4) becomes strongly depleted by diatom growth (Sarmiento et al., 2004). The balance between the efficiency of the biological pump (e.g. its ability to strip nutrients out of the mixed layer) and deep ocean ventilation releasing CO2 are both likely to control the atmospheric CO2 concentrations on glacial/interglacial timescales (Anderson et al., 2002, 2009; Brzezinski et al., 2002; Sigman et al., 2010).

The δ30Si signatures of biogenic silica (=30R_{Si}O_{2}−30R_{standard}−1, reported in permil units, ‰, where 30R = 30Si:28Si) are a proxy of Si-nutrient utilization in paleoceanography, and have been used to formulate and
test hypotheses on both Southern Ocean productivity and fluctuations in atmospheric $pCO_2$ over glacial cycles (De La Rocha et al., 1998; Anderson et al., 2002; Brzezinski et al., 2002; De La Rocha, 2006; Beucher et al., 2007). During silicic acid consumption by diatoms, the lighter Si isotope ($^{28}$Si) is preferentially consumed, leaving the silicic acid pool enriched in heavy Si-isotope ($^{30}$Si) (De La Rocha et al., 1997). Such preferential incorporation of $^{28}$Si into biogenic silica (bSiO$_2$) is defined by a fractionation factor ($^{30}$ε), which is equivalent to the ratio of the reaction rates of the heavy ($^{30}$k) and light ($^{28}$k) Si-isotopes ($=^{30}$k/$^{28}$k − 1, reported in permil units, ‰). Therefore, for a given set of conditions setting the supply of Si(OH)$_4$, a relationship is expected between the decrease of Si(OH)$_4$ concentration due to the Si-uptake and the increase of $^{30}$Si of both bSiO$_2$ and Si(OH)$_4$ (De La Rocha et al., 1997). However, the calibration of this proxy in the modern ocean has still not been fully achieved and processes such as Si(OH)$_4$ supply and bSiO$_2$ dissolution can bias the expected relationship between Si(OH)$_4$ concentration and $^{30}$Si composition (Demarest et al., 2009). For example, contemporary isotopic compositions of biogenic silica vary widely in the Southern Ocean (−0.5 to +2.5 ‰), while the isotopic range of silicic acid is smaller (from +1.5 to 3.4 ‰) (Varela et al., 2004; Cardinal et al., 2005, 2007; Fripiat et al., 2011a; Cavagna et al., 2011). None of the existing isotopic fractionation models are capable of reproducing these differences. This highlights the significant importance of fully understanding the different processes involved in contemporary Si-isotopic balances before applying this proxy to reconstruct past ocean environments.

New data from the summer Antarctic Circumpolar Current (ACC) are presented here, looking at the relationship between $^{30}$Si$_{Si(OH)}$ and $^{30}$Si$_{SiO_2}$ in the mixed layer. Primary objectives are to (1) assess the processes controlling mixed layer $^{30}$Si$_{SiO_2}$ and across the water column, and (2) the impact of such processes on the relationship between the relative silicic acid utilisation and $^{30}$Si.

2 Materials and methods

2.1 Sampling and hydrology

Between 8 February and 8 March 2008, the GEOTRACES-International Polar Year (IPY) BONUS-GoodHope (BGH) cruise aboard R/V Marion Dufresne covered a transect from Cape Basin to the north of the Weddell Gyre (up to 58° S) centred around the 0° meridian (Fig. 1). Biogenic silica was sampled (1) in the mixed layer at 7 stations crossing the different hydrographic regimes of the ACC, and (2) from a profile of the surface to the deep ocean at 3 stations. For these stations and others from the same cruise, the water column profiles of silicic acid $^{30}$Si are discussed in an associated paper (Fripiat et al., 2011b). In the latter study, water masses dynamics is suggested to be the main driving processes setting the distribution of the whole water-column $^{30}$Si$_{Si(OH)}$.

Seawater was collected using a CTD Rosette equipped with 121 Niskin bottles. Water samples (∼0.25 to ∼10 l) were immediately filtered on 0.4 µm Nuclepore polycarbonate membranes, using Perspex filtration units under the pressure of filtered air (0.4 µm Nuclepore; <2 bar). Filtered water samples for silicic acid analyses were stored in acid-cleaned polypropylene (PP) bottles at room temperature in the dark. Nuclepore membranes were stored in polycarbonate Petri dishes and dried overnight at 50°C. Large Volume Filtration Systems, hereafter referred to as “in situ pumps” (Challenger Oceanic Systems and Services, Surry, UK), sampled the complete water column by filtering 15 to 2501 of water through hydrophylc polyethersulfone membranes (SUPOR, 293 mm Ø, 0.45 µm). After partitioning the filters amongst the different end-users knowing the used filter surface, the pieces of SUPOR membranes were dried overnight at 50°C and stored in Petri dishes in the dark at ambient temperature.

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www.biogeosciences.net/9/2443/2012/
2.2 Analyses

Samples were processed ashore (RMCA, Tervuren). For bSiO_2 the membranes were subjected to a wet-alkaline digestion (adapted from Ragueneau et al., 2005) whereby bSiO_2 is dissolved with a 0.2 µmol l⁻¹ NaOH solution (pH 13.3) at 100 °C for 40 min. As this digestion can also dissolve some lithogenic silica, we also analyzed aluminum (Al) in the digested solution to check for possible lithogenic contamination. Aluminum was determined using a Sector Field Inductively Coupled Plasma Mass Spectrometer (SF-ICP-MS Element2, detection limit = 3 ppb, = 3*blank standard deviation). Using a Si:Al mass ratio of 3.74 for average crust (Taylor and McLennan, 1985), the lithogenic silica contributed to less than 2.5 % of the Si in digested samples. The average distribution of clays in marine sediments along the BONUS-GoodHope transect is <10 % kaolinite, >40 % illite, <20 % smectite and 20–30 % chlorite (Füttner, 2006). The δ³⁰Si signatures for such different types of clays published in previous studies are the following: kaolinite −2.2 ± 0.4 ‰ (Ziegler et al., 2005; Opfergelt et al., 2010); illite: −0.9 ± 0.2 ‰ (Douthitt, 1982) smectite: −0.3 ± 0.2 ‰ (Georg et al., 2009). No δ³⁰Si value is available for chlorite. Setting the chlorite δ³⁰Si values to the one of kaolinite which maximizes the potential clay δ³⁰Si contamination, we estimate an average clay δ³⁰Si value for the BONUS-GoodHope transect of −1.3 to −1.0 ‰ using mass and isotopic balances from the expected distribution of clays. Taking the heavier BGH δ³⁰Si_bSiO² and the calculated clay δ³⁰Si, with a lithogenic silica contamination of 2.5 %, the maximum potential contribution of such lithogenic Si to the measured δ³⁰Si is 0.1 %. Such a lithogenic contribution can be neglected since it is similar to the achieved analytical precision for δ³⁰Si determination (Abraham et al., 2008).

