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Seasonal evolution of net and regenerated silica
 1
     production around a natural Fe-fertilized area in the
 2
     Southern Ocean estimated from Si isotopic approaches
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18
      Abstract
      A massive diatom-bloom is observed each year in the surface waters of the naturally Fe
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      fertilized Kerguelen Plateau (Southern Ocean). We measured biogenic silica production and
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      dissolution fluxes (pSi and pDiss respectively) in the mixed layer in the vicinity of the
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      Kerguelen Plateau during austral spring 2011 (KEOPS-2 cruise). We compare results from a
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      High-Nutrient Low-Chlorophyll reference station and stations with different degrees of iron
      enrichment and bloom conditions. Above the Plateau biogenic pSi are among the highest
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     reported so far in the Southern Ocean (up to 47.9 mmol m^{-2} d^{-1}). Although significant (10.2
25
      mmol m^{-2} d^{-1} in average), pDiss were generally much lower than production rates. Uptake
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     ratios (pSi:pC and pSi:pN) confirm that diatoms strongly dominate the primary production in
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28 this area. At the bloom onset, decreasing dissolution to production ratios (D:P) indicate that 29 the remineralization of silica could sustained most of the low silicon uptake and that the 30 system progressively shifts toward a silica production regime which must be mainly 31 supported by new source of silicic acid. Moreover, by comparing results from the two 32 KEOPS-expeditions (spring 2011 and summer 2005), we suggest that there is a seasonal 33 evolution on the processes decoupling Si and N cycles in the area. Indeed, the consumption of 34 H₄SiO₄ standing stocks occurs only during the growing stage of the bloom when strong net silica production is observed, contributing to a higher H₄SiO₄ depletion relative to NO₃⁻. 35 36 Then, the decoupling between H₄SiO₄ and NO₃⁻ is mainly controlled by the more efficient 37 nitrogen recycling relative to Si. Gross-Si:N uptake ratios were higher in the Fe-rich regions 38 compared to the HNLC area, likely due to different diatoms communities. This suggests that 39 the diatom responses to natural Fe fertilization are more complex than previously thought, and 40 that natural iron fertilization over long time scales does not necessarily decrease Si:N uptake 41 ratios as suggested by the Silicic Acid Leakage Hypothesis. Finally, we propose the first 42 seasonal estimate of Si-biogeochemical budget above the Kerguelen Plateau based on direct 43 measurements. This study points out that naturally iron fertilized areas of the Southern Ocean 44 could sustain very high regimes of biogenic silica production, similar to those observed in 45 highly productive upwelling systems.

46

47 **1** Introduction

48 Covering 20 % of the World Ocean, the Southern Ocean is considered as a crucial component 49 of the climate system since it represents a net sink for atmospheric CO₂ (Takahashi et al.,

50 2009). It also plays a key role in the global silicon (Si) biogeochemical cycle because diatoms,

a siliceous phytoplankton group, are one of the major primary producers in this area

52 (Buesseler et al., 2001; Quéguiner and Brzezinski, 2002; Tréguer and De la Rocha, 2013). As

53 their cell wall is composed of biogenic silica (opal, amorphous SiO₂.nH₂O, hereafter referred

54 to as BSi), diatoms take up dissolved silicon (hereafter referred to as DSi) in the form of

silicic acid (H_4SiO_4), to produce their siliceous frustules. At global scale, 56 % of this gross

56 production is estimated to be directly recycled in the upper 100 m (Tréguer and De La Rocha,

57 2013) due to the combined effects of both physico-chemical and biological processes

58 (Kamatani, 1982; Bidle and Azam, 1999; Ragueneau et al., 2000). Only the material escaping

59 dissolution is exported toward the deep ocean and eventually buried in sediments.