Si(OH)₄ and bSiO₂ concentrations were determined via a colorimetric method (Grasshoff et al., 1983) on the same samples as for Si-isotopic composition. Since Si(OH)₄ concentrations of surface and some subsurface samples at stations north of the Polar Front (PF) were too low (<10 µmol l⁻¹) to directly apply the Si purification procedure required for Si-isotopic measurement (De La Rocha et al., 1996), a Si(OH)₄ preconcentration step was performed. It was achieved using a protocol adapted by Brzezinski et al. (2003) and Reynolds et al. (2006) from the MAGIC method (Karl and Tien, 1992). Briefly, the method consists of one or two steps (depending of the Si-recovery of the first step) of the co-precipitation of Si(OH)₄ with brucite (Mg(OH)₂), by increasing pH either with NaOH 1 µmol l⁻¹ (20 ml l⁻¹ seawater, Reynolds et al., 2006) or NH₄OH 13.5 µmol l⁻¹ (6 ml l⁻¹ seawater Brzezinski et al., 2003). The precipitates were recovered by centrifugation and then redissolved with HCl.

Silicon was co-precipitated with triethylamine molybdate (De La Rocha et al., 1996) with a minimum Si requirement of ~1.5 µmol Si. After combustion of the siliconolybdate precipitate in covered Pt crucibles at 1000 °C, the pure cristobalite phase was transferred to pre-cleaned polypropylene vials. Dissolution of cristobalite was done in a dilute HF/HCl mixture as described in Cardinal et al. (2003). Isotopic measurements were carried out on a Nu Plasma MC-ICP-MS (ULB, Brussels) using Mg external doping in dry plasma mode following Abraham et al. (2008). The average precision and reproducibility of the measurements are ±0.1 ‰ (±1 sd) for δ³⁰Si (Reynolds et al., 2007). The accuracy of the measurements is checked on a daily basis on secondary reference materials (e.g. Diatomite) with known Si isotopic compositions resulting from an inter-comparison exercise (Reynolds et al., 2007).

3 Results

Surface δ³⁰Si_{Si(OH)₄} and [Si(OH)₄] display an opposite pattern with latitude (Table 1 and Fig. 2a). There is a 30Si enrichment associated with the northward decreasing Si(OH)₄ gradient across the Polar Front (PF), as observed earlier for δ³⁰Si_{Si(OH)₄} (Varela et al., 2004; Cardinal et al., 2005) and Si(OH)₄ concentrations (Brzezinski et al., 2001; Quégneux and Brzezinski, 2002). δ³⁰Si_{SiO₂} values are systematically lighter than δ³⁰Si_{Si(OH)₄} values in agreement with the preferential uptake of 28Si by diatoms. From the WG to the PFZ we observe a trend of increasing δ³⁰Si_{SiO₂}, following the δ³⁰Si_{Si(OH)₄} gradient, although the trend is much steeper for the former, as also observed by Varela et al. (2004) and Cardinal et al. (2007). North of the PFZ and SAZ, respectively, the δ³⁰Si_{SiO₂} and δ³⁰Si_{Si(OH)₄} trend reverses, again displaying a steeper gradient for δ³⁰Si_{SiO₂} values. This decoupling is reflected in the apparent fractionation factor (Δ³⁰Si = δ³⁰Si_{Si(OH)₄} − δ³⁰Si_{SiO₂}) which varies widely with latitude (Fig. 2b). Lower Δ³⁰Si values (∼0.35 ± 0.10 ‰) are observed at the PF and in the PFZ. Such low values seem to be a recurrent feature in the Southern Antarctic Circumpolar Current at low Si(OH)₄ concentrations (Fig. 3). The mixed layer BONUS-GoodHope results for δ³⁰Si_{Si(OH)₄} (+1.8 to +3.2 ‰) and for δ³⁰Si_{SiO₂} (+0.3 to +2.5 ‰) fall in the range reported for previous Southern Ocean studies (+1.5 to +3.1 ‰; for δ³⁰Si_{Si(OH)₄}; −0.5 to +2.6 ‰ for δ³⁰Si_{SiO₂}) (Varela et al., 2004; Cardinal et al., 2005, 2007; Fripiat et al., 2011a; Cavagna et al., 2011). For bSiO₂, the previous studies are restricted to the Seasonal Ice Zone (SIZ), AZ, and PFZ. It is interesting to note that the range of δ³⁰Si_{SiO₂} values are roughly two times those of δ³⁰Si_{Si(OH)₄} (Fig. 3a). A clear trend between Δ³⁰Si and silicic acid concentration is observed in the mixed layer (Fig. 3b, non-linear regression using a power function, Δ³⁰Si = 0.404°[Si(OH)₄]°³⁶, R² = 0.61, p-value < 0.01), similar to the one obtained with sponge spicules (Wille et al., 2010; Hendry and Robinson, 2012).

The water column profiles of Si(OH)₄ and δ³⁰Si_{Si(OH)₄} clearly display 30Si enrichment and Si(OH)₄ depletion...
Table 1. Si(OH)$_4$ and bSiO$_2$ concentrations and isotopic compositions. Only the standard deviations for duplicates are shown. Bold numbers represents samples from in situ pumps. The mixed layer depth (MLD) has been determined using a combination of threshold density/temperature criterions relative to 10 m ($\Delta T$ °C = 0.2 °C; $\Delta \sigma$ = 0.03 and 0.125 kg m$^{-3}$; De Boyer Montégut et al., 2004). $\delta^{30}$Si-Si(OH)$_4$ data are from Fripiat et al. (2011b).

<table>
<thead>
<tr>
<th>Station</th>
<th>Depth m</th>
<th>Si(OH)$_4$ µmol l$^{-1}$</th>
<th>$\delta^{30}$Si-Si(OH)$_4$ ‰ ± sd</th>
<th>Depth m</th>
<th>bSiO$_2$ µmol l$^{-1}$</th>
<th>$\delta^{30}$Si-bSiO$_2$ ‰ ± sd</th>
</tr>
</thead>
<tbody>
<tr>
<td>Super 5</td>
<td>29</td>
<td>65.2</td>
<td>1.85 ± 0.03</td>
<td>30</td>
<td>0.36</td>
<td>0.28 ± 0.04</td>
</tr>
<tr>
<td>16 March 2008</td>
<td>88</td>
<td>64.2</td>
<td>2.00 ± 0.03</td>
<td>88</td>
<td>1.23</td>
<td>0.28</td>
</tr>
<tr>
<td>00.02° E–57.32° S</td>
<td>151</td>
<td>84.1</td>
<td>1.50 ± 0.03</td>
<td>140</td>
<td>0.12</td>
<td>$-0.40 ± 0.10$</td>
</tr>
<tr>
<td>MLD = 58–115 m</td>
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<td>96.2</td>
<td>1.35 ± 0.01</td>
<td>250</td>
<td>0.23</td>
<td>0.16 ± 0.01</td>
</tr>
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<td>251</td>
<td>101.5</td>
<td>1.37 ± 0.05</td>
<td>400</td>
<td>0.18</td>
<td>0.27 ± 0.03</td>
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<td>299</td>
<td>104.2</td>
<td>1.41 ± 0.05</td>
<td>700</td>
<td>0.10</td>
<td>0.26 ± 0.02</td>
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<tr>
<td></td>
<td>400</td>
<td>110.5</td>
<td>1.44 ± 0.36</td>
<td>1800</td>
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<td>0.68 ± 0.11</td>
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<tr>
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<td>499</td>
<td>117.2</td>
<td>1.47</td>
<td>2500</td>
<td>0.04</td>
<td>0.65 ± 0.15</td>
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<tr>
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<td>700</td>
<td>116.5</td>
<td>1.44 ± 0.20</td>
<td>3200</td>
<td>0.04</td>
<td>0.72 ± 0.12</td>
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<td>1.95</td>
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<td>n.a.</td>
<td>n.a.</td>
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<td>0.72</td>
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<td>11</td>
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<td>1.38</td>
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<td>n.a.</td>
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<td>1.37 ± 0.10</td>
</tr>
<tr>
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<td>22.3</td>
<td>2.37 ± 0.06</td>
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<td>0.29</td>
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<td>n.a.</td>
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<td>170</td>
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<td>1.48 ± 0.00</td>
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<td>1.36 ± 0.06</td>
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<td>2488</td>
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<td>1.42 ± 0.20</td>
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<td>1.47</td>
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<td>5.1</td>
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<td>3.24 ± 0.01</td>
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<td>2.85</td>
<td>41</td>
<td>0.32</td>
<td>2.50 ± 0.06*</td>
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<td>2.36</td>
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<td>2.57 ± 0.19</td>
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<td>9.2</td>
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<td>600</td>
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<td></td>
<td>401</td>
<td>25.3</td>
<td>1.93 ± 0.35</td>
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<td>MLD = 13–19 m</td>
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* Samples in the mixed layer have been merged for Si-isotopic analysis.
towards the surface (Table 1; see Fripiat et al., 2011b for a more complete discussion on water-column profiles of $\delta^{30}$Si$_{Si(OH)_4}$), similar to previous studies (e.g. Cardinal et al., 2005). While biogenic silica concentrations exhibit an inverse pattern (Fig. 4a, c), $\delta^{30}$Si$_{bSiO_2}$ remains constant with depth in the AZ and in the upper $\sim$1000 m of the northern WG and the PFZ (Fig. 4b, d). In the WG, $\delta^{30}$Si$_{bSiO_2}$ decreases just below the mixed layer, coinciding with a decrease in bSiO$_2$ and $\delta^{30}$Si$_{Si(OH)_4}$, while below 1000 m $\delta^{30}$Si$_{bSiO_2}$ increases.

4 Discussion

Proxies can be influenced by multiple factors, and the sensitivity of a specific proxy to these factors likely changes in space and time. In the following sections, we discuss the processes controlling the origin of the isotopic composition of biogenic silica in the mixed layer (Sect. 4.1) and subsequently the fate of the isotopic composition of biogenic silica across the water column (Sect. 4.2). The link between the isotopic composition and the relative silicic acid utilization in the surface waters will be investigated.

4.1 Origin of the mixed layer biogenic silica isotopic composition

Two models are commonly applied to describe Si-isotopic fractionation during Si-consumption by diatoms, directly linking the $\delta^{30}$Si to the relative utilisation of Si(OH)$_4$. If a pool of silicic acid is consumed without resupply during the consumption process and with a constant fractionation factor ($^{30}$e), then the isotopic evolution of the remaining Si(OH)$_4$, instantaneous bSiO$_2$ (hereafter referred to with the superscript “inst”), and accumulated bSiO$_2$ (hereafter referred to
with the superscript “acc”) should follow the Rayleigh fractionation equations (De La Rocha et al., 1997):

\[
\delta^{30}\text{Si}_{\text{Si(OH)4}}[t] = \delta^{30}\text{Si}_{\text{Si(OH)4}}[t_0] + 30\varepsilon \cdot \ln(f)
\]  

(1)

\[
\delta^{30}\text{Si}_{\text{bSiO}_2[\text{inst}]\,[t]} = \delta^{30}\text{Si}_{\text{Si(OH)4}}[t_0] + 30\varepsilon
\]  