60 Consequently, the marine Si biogeochemical cycle is dominated by biogenic silica production

61 and dissolution in the surface mixed layer, and one atom of Si undergoes a cycle of biological 62 uptake by diatom and subsequent dissolution about 25 times before being removed to the 63 seabed (Tréguer and De La Rocha, 2013). Thus, it is essential to estimate the balance between 64 silica production and dissolution in the euphotic zone which is best illustrated by the integrated dissolution to production rate ratio (D: P; Brzezinski et al., 2003) or by integrated 65 net production rate ($\int \rho Si_{Net}$ = production minus dissolution). Globally, the $\int D$: P values present 66 67 an annual mean of 0.56 (Tréguer and De La Rocha, 2013) and range from < 0.1 to > 1, with 68 low D:P values associated to diatom bloom events, and D:P values exceeding 0.5 occurring 69 during non-bloom periods (Brzezinski et al., 2001). However, the number of D:P estimates, 70 due to a small number of Si uptake measurements and an even lower number of Si dissolution 71 measurements, is insufficient compared to the high variability observed regionally and 72 seasonally in the ocean which implies high uncertainty in the global D:P estimate and overall 73 on the marine silicon budget.

74 Diatoms are ecologically widespread and dominate the primary production in the Antarctic 75 Circumpolar Current (ACC), especially south of the Polar Front (PF), where their productivity 76 accounts for 1/3 of the global marine silica production (Pondaven et al. 2000; Buesseler et al. 77 2001). Consequently, this part of the Southern Ocean represents a key study area to improve 78 our understanding of the global biogeochemical cycles of both carbon and silicon. Biological 79 processes occurring in the Southern Ocean have indeed a significant impact on global 80 biogeochemistry. For example, the large H₄SiO₄ utilization by diatoms in the ACC, combined 81 to the global overturning circulation would determine the functioning of the biological pump 82 of low latitude areas by inducing a strong silicic acid limitation (Sarmiento et al., 2004). In the 83 Southern Ocean, a much larger depletion of silicic acid than nitrate in surface waters occurs 84 (Trull et al, 2001), which results from the action of a silicon pump, i.e. the preferential export 85 of BSi compared to particulate organic nitrogen (PON; Dugdale et al., 1995). This area is also 86 the largest High Nutrient Low Chlorophyll (HNLC) zone of the global ocean where dissolved 87 iron limitation plays a fundamental role in regulating the primary production and the carbon 88 cycle (De Baar et al., 2005; Boyd et al., 2007; Tagliabue et al., 2012). Indeed, phytoplankton 89 community structure and nutrient cycling could be largely controlled by Fe availability, with 90 highest growth rates located close to iron sources such as continental margins, island systems 91 and frontal regions (Blain et al., 2007; Tagliabue et al., 2012).

- 92 In this context, the KErguelen Ocean and Plateau compared Study (KEOPS) program,
- 93 consisting of two expeditions (late summer 2005 and early spring 2011), was conducted to

investigate a naturally iron-fertilized area located in the Indian sector of the Southern Ocean, 94 95 where the iron availability could potentially favor the carbon and silicon biological pumps 96 (Fig. 1; Blain et al., 2007). Two massive and complex blooms, which are clearly constrained 97 by the local bathymetry, are observed annually over the Kerguelen Plateau and contrast with 98 the HNLC character of surrounding waters (Pollard et al., 2002; Mongin et al., 2008). The 99 first KEOPS expedition (KEOPS-1, January-February 2005) has highlighted the impact of 100 natural iron fertilization on primary production and nutrient cycling, as well as the advantages 101 to study natural laboratories in the context of such ocean fertilization (Blain et al., 2008). The 102 general purpose of KEOPS-2 (October-November 2011) was to improve our knowledge about 103 the processes responsible for this iron fertilization and its impact on the seasonal variations of 104 the mechanisms controlling the primary production and carbon export. While the KEOPS-1 105 cruise was mainly directed towards the study of the bloom in the South-East area of the 106 Plateau, KEOPS-2 focused mainly on the bloom located North-East of the Kerguelen Islands

107 above the Kerguelen abyssal plain.

In this paper, we investigate the spatial and seasonal variability of silica production and dissolution in the surface waters of the Kerguelen area. The specific objectives are the following:

111 - Compare the Si cycle dynamics in contrasting productive environments such as the

112 southeastern Kerguelen Plateau bloom, the northeastern Kerguelen bloom in a stationary

113 meander southward the PF and the warmer waters located north of this front, relative to the

114 upstream HNLC area south-west of Kerguelen Islands, and identify controlling processes.