(2)

\[
\delta^{30}\text{Si}_{\text{bSiO}_2[\text{acc}]\,[t]} = \delta^{30}\text{Si}_{\text{Si(OH)4}}[t_0] - 30\varepsilon \cdot f \cdot \ln(f) \left( \frac{1}{1-f} \right)
\]  

(3)

where \( f \) is the \([\text{Si(OH)}_4]\,[t]/[\text{Si(OH)}_4]\,[t_0]\) ratio. The subscript “[t]” is the sampling time and “[t0]” the initial time. Note that in such a system, the apparent fractionation factor (\(\Delta^{30}\text{Si} = \delta^{30}\text{Si}_{\text{Si(OH)4}} - \delta^{30}\text{Si}_{\text{bSiO}_2}\)) is either constant (Eq. (2) for instantaneous \(\text{bSiO}_2\)) or increases (Eq. (3) for accumulated \(\text{bSiO}_2\)) with silicic acid consumption.

The alternative to this closed system is the open, flow-through system, where the substrate is continuously supplied where the reaction continuously occurs at the same rate (also referred to as steady state system, Fry, 2006).

\[
\delta^{30}\text{Si}_{\text{Si(OH)4}}[t] = \delta^{30}\text{Si}_{\text{Si(OH)4}}[t_0] - 30\varepsilon \cdot (1 - f)
\]  

(4)

\[
\delta^{30}\text{Si}_{\text{bSiO}_2[\text{inst}]\,[t]} = \delta^{30}\text{Si}_{\text{Si(OH)4}}[t_0] + 30\varepsilon \cdot f = \delta^{30}\text{Si}_{\text{Si(OH)4}}[t_0] + 30\varepsilon
\]  

(5)

The steady state system assumes equilibrium and does not represent a temporal evolution. Conceptually, for the Southern Ocean this poses a problem, since Si-depletion is observed in the mixed layer as a result of varying growth and export of diatoms (Brzezinski et al., 2001). Note that in such a system, \(\Delta^{30}\text{Si}\) is constant (Eq. 5) along with silicic acid consumption.

Neither of these models can explain the large variability that is observed in \(\delta^{30}\text{Si}_{\text{bSiO}_2}\) values compared to \(\delta^{30}\text{Si}_{\text{Si(OH)4}}\) (Fig. 3a), and therefore the associated low and high \(\Delta^{30}\text{Si}\) (Fig. 3b). These model failures may result from the non-validity of one of the underlying model assumptions. We will investigate the impact of the following assumptions: (1) the fractionation factor is constant all along the seasonal Si(OH)4 depletion (Sect. 4.1.1), and (2) the seasonal expression of the different isotopic effects is under-resolved in the model equations (Sects. 4.1.2 and 4.1.3).
4.1.1 Variability in the fractionation factor

The mean apparent fractionation factor ($\Delta^{30}\text{Si} = \delta^{30}\text{Si}_{\text{Si(OH)}_4} - \delta^{30}\text{Si}_{\text{bSiO}_2}$) south of the Subantarctic Front is $1.3 \pm 0.5\%$ ($n = 90$; Fig. 3b). Assuming that there is no accumulation of biogenic silica in the mixed layer, $\Delta^{30}\text{Si}$ should be equal, but with an opposite sign, to the fractionation factor associated with biogenic silica production ($^{30}\varepsilon$; Eqs. 2 and 5). The latter is not significantly different from the in vitro $^{30}\varepsilon$ estimation with temperate diatom species ($-1.1 \pm 0.4\%$; De La Rocha et al., 1997), the in situ $^{30}\varepsilon$ compilation in the ACC ($-1.2 \pm 0.2\%$; Fripiat et al., 2011a), or the different $^{30}\varepsilon$ estimations reported in other oceanic regions (De La Rocha et al., 2000; Reynolds et al., 2006; Beucher et al., 2008). Despite that these different estimations are roughly centered on the in vitro estimations from De La Rocha et al. (1997; $-1.1 \pm 0.4\%$), the large dispersion (from $-0.4$ to $-2.0\%$) suggests that the constancy of the fractionation factor can be discussed.

In the Southern Ocean, a clear trend between $\Delta^{30}\text{Si}$ and silicic acid concentration is observed in the mixed layer (Fig. 3b). If indeed $\Delta^{30}\text{Si}$ can be taken as an estimate of $-^{30}\varepsilon$, then such dataset can result from a causal link between silicic acid concentration and the extent of Si-isotopic fractionation. However, the physiological mechanisms leading to the fractionation of Si-isotopes by diatoms are still poorly assessed and understood. Several mechanisms can induce a variation in $^{30}\varepsilon$: (1) an inter-specific variation, and an impact of (2) the Si-cellular metabolic balance and (3) the bSiO$_2$ dissolution. In the following, these three points will be discussed:

1. Recently, Sutton et al. (2011) suggested an inter-specific variation in Southern Ocean diatoms based on in vitro incubations. Such observations are not in agreement with De La Rocha et al. (1997)'s hypothesis showing no inter-specific variation with temperate diatom species. Despite of being very hypothetical, different diatom species adapted to different silicic acid availability, together with different fractionation factors, could explain the observed relationship between $\Delta^{30}\text{Si}$ and silicic acid concentration (Fig. 3b).

2. If the main Si-isotopic fractionation is occurring within the diatom cell (e.g. during the polymerization of biogenic silica), the isotopic fractionation will be more expressed when the efflux/influx ratio increases (Milligan et al., 2004), as expressed in the following equation:

$$^{30}\varepsilon_{\text{net}} = \frac{^{30}\varepsilon_{\text{inf}}}{\rho_{\text{eff}}} + \rho_{\text{eff}} \left( {^{30}\varepsilon_{\text{poly}} - ^{30}\varepsilon_{\text{eff}}} \right)$$

where the subscript “Inf” (=influx) is the transport rate across the plasmalemma from outside to inside the cell; “poly” (=polymerisation) the rate of frustule formation inside the silicon vesicle deposit (SDV); “Eff” (efflux) the transport rate across the plasmalemma from inside to outside the cell.