115 - Determine and quantify the seasonal evolution of processes which drive the Si

116 biogeochemical budget in the upper layer of the Kerguelen Plateau, using the ³⁰Si stable

117 isotope method (Nelson and Goering, 1977a; Fripiat et al, 2009) applied during KEOPS-2 and

118 other techniques of mass and isotopic balance used during KEOPS-1 (³²Si radiogenic tracer

119 incubations; Mosseri et al., 2008 and natural silicon isotopic composition, δ^{30} Si; Fripiat et al.,

120 2011a), in order to fully characterize the silicon cycle above the Kerguelen Plateau.

121 - Discuss the potential role of Fe on silica production – dissolution and on Si:N uptake ratios

122 in the context of the Silicic Acid Leakage Hypothesis (Brzezinski et al., 2002; Sarmiento et

123 al., 2004).

- 124 - Finally, compare our dataset with previous results in other productive regions of the global 125 ocean and discuss the different types of diatom dominated regimes.
- 126

127 2 Material and methods

KEOPS-2 sampling campaign 128 2.1

The KEOPS-2 cruise was conducted in the Indian sector of the Southern Ocean during the 129 austral spring 2011 (from October 10th to November 20th) on board the R/V Marion Dufresne 130 (TAAF/IPEV) and was focused on the iron-fertilized blooms observed around the Kerguelen 131 132 Plateau region. This plateau is a large area of relatively shallow seafloor that acts as a barrier 133 to the circumpolar flow of the ACC, forcing a large part of the current to pass north of the 134 plateau. The remaining flow passes south of the Kerguelen Islands and forms the jet of the PF, 135 which exhibits strong meandering and eddy activity (Park et al., 1998, 2008, Roquet et al., 136 2009). As a consequence, the shallow region located south of Kerguelen Island represents a 137 zone of weak north-eastward circulation (Park et al., 2008; Roquet et al, 2009) and bears a 138 capacity of high chlorophyll a and BSi accumulation during phytoplankton blooms (Blain et 139 al., 2001; Mosseri et al., 2008; Mongin et al., 2008). Over the plateau, enhanced vertical 140 mixing associated to internal waves interact with the local bathymetry (Park et al., 2008) and 141 supply iron and macronutrients from depth to surface waters, enabling to fuel phytoplankton 142 bloom (Blain et al., 2007; Fripiat et al., 2011a). 143

- The cruise included 8 long-term stations devoted to process studies with incubation
- 144 experiments (Fig. 1). Except for one station (E4E) where we were not able to measure silica
- 145 dissolution, Si fluxes were investigated in all these process-stations which characteristics are

146 presented in table 1:

- 147 - A HNLC reference station (R-2) located in deep waters south-west of Kerguelen Islands.
- 148 - The Kerguelen Plateau bloom reference station of KEOPS-1 (A3-2).
- 149 - A productive open ocean station (F-L) influenced by warmer Sub-Antarctic Surface Water, 150 located north of the Polar Front.
- 151 - A productive station (E-4W) located in the plume of chlorophyll observed downstream of 152 the plateau and close to the jet induced by the PF.

153 - 4 stations (E-1 to E-5) constituting a pseudo-lagrangian survey located in a complex

recirculation zone in a stationary meander of the Polar front characterized by strong

155 mesoscale activity (Zhou et al., 2014).

156 **2.2** Sample collection, spike and incubation conditions

157 The isotopic dilution technique adapted by Fripiat et al. (2009) from Corvaisier et al. (2005) 158 aims at simultaneously determining the rates of Si uptake (i.e. silica production) and of 159 biogenic silica dissolution in the same seawater sample. After spiking with a solution enriched 160 in ³⁰Si followed by incubation of the samples, the production rate is estimated from the 161 change in isotopic composition of the particulate phase (increase in ³⁰Si). Similarly, the 162 isotopic dilution (increase in ²⁸Si) in the ³⁰Si enriched seawater, due to the dissolution of 163 naturally ²⁸Si enriched BSi initially present, is used to estimate the dissolution rate.

164 Production and dissolution rates were determined at 7 and 5 depths respectively,

165 corresponding to different levels of Photosynthetically Active Radiation (PAR), from 75 % to

166 0.1 % of surface irradiance. Seawater was collected at defined depths in the euphotic layer

167 using Niskin bottles mounted on a CTD-rosette. For each depth, 5 l of seawater were sampled.