The efflux has never been measured in situ, precluding any comparison with oceanic data. It has been suggested to be a consequence of transient imbalances between uptake and polymerization caused by a level of transport that exceeds the capacity of intracellular silicon-binding component (Martin-Jézéquel et al., 2000; Thamatrakoln and Hildebrand, 2008). Average concentration of Si(OH)$_4$ in most of the ocean’s euphotic zone is less than 10 µmol Si l$^{-1}$ (Garcia et al., 2005). Si(OH)$_4$ uptake by field assemblages would mostly be directly controlled by silicic acid transporters (SITs), which would minimize the associated costs of excess uptake and efflux (Thamatrakoln and Hildebrand, 2008). On the other side, our data are all from the Southern Ocean where high Si(OH)$_4$ concentrations prevail. These concentrations are expected to be in a range favoring diffusion mediated uptake, incidentally increasing the probability of efflux to occur (Thamatrakoln and Hildebrand, 2008).

If (i) the isotopic fractionation associated with the polymerization is larger than the ones occurring during both influx and efflux, and (ii) the efflux/influx ratio is proportional to the silicic acid concentration, then the isotopic fractionation associated with biogenic silica production should increase together with silicic acid concentration (Eq. 6), as observed in Fig. 3b.

It seems unlikely that (1) and (2) are sufficient to fully explain the observed trend in Fig. 3b since the $\Delta^{30}\text{Si}$ Front is approaching zero at low concentrations. Indeed, it would suggest that either some diatom species or diatoms growing in environmental conditions unfavorable to efflux, specifically low Si(OH)$_4$ concentration (Thamatrakoln and Hildebrand, 2008), do not fractionate the silicon isotopes. For (2), it implies that the fractionation associated with influx is 0‰ (Eq. 6). This is in disagreement with the observations in other oceanic regions or with in vitro incubations: (i) $\Delta^{30}\text{Si}$ north of the Subantarctic Front is larger than 1‰ despite low silicic acid concentrations, (ii) an imprint from isotopic fractionation is clearly observed on the surface $\delta^{30}\text{Si}_{\text{Si(OH)}_4}$ distribution in every oceanic areas measured to date, including the ones where low silicic acid concentration prevails all year (De La Rocha et al., 2000; Cardinal et al., 2005; Reynolds et al., 2006; Beucher et al., 2008; de Souza, 2011; this study), and (iii) Milligan et al. (2004) with in vitro incubations measured an isotopic fractionation ($^{30}\varepsilon \approx -1.6\%$) which seemed not to be affected by the efflux/influx ratio and requiring a significant isotopic fractionation during influx.

3. Demarest et al. (2009) report that during dissolution of biogenic silica, light Si-isotopes are released preferentially with a fractionation factor ($^{30}\varepsilon_{\text{diss}}$) estimated to be at $-0.55 \pm 0.05\%$. This effect partly counteracts the
fractionation occurring during bSiO$_2$ production. By increasing the mixed layer $D:P$ ratio, the net fractionation ($^{30}\epsilon_{\text{net}}$) factor associated with net bSiO$_2$ production ($\approx \text{net} P$) would be dampened as follows (Demarest et al., 2009):

$$^{30}\epsilon_{\text{net}} = {^{30}\epsilon_{\text{upt}}} + \frac{D}{P} \cdot {^{30}\epsilon_{\text{diss}}}. \quad (7)$$

Brzezinski et al. (2003) observed a shift in the mode of functioning of the surface silicon cycle, from bSiO$_2$ dissolution supporting a small fraction of gross bSiO$_2$ production during diatom bloom to bSiO$_2$ dissolution supporting the majority of gross bSiO$_2$ production during non-bloom periods when silica production is low. Applying strictly the Eq. (7), such isotopic effect could explain at the maximum only 0.55% of the variation observed in Fig. 3b (being $\approx 2\%$).

To conclude this section, the seasonal and spatial variations of the fractionation factor are not sufficient to fully explain the observed trends between $\Delta^{30}\text{Si}$ and silicic acid concentration (Fig. 3b). Additional processes are required. In the following (Sects. 4.1.2 and 4.1.3), the seasonal variation in the expression of the different isotopic effects (dissolution and mixing) will be discussed together with induced disequilibriums between biogenic silica and silicic acid. A box model based on conservation of Si mass and isotopic composition during the productive period for the mixed layer is implemented in Sect. 4.1.2. The different hypotheses are tested in the Sect. 4.1.3.

4.1.2 Model description

Figure 5a shows the model structure with the associated fluxes: Si(OH)$_4$ supply, bSiO$_2$ production, dissolution, and export. The box represents a 100 m deep mixed layer and has two compartments: the Si(OH)$_4$ and bSiO$_2$ pools. South of Africa, Faure et al. (2011) estimated a mean mixed layer depth of $\pm100\text{ m}$ for the productive period across the ACC (including the AZ, PFZ, and SAZ), in agreement with our assumption. The accumulated exported bSiO$_2$ is also simulated, to assess the effects of seasonal changes upon the isotopic composition of exported $\delta^{30}\text{Si}$. Si(OH)$_4$, i.e. the variable measured in paleoceanography. Here, the productive period is assumed to have a duration of 90 days, with the productive peak roughly centered in December, as shown by chl $a$/POC satellite data and modeling studies (Pondaven et al., 1998; Moore and Abbot, 2000; Allison et al., 2010). Across the BGH transect which took place in late summer 2008, mixed layer silicon properties can be considered as representative of final summer conditions. This is supported by Si-uptake in the mixed layer being in the lower range of published values, 0.02 ± 0.03 mmol Si l$^{-1}$ d$^{-1}$ (mean Si-uptake from $^{30}$Si tracer incubations for the BGH transect, $n = 27$ samples, each incubated as triplicates; Fripiat, 2010). This indicates that the end of the diatoms’ growth period has been reached. The PFZ station was chosen as a case study for this simulation as both (1) near complete Si-depletion (Fig. 2a) and (2) low $\Delta^{30}\text{Si}$ are observed (Fig. 2b). The model initial conditions (i.e. before the start of the growth season) are set using the Si-characteristics of subsurface waters in the PFZ by Fripiat et al. (2011b; median for the halocline in the PFZ, see Table 2 for the initial and final conditions), argued here to represent the winter mixed layer (Tomczak and Godfrey, 2001).

The vertical supply of Si to the mixed layer is described either by a constant flux (zero order equation) or by 10 discrete mixing events (every 9 days) supplying an equal quantity of Si into the mixed layer. The gross uptake rate of Si (=gross bSiO$_2$ production) during the growth period is described by a Michaelis-Menten saturation function:

$$V_{\text{Si}}(t) = \frac{V_{\text{max}} \cdot [\text{Si(OH)}_4]}{k_{\text{si}} + [\text{Si(OH)}_4]} \quad (8)$$

where $V_{\text{Si}}$ is the specific uptake rate (d$^{-1}$), $V_{\text{max}}$ the maximum specific uptake rate, and $k_{\text{si}}$ (mmol l$^{-1}$) the Si(OH)$_4$ concentration when $V_{\text{Si}} = V_{\text{max}}/2$. In the ACC silicon uptake is well described using the Michaelis-Menten relationship (Pondaven et al., 1998, 2000). The integrated (mixed layer depth = MLD) gross bSiO$_2$ production ($P$; mmol m$^{-2}$ d$^{-1}$) is described as follows:

$$P = \int_{0}^{\text{MLD}} V_{\text{Si}(t)} \cdot [\text{bSiO}_2\text{ML}(t)] \cdot dz. \quad (9)$$

The “ML” superscript indicates mixed layer biogenic silica (being the initial bSiO$_2$+ net bSiO$_2$ production, at each time step), which is a mixture of detrital and living diatom cells (Krause et al., 2010). The bSiO$_2$ export (= $E$, mmol m$^{-2}$ d$^{-1}$) is described by a first order equation (first order constant $= k_{\text{export}}$):

$$E = \int_{0}^{\text{MLD}} k_{\text{export}} \cdot [\text{bSiO}_2\text{ML}(t)] \cdot dz. \quad (10)$$

The bSiO$_2$ dissolution:production ($D:P$) ratio is assumed to increase linearly (from 0.0 to 1.0 over the expected range Brzezinski et al., 2003; Fig. 5d):

$$\frac{d(D/P)}{dt} = \frac{1}{N_{\text{season}}} \quad (11)$$

where $N$ is the number of days. The model could be over-simplified as in the ACC, $D:P$ ratios exceeding 1 have already been documented at the end of summer (Beucher et al., 2004; Fripiat et al., 2011c). This has been explained by an accumulation of detrital diatom cells in the mixed layer after a productive period.

The fractionation factor associated with net bSiO$_2$ production is determined following equation 7 where $^{30}\epsilon_{\text{upt}}$ has
been compiled for the ACC at $-1.2 \pm 0.2\%$ in Fripiat et al. (2011a) and $30 \varepsilon_{\text{diss}}$ has been measured experimentally at $-0.55 \pm 0.05\%$ by Demarest et al. (2009). Assuming a $D:P$ ratio evolution as described in Eq. (7), the $30 \varepsilon_{\text{net}}$ at the beginning of the productive period will be equal to the $30 \varepsilon_{\text{upt}}$ ($-1.2\%$) and decrease to $-0.65\%$ ($30 \varepsilon_{\text{upt}} - 30 \varepsilon_{\text{diss}}$) at the end, in agreement with Eq. (11). In the following discussion, model results with and without an isotopic fractionation during bSiO$_2$ dissolution will be compared. Given the lack of information on the role Si-efflux on isotopic fractionation, we decided to not include Eq. (6) in this model.

Using the above parameters and assuming the silicic acid pool is homogeneous and is fed with silicic acid of constant $30 \delta_{\text{Si}}$ composition, the seasonal evolution can be described as follows:

$$\frac{d(m_{\text{Si(OH)}_4})}{dt} = F_{\text{supply}} + \left(\frac{D}{P_{[t]}} - 1\right) \cdot P_{[t]} - E_{[t]} = F_{\text{supply}} - \text{net} P_{[t]} \quad (12)$$

$$m_{\text{Si(OH)}_4} \cdot \frac{d(\delta_{30} \text{Si(OH)}_4)}{dt} = F_{\text{supply}} \cdot \delta_{30} \text{Si}_{\text{supply}} - 30 \varepsilon_{[t]} \cdot \text{net} P_{[t]} \quad (13)$$

$$\frac{d(m_{\text{Si(OH)}_4})}{dt} = F_{\text{supply}} + \left(\frac{D}{P_{[t]}} - 1\right) \cdot P_{[t]} - E_{[t]} = F_{\text{supply}} - \text{net} P_{[t]} \quad (14)$$

$$m_{\text{Si(OH)}_4} \cdot \frac{d(\delta_{30} \text{Si(OH)}_4)}{dt} = F_{\text{supply}} \cdot \delta_{30} \text{Si}_{\text{supply}} - 30 \varepsilon_{[t]} \cdot \text{net} P_{[t]} \quad (15)$$

where $m$ is the molar quantity of silicon (mmol m$^{-2}$) and $F_{\text{supply}}$ the Si(OH)$_4$ supply (mmol m$^{-2}$ d$^{-1}$).

The biogenic silica pools are described as follows:

$$\frac{d(m_{\text{bSiO}_2})}{dt} = \text{net} P_{[t]} - E_{[t]} \quad (16)$$

$$m_{\text{bSiO}_2} \cdot \frac{d(\delta_{30} \text{Si}_{\text{bSiO}_2})}{dt} = \text{net} P_{[t]}$$

$$\left(\delta_{30} \text{Si}_{\text{bSiO}_2} + 30 \varepsilon_{[t]}\right) - E_{[t]} \cdot \delta_{30} \text{Si}_{\text{bSiO}_2} \quad (17)$$
Superscript “export” is the biogenic silica exported out of the mixed layer assuming that the exported fraction of biogenic silica has an isotopic composition similar to the one in the mixed layer. This could be an oversimplification since Krause et al. (2010) suggested that for the Equatoriocal Pacific exported biogenic silica can be composed of mainly detrital diatom cells.