168 1 l was subsampled to obtain a natural silicon isotopic standard (i.e. not spiked with 30 Si) to be

169 processed along with the samples to correct for the matrix effect and mass bias during isotopic

analysis (Fripiat et al., 2009). These unspiked samples were immediately filtered on 0.8 μ m

171 Nuclepore polycarbonate membranes to separate biogenic silica from silicic acid. The

172 membrane was dried at 50 $^{\circ}$ C overnight and the filtrate was directly preconcentrated (see

173 section – Sample preparation and isotopic measurements) and stored at room temperature in

the dark.

175 The remaining seawater volume was subsampled in 2 l aliquots spiked with $H_4^{30}SiO_4$ -

176 enriched solution (99 % ³⁰Si). Aliquots devoted to production measurements were spiked with

a spike contribution representing usually less than 10 % of natural concentrations to minimize

178 the perturbation of the natural DSi contents (Nelson & Goering, 1977a). In order to improve

the detection limit of the method for dissolution, a second 21 aliquot was spiked by adding

³⁰Si in the same amount as natural DSi (i.e. DSi spike addition at 100 % of the initial DSi).

181 This provided sufficient sensitivity for the isotopic measurements of dissolution (see section –

182 Accuracy of the model, detection limit and standard deviation).

183 Immediately after spike addition and gentle mixing, 1 l was filtered following the same 184 procedure than for the unspiked standard, to determine the initial conditions (t_0) . The second 185 half of the sample was poured into polycarbonate incubation-bottles and incubated under light 186 conditions simulating those prevailing *in situ* for 24 h (10 % spiked samples) and for 48 h 187 (100 % spiked samples). Deck-incubators were fitted with blue plastic optical filters to 188 simulate the light attenuation of the corresponding sampling depths, and temperature was 189 regulated by circulating surface seawater. At the end of the incubation period, samples were 190 filtered and treated as described above to characterize the final conditions of the incubation 191 $(t_{24} \text{ or } t_{48}).$

192 **2.3** Sample preparation and isotopic measurements

193 Preconcentration of H₄SiO₄ in the seawater samples (for both production and dissolution 194 measurements) was applied on-board to increase the Si:salinity ratio, because the maximum 195 salinity of the solution that can be introduced in the mass spectrometer is about 2 % (Fripiat 196 et al., 2009). This step was achieved using a protocol adapted from the MAGIC method (Karl 197 & Tien, 1992; Reynolds et al., 2006). The H₄SiO₄ in seawater was scavenged by the brucite 198 precipitate (Mg(OH)₂) obtained by adding 1 ml of 14 N NaOH to the 1 l of seawater sample 199 and strong stirring. The precipitate was recovered by decantation and centrifugation, and was 200 then dissolved in 3 ml of 3 N HCl.

201 In the shore based laboratory, polycarbonate membranes (t_0 , t_{24} and t_{48} for both production and

202 dissolution measurements) were digested in one step using a protocol adapted from

203 Ragueneau et al. (2005) with 4 ml of 0.2 N NaOH during 40 min at 100 °C to hydrolyse BSi.

204 Samples were then neutralized with 1 ml of 1 N HCl to stop the reaction.

205 An aliquot of the solutions obtained after preconcentration and digestion was used to

206 determine colorimetrically the DSi and BSi concentrations, following the method of

207 Strickland and Parsons (1972). The remaining sample was diluted to 100 ppb Si in a 2 %

208 HNO₃ solution to determine the initial and final Si isotopic composition of the dissolved and

209 particulate phases using a Element 2 (Thermo-Fischer) HR-SF-ICP-MS (High Resolution –

210 Sector Field – Inductively Coupled Plasma – Mass Spectrometer) with the same configuration

211 used by Fripiat et al. (2009). The sequence of analysis consists in: blank – natural standard –

- spiked sample 1 natural standard spiked sample 2 natural standard spiked sample 3 –
- 213 natural standard blank. The average of the two blanks were subtracted to each standard and