Small errors arise for both mixing and fractionation when the δ notation is used instead of isotopic ratios. However, for δ values close to 0, as is the case for Si isotopic system, these sources of errors are negligible (Fry, 2006). The aim here is to assess, with the current knowledge of the Si biogeochemical dynamic, how the seasonal variability of the different isotopic effects potentially affects the mass and isotopic balances. This is of crucial importance for palaeoceanographic reconstructions of past ocean conditions. This approach should not be taken as an attempt to quantify fluxes (uptake, dissolution, export, and Si(OH)₄ supply) or to constrain the complexity of the seasonal silicon cycle.

### 4.1.3 Model outputs

With the current knowledge of Si biogeochemical dynamic in the Southern Ocean, the model seems to adequately reproduce the seasonal evolution of silicic acid and biogenic silica concentrations in the PFZ mixed layer (Fig. 5; Brzezinski et al., 2001; Quégúiner and Brzezinski, 2002). The integrated Si-uptake of 1.6 mol Si m⁻² yr⁻¹ fits relatively well with the mean expected values in the Polar Front Zone, 1.2 mol Si m⁻² yr⁻¹ (Fripiat et al., 2011c).

The simulated isotopic trends for δ³⁰Si_bSiO₂ differ from the δ³⁰Si₅SiOH₄ trends, especially at the end of the productive period (Fig. 6a). In the PFZ the Si uptake:supply ratio is high at the beginning of the growth period, and, consequently, the δ³⁰Si₅SiOH₄ and δ³⁰Si_bSiO₂ evolutions roughly follow a Rayleigh distillation (Figs. 5d and 6c). Note that the steady-state model is unable to reproduce the measured ML δ³⁰Si_bSiO₂ value and is in disagreement with the present simulation (Fig. 6d). Once the silicic acid concentration has reached a level that limits Si-uptake (Eq. 8), the Si uptake:supply ratio decreases and the supply by vertical mixing of lighter Si isotopic signatures into the Si-depleted mixed layer decreases the δ³⁰Si₅SiOH₄ (Figs. 5d and 6). The δ³⁰Si_bSiO₂ is affected differently, since both new and preceding diatom generations set the ML δ³⁰Si_bSiO₂ (Eq. 14), especially at the end of the productive period when bSiO₂ production is low. Such process decreases the Δ³⁰Si and is consistent with our observations (Figs. 2b, 3b, and 7a). The
Importantly, δ30Si values observed only in the PFZ and PF (Fig. 2b), both characterized by heavily Si-depleted mixed layer together with a large vertical silicic acid gradient. The relationship between the accumulated exported δ30SiO2 and relative Si-utilization was not significantly different from the simulation with the accumulated product (sensu Rayleigh; Fig. 7b), both with and without isotopic fractionation during bSiO2 dissolution. Therefore, for reconstructing past Si(OH)4 utilisation in the ACC surface waters from sedimentary δ30SiO2 data, our results indicate that a Rayleigh bSiO2 accumulated product model is more appropriate than a steady-state model. This mainly results from the

Fig. 7. (a) Simulated Δ30Si (= δ30Si(OH)4 – mixed layer δ30Si(OH)2) vs. relative Si-utilisation in the mixed layer (=|Si(OH)4|/|Si(OH)4|) without (green line) and with (red line) an isotopic fractionation during bSiO2 dissolution. The isotopic difference between the Rayleigh δ30Si(OH)4 and either Rayleigh instantaneous δ30SiO2 (full blue line) or Rayleigh accumulated δ30SiO2 (dashed blue line) are also shown. (b) Simulated relationship between accumulated exported δ30SiO2 and relative Si-utilization in the mixed layer without (green line) and with (red line) an isotopic fractionation during bSiO2 dissolution. The relationship with the Rayleigh accumulated δ30SiO2 (full blue line; 1 standard deviation = dashed blue lines) is also shown to compare.

Impact of isotopic fractionation during dissolution – by adding the isotopic fractionation of bSiO2 dissolution (Demarest et al., 2009), the δ30Si(OH)4 gets slightly lighter and the δ30SiO2 slightly heavier (Fig. 6a), relative to the model runs without such fractionation, throughout the productive period, in agreement with a decrease in the net fractionation extent (Eq. 7). This process together with the low Si-uptake:supply ratio could explain the lowest Δ30Si measured at the end of the productive period (Fig. 7a). The latter could bias the relationship between the exported δ30SiO2 and the relative silicic acid utilization (= f). Nevertheless, such bias is lying within isotopic analytical precision (Fig. 7b) and therefore should not impact on the use and interpretation of this proxy in palaeoceanography. Furthermore, the isotopic effect of dissolution could have been over-expressed in this simulation since the D:P ratio at the bSiO2 productive peak was already high (0.4) in comparison with published values, which are close to 0.1 (Brzezinski et al., 2003).

Impact of mixing events – by switching from a continuous supply to a pulsed supply mode, consisting of 10 consecutive instantaneous mixing events, the simulation is scattered around the continuous supply with no effect on biogenic silica isotopic composition (Fig. 6b). Moreover, in reality, mixing is probably spatially heterogeneous and lateral mixing should smooth the trends toward the mean continuous supply.

As a comparison, similar simulations have been carried out in the SAZ, an area characterized, in contrast to PFZ, by a lower vertical silicic acid gradient, and in the AZ characterized by an incomplete Si depletion of the summer mixed layer (Fripiat et al., 2011b). The model’s initial conditions (i.e. before the start of the growth season) are set using the Si-characteristic of subsurface waters, respectively, in the SAZ and AZ by Fripiat et al. (2011b; median for the thermocline and the halocline, see Table 2 for initial and final conditions), assumed here to represent the winter mixed layer in both areas (Tomczak and Godfrey, 2001). In order to agree as much as possible with the PFZ simulation, the Si-supply over bSiO2 production ratio has been fit at ~0.3 and the D:P ratio at the productive peak at ~0.4. For the same relative vertical mixing, the vertical supply of light Si-isotopes at the end of summer (low Si-uptake:supply ratio) is not sufficient to lower the mixed layer δ30Si(OH)4 (Fig. 8). Therefore, it enables us to explain low Δ30Si values observed only in the SAZ and PFZ (Fig. 2b), both characterized by heavily Si-depleted mixed layer together with a large vertical silicic acid gradient.