- sample. To test whether our dissolution measurements were biased by a ³⁰Si contamination
- 215 linked to a possible memory effect in the HR-SF-ICP-MS, we compared the average
- 216 composition of the first natural standards (i.e., without contamination from memory effect, n
- 217 = 55) with the composition of natural standards analyzed after a spiked sample (n = 102).
- 218 There was no significant difference between natural standards passed before and after a 100
- 219 % DSi spiked sample (T-test, p-value <0.001). We can thus exclude significant memory effect
- 220 when applying the analytical sequence described above.
- 221

222 **3 Results**

3.1 Accuracy of the model, detection limit and standard deviation

224 To estimate the production and dissolution of biogenic silica (pSi and pDiss, respectively), 225 two different models are available: the linear one-compartmental model described by Nelson 226 and Goering (1977a, b) and the non-linear two-compartmental model described in de 227 Brauwere et al. (2005) and Elskens et al. (2007). In the latter, both isotopic composition and 228 concentration changes occurring during the incubation time are taken into account to estimate 229 production and dissolution rates simultaneously. Lack of consideration of these changes could 230 induce significant biases in the estimated fluxes (Elskens et al., 2007). In this model the fluxes 231 are calculated by resolving a system of 4 equations given by:

232
$$[DSi]_t = [DSi]_{t0} + (\rho Diss - \rho Si) \times t$$
(3)

233
$$[BSi]_t = [BSi]_{t0} \times (\rho Si - \rho Diss) \times t$$
(4)

234
$$\alpha DSi_t = \alpha DSi_{t0} \times \left(1 + \frac{\rho Diss - \rho Si}{[DSi]_{t0}} \times t\right)^{\frac{\rho Diss}{\rho Si - \rho Diss}}$$
(5)

where [BSi] and [DSi] are the dissolved silicon and biogenic silica concentrations (in µmol l⁻
¹); æBSi and æDSi are the abundance in excess of ³⁰Si (measured minus natural abundances)
in the particulate and dissolved phase respectively; the subscribes t0 and t refer to the initial
and final incubation values.

The best solution is found numerically by optimizing parameter values (ρ Si and ρ Diss) and minimizing the cost function (weighted sum of squared differences between calculated and measured variables, [BSi], [H₄SiO₄], æBSi and æDSi for the four equations simultaneously; de Brauwere et al., 2005; Elskens et al., 2007).

244 The relevance of the 2 models against a given data set has already been discussed by Elskens 245 et al. (2007) and Fripiat et al. (2011b). Taking into account these considerations, and after 246 testing the accuracy and the sensitivity of each model, we use preferentially the non-linear 2 247 compartmental model to estimate the biogenic silica production and dissolution during 248 KEOPS-2. This model was tested according to the four criteria and the residual of the cost function was checked to follow a Chi² distribution as detailed in Elskens et al. (2007). Due to 249 250 unexpected sampling problems on-board, we were not able to measure [DSi]_t. Thus, in 251 addition to the biogenic silica production and dissolution rates, this variable was also 252 estimated by the model (Eqs. 3-6). Under these conditions, one degree of freedom is lost but 253 the system remains identifiable with 3 unknowns and 4 equations.

KEOPS-2 took place during the onset of the blooms, the biogenic silica production rates were
quite high and far above the detection limit, except for 3 depths of the HNLC reference station
(R-2) and for the deepest value at each station (0.01 % PAR attenuation depth, 8 samples).
However, since biogenic silica dissolution rates were expected to be low in early spring, it is
essential to determine the limit of detection for the ³⁰Si isotopic dilution.

259 In most cases, final æDSi were significantly different from initial æDSi (paired T-test, p-value < 0.001). The detection limit for isotopic dilution was then estimated as being the lowest 260 difference between initial and final ³⁰Si isotopic abundances ($\Delta \alpha$ DSi) measurable by the 261 262 instrument. Every æDSi solutions have been analyzed in duplicates with a pooled standard 263 deviation of 0.32 % (n = 35). In addition, we analyzed the same in-house standard several 264 times during every analytical session. This solution was a 10 % spiked seawater from 265 Southern Ocean analyzed since several years with a α DSi at 11.83 \pm 0.43 % (n = 40). The 266 relative standard deviation (RSD) on α DSi of this standard solution is 0.43 % (n = 40) and 267 represents the long-term reproducibility of HR-SF-ICP-MS measurements. Therefore, each 268 KEOPS-2 incubation with a $\Delta \approx$ DSi between t₀ and t₄₈ higher than this RSD was considered to be significantly different from zero, and hence above the detection limit. This was the case for 269 almost all the KEOPS-2 dataset (see e.g. Fig. 2), except for 7 values showing a change in ³⁰Si 270