Note that this simulation is already sufficient to explain partly the measured range in the Δ30Si (Figs. 3 and 7a).

exported bSiO2 out of the mixed layer fits with an accumulation product (sensu Rayleigh; Fig. 6c). This is seen from (1) Fig. 5d showing that ~85% of the export of biogenic silica occurs before the decreasing trend in δ30Si(OH)4 and, incidentally, the exported δ30SiO2 is not significantly affected by processes occurring at the end of the productive period when the Si uptake:supply ratio is low; (2) Fig. 7b showing the relationships (non-significantly different) between the simulated exported δ30SiO2 and the accumulated biogenic silica (sensu Rayleigh) with relative silicon utilisation (= f).

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The fate of biogenic silica δ^{30}Si through the water column is of prime importance to link the sedimentary δ^{30}Si to the surface environmental conditions in which they were generated. Dissolution during settling could affect the δ^{30}Si subSiO₂ distributions by shifting the preserved material toward more positive values (Demarest et al., 2009). By analysing the δ^{30}Si of sinking biogenic silica (two sediment traps in the AZ at 1031 and 2182 m in the Pacific sector; Varela et al., 2004) a remarkable agreement with surface values was observed. This is further confirmed by the present study which present the first whole water column profiles of suspended biogenic silica δ^{30}Si signatures (Fig. 4). The isotopic composition of subsurface suspended biogenic silica generally reflects that of the mixed layer. It thus seems that dissolution does not significantly affect the δ^{30}Si signal of exported bSiO₂. However, there are two notable exceptions:

1. In the WG, the δ^{30}Si of bSiO₂ is becoming heavier by 0.4 to 0.8‰ below 1800 m (Fig. 4). Since no values are available between 700 and 1800 m we do not know whether this shift is progressive or only located at a specific depth. Cavagna et al. (2012) for the same stations, observed that the δ^{13}C composition of deep ocean particles during BGH is consistent mostly with a 1-D vertical transport. This would preclude lateral advection of particles as the reason for such offset in Si isotopes. Consequently, it cannot be ruled out that the δ^{30}Si of bSiO₂ particle water column profile of the WG station is biased by a progressive dissolution of biogenic silica, preferentially releasing light silicon isotopes.

2. The strong δ^{30}Si lightening, by 0.6‰, below the mixed layer in the WG might result from biogenic silica produced from a subsurface Si(OH)₄ pool with lighter δ^{30}Si (OH)₄, as is present at such depths (Table 1; Fripiat et al., 2011b). These depths extend below the euphotic layer (1 % PAR). Several studies have shown that energy involved in the silification process is mainly of respiratory origin and therefore decoupled from photosynthesis (Martin-Jézéquel et al., 2000).
allowing Si-uptake below the euphotic layer (Brzezinski et al., 1989).

5 Conclusions

With a palaeoceanography perspective this study has attempted to assess the controls upon, and the seasonal evolution of, biogenic silica isotopic compositions in the mixed layer and its transfer across the water column. This data set acquired in late austral summer highlights two main points:

1. A large latitudinal variation in $\delta^{30}$Si$_{bSiO_2}$ is observed in the mixed layer across the meridional BGH transect, contrasting with a narrower variation for $\delta^{30}$Si$_{Si(OH)_4}$. This decoupling cannot be reproduced by Rayleigh and steady-state isotopic fractionation models, which directly link the $\delta^{30}$Si to relative Si(OH)$_4$ utilisation. These model failures probably result from the fact that (i) a constant isotopic fractionation is assumed over the entire period of silicate depletion and/or (ii) the seasonal expressions of the different isotopic effects are not appropriately taken into account (i.e. only Si-uptake is considered). We implemented an isotopic fractionation model to describe the seasonal $\delta^{30}$Si variation in the mixed layer. From this analysis, it seems that the exported biogenic silica is rather well described by the Rayleigh accumulated product. The failures of the existing models are especially significant at the end of the productive period when (i) biogenic silica production and export are low; (ii) bSiO$_2$ dissolution:production ratio is high, imposing a lower apparent fractionation factor; and (iii) Si-supply:Si-uptake ratio is high. The latter effect is especially expressed when the summer mixed layer is strongly Si-depleted together with a large silicic acid vertical gradient, as is the case in the southern ACC. Despite being very hypothetical, we cannot rule out a spatial and seasonal variation in the fractionation factor associated with the net bSiO$_2$ production. However, it seems unlikely that such variation alone is sufficient to explain the observations.

2. Except below 1800 m in the Weddell Gyre, where we cannot rule out a potential bSiO$_2$ dissolution effect, the deeper $\delta^{30}$Si$_{bSiO_2}$ is generally similar to the mixed layer $\delta^{30}$Si$_{bSiO_2}$, which supports the absence of an isotopic effect during particle settling through the water column in most cases.

Acknowledgements. Our warm thanks go to the officers and crew of the R/V Marion Dufresne during the BONUS-Goodhope cruise, as well as to S. Speich and M. Boyé as program leaders. We are also grateful to J. De Jong and N. Mattielli for the management of the MC-ICP-MS laboratory at ULB and to L. Monin and N. Dahkani (RMCA) for their help in sample processing and to Virginia Panizzo (ULB) for correction of English. This work was conducted within the BELCANTO III – BIGSOUTH networks (contracts SD/CA/03A of SPSDIII, Support Plan for Sustainable Development) funded by BELSPO, the Belgian Science Policy. Luc André thanks the “Fonds National de la Recherche Scientifique” (FNRS, Belgium) for its financial support (FRFC project 2.4512.00). François Fripiat and Anouk de Brauwere are post-doctoral fellows with the FNRS.

Edited by: A. Shemesh

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