- abundance below the detection limit. This included 4 samples from the HNLC reference
- station R-2 where biological activity was extremely low.
- 273 Due to time and sampled water volume constraints, the sampling strategy adopted for
- 274 KEOPS-2 gave the priority to highest vertical resolution instead of replicate incubations.
- 275 Since only the analytical reproducibility was taken into account in the model, the standard
- 276 deviations on Si uptake and dissolution rates were likely to be underestimated. Therefore we
- will use a theoretical relative precision for the whole incubation experiments of 10 %, as
- estimated for Si uptake rates by Fripiat et al. (2009).

279 **3.2** Physical, chemical and biological parameters

The vertical structure of upper layer waters in the area was characteristic of the Antarctic Surface Water in the vicinity of the Polar Front (Park et al., 1998, 2008). The Winter Water (WW), identified by the minimum of temperature centered around 200 m, was capped by a homogeneous mixed layer (ML) induced by seasonal stratification. The boundary between the surface ML and the WW is usually marked by a strong seasonal pycnocline. However, at some stations, the stratification of the surface layer was relatively complex and showed two successive discontinuities evidenced by two different density gradients as indicated in Fig. 3.

During KEOPS-2, the surface ML depth, defined by the density difference of 0.02 from the surface (Park et al, in prep.), showed a large variability between stations (Fig. 3). A strong and shallow stratification was measured north of the polar front, while wind events induced weak stratification and deep ML in the stations above the plateau and in the HNLC area. Stations in the recirculation zone (E-1 to E-5) supported a complex stratification due to their highly spatial and temporal dynamic and were characterized by 2 distinct density discontinuities.

- All the stations located south of the Polar Front had quite homogeneous Chl-*a*, BSi and DSi
- stocks from the surface to the deepest density discontinuity (below the so-called ML; Fig. 3).
- 295 North of the Polar Front, these stocks were higher at the surface and decreased with depth
- 296 (Fig. 3c). Stations A3-2 and E-4W present similar BSi and DSi surface concentrations (Fig.
- 297 3e, g). At these 2 stations, DSi concentrations increase gradually while Chl-a and BSi
- 298 decrease drastically below the deepest density discontinuity. Station R-2 contrasted from the
- 299 latter stations by its low BSi, low Chl-*a* content and relatively high DSi concentrations,
- 300 confirming its HNLC character (Fig. 3a). During the lagrangian survey (stations E-1, E-3, E-
- 4E and E-5), we observed a DSi depletion from ≈ 15 to $\approx 10 \ \mu mol \ l^{-1}$ in surface waters, an

302 increase of Chl-*a* concentrations from < 1 to > 1 mg m⁻³ and a doubling of the BSi content 303 from ≈ 1.5 to > 3 µmol l⁻¹ (Fig. 3b, d, f, h). Such temporal variations were mainly driven by 304 diatom production as described below.

305 3.3 Biogenic silica production and dissolution rates

Silica production rates were quite homogeneously distributed in the euphotic layer with an exception for the station F-L located north of the Polar Front where it decreases progressively with depth (Fig. 4a). Surface ρ Si varied from $0.036 \pm 0.003 \mu mol l^{-1} d^{-1}$ (R-2 in the HNLC area) to $1.28 \pm 0.12 \mu mol l^{-1} d^{-1}$ (A3-2, above the Plateau). All over the study area, Si uptake rates reached very low values at the base of the euphotic layer. Note that the same decreasing trend was also observed in primary production experiments performed in parallel (see e.g. Cavagna et al., in prep.).

313 BSi dissolution rates were considerably lower than Si uptake rates except in the HNLC area 314 (R-2) and at station E-3 where pSi was in the lower range of the KEOPS-2 dataset. Vertical 315 profiles of pDiss (Fig. 4b) were quite homogeneous from the surface to the base of the 316 euphotic layer and did not increase at depth. This indicates that, the physical and 317 biogeochemical processes affecting BSi dissolution did not vary significantly over the water 318 column. This is also consistent with the low accumulation of biogenic silica observed at depth 319 in spring (Lasbleiz et al., 2014) which contrasts with the occurrence of deep BSi maxima at 320 the end of summer (Mosseri et al., 2008). Moreover, silica dissolution rates were not 321 significantly different between bloom stations, and were comparable to those measured by 322 Brzezinski et al. (2001) for the same season in the Pacific sector and by Beucher et al. (2004) 323 and Fripiat et al. (2011b) for the end of summer in the Australian sector.

324 As silica production was close to zero below the euphotic layer, all the vertically integrated

325 values presented in table 2 were calculated from 100 % to 1 % of the surface PAR. The

326 integrated Si uptake rates ($\int \rho Si$) varied from 3.09 ± 0.01 mmol m⁻² d⁻¹ (R-2, in the HNLC

327 area) to 47.9 ± 0.4 mmol m⁻² d⁻¹ (A3-2, above the Plateau), and were among the highest

reported so far in the Southern Ocean (see review in Fripiat, 2010). Integrated BSi dissolution

329 rates ($\int \rho Diss$) were generally much lower than integrated production rates with values ranging

330 from 3.79 \pm 0.03 mmol m⁻² d⁻¹ north of the Polar Front (F-L) to 9.99 \pm 0.03 mmol m⁻² d⁻¹ at E-

331 3. Because pDiss did not vary over depth, or between stations, integrated dissolution

- estimates were correlated with the depth of the euphotic layer (Ze), with higher values in
- stations with deeper Ze, e.g. E-1 and E-3 ($R^2 = 0.83$, not shown).
- 334 Net production rate of BSi in the euphotic layer ($\int \rho Si_{net}$) represents the difference between
- 335 gross silica production and dissolution rates (Fig. 5) and could be associated to an uptake of
- 336 "new-H₄SiO₄" i.e. uptake that does not come from remineralisation processes within the ML.
- 337 As for the net primary production, the net silica production could be defined as the part of the
- 338 BSi that accumulates in the surface layer during the productive period, which would then be
- potentially available later for export to the mesopelagic layer (Brzezinski et al., 2001;
- 340 Quéguiner, 2013). During the pseudo-lagrangian survey, net silica production was quite low
- during the first 2 visits (E-1 and E-3 with respectively 9.6 ± 0.1 and 0.5 ± 0.1 mmol m⁻² d⁻¹)
- and reached the maximal value at the last visit (E-5, $20.5 \pm 0.2 \text{ mmol m}^{-2} \text{ d}^{-1}$). The highest net
- 343 production rate was observed above the Kerguelen Plateau (A3-2, 43.4 \pm 0.4 mmol m⁻² d⁻¹). In
- 344 the HNLC area (Station R-2), silica dissolution was higher than silica production, leading to a
- 345 negative $\int \rho Si_{net}(-1.78 \pm 0.02 \text{ mmol m}^{-2} \text{ d}^{-1})$.

346 3.4 Specific rates of production and dissolution

347 The specific Si uptake rate (VSi, d^{-1}) and dissolution rate (VDiss, d^{-1}) give the fraction of the 348 BSi pool produced or dissolved in one day as follows:

$$349 \quad VSi = \frac{\rho Si}{[BSi]} \tag{7}$$

350 and
$$VDiss = \frac{\rho Diss}{[BSi]}$$
 (8)

351 VSi is mainly impacted by nutrient and/or light limitation (Frank et al., 2000; Claquin et al.,

352 2002) and by the diatom community composition (Leynaert et al., 2004). During KEOPS-2,

353 VSi values (profiles not shown) presented the same decreasing trends with depth as Si uptake,

- 354 which is consistent with an impact of light limitation on silica production. Globally, relatively
- high "integrated specific Si-uptake rates" (JVSi, calculated by the averaged integrated ρ
- divided by the integrated BSi) prevailed for KEOPS-2 (≈ 0.1 to ≈ 0.3 d⁻¹; table 2). Such values
- are not different from those of nutrient-replete diatoms growing in the open ocean zone of the
- 358 Southern Ocean (Brzezinski et al., 2001). By contrast, the HNLC area showed a JVSi value
- below 0.1 d⁻¹, suggesting non-optimal conditions for the growth of diatoms and/or artifact of

siliceous detritus, which is important in other HNLC regions (e.g. Krause et al., 2010; Fripiatet al., 2011b).

 $\int VDiss varied around one order of magnitude during KEOPS-2 with low specific rates in$ productive stations (e.g. 0.03 d⁻¹ above the Plateau), and higher values in the HNLC area (up $to 0.15 d⁻¹). Interestingly, E-3 showed unexpected high <math>\int VDiss$ (0.12 d⁻¹; table 2).

365

366 **4 Discussion**

367 **4.1** Seasonality of the balance between silica production and dissolution

The D:P ratios integrated between the surface and the 1 % PAR attenuation depth ([D:]P; also 368 369 summarized in table 3) are presented in figure 5. At the HNLC reference station R-2, the 370 D:P value >1 indicates that the integrated dissolution rate exceeds the measured integrated 371 production rate (note that both fluxes were very low at R-2). This situation leads to a net loss 372 of biogenic silica by dissolution in the euphotic zone and suggests that a short development of 373 diatoms could have occurred before our sampling. This observation is in accordance with the 374 high barium excess measured between 200 and 400 m at R-2 (Jacquet et al, 2014), indicating 375 a high carbon mineralization activity in the mesopelagic zone which could be likely 376 associated to a surface production event prior sampling. High D values have already been 377 measured occasionally in the Southern Ocean during the summer bloom (review in Tréguer 378 and De La Rocha, 2013).

379 In the Kerguelen bloom area, $\int D$: $\int P$ ratios ranged from 0.09 (station A3-2) to 0.95 (station E-

380 3) and depended on the stage of the blooms. The $\int D: \int P$ ratios were relatively high at stations

381 visited in the beginning of the cruise, indicating that a significant fraction of silica was

recycled in the surface waters in early spring, and then decreased as the bloom took place.

383 The highest $\int D: \int P$ ratio occurred at station E-3. This station was characterized by a low BSi

384 stock (83.6 mmol Si m⁻²), a low integrated BSi production rate (10.5 mmol Si m⁻² d⁻¹

integrated over the euphotic layer; table 2), a dissolution rate close to the mean for all stations,

386 and high specific dissolution rate. This may evidence a higher relative proportion of detrital

387 silica free of organic matter at this station which could be due to stronger bacterial and/or

388 grazing activities inducing a top-down control on diatom growth. Without considering E-3,

 $\int D P$ ratios decreased progressively from E-1 to E-5 and showed low values at the most

productive stations E-4W, A3-2 and F-L. Here, ∫D:∫P ratios were similar to those measured in
nutrient-replete conditions such as productive upwelling regions (Brzezinski et al., 2003).

392 High D Pratios in winter and in early spring indicate that silica dissolution is sufficient to 393 sustain a large fraction of the low Si uptake rates observed during non-bloom conditions and 394 during the bloom onset, i.e. when primary production is still low. Indeed there is a temporal 395 decoupling between silica production and dissolution since the dissolution kinetic is slow. It is 396 only after diatom death and removal of their protecting organic coating by micro-organisms 397 that the silica frustules can dissolve (Kamatani, 1982; Bidle and Azam, 1999; Bidle et al., 2003). By contrast, the progressive decrease of the D: P values implies that the majority of 398 399 gross silica production is sustained by the silicic acid pool supplied from below (winter water) 400 as the bloom develops. This pool can be regarded as the "new" Si reservoir, similar to nitrate 401 for N. Thus, we observe a seasonal shift from Si uptake behaving mainly as a regenerated 402 production before the bloom onset, when silica production is still very low, and then behaving 403 more like a new production during bloom, when we observe higher Si uptake rates.

404 An opposite shift at the end of the productive period was suggested by Brzezinski et al. (2001) in the upwelling system of the Monterey Bay with D P ratios increasing following the bloom 405 development. In this case, higher [D:]P values were associated to an increase of the relative 406 407 proportion of detrital BSi in the water column. Similarly, the occurrence of an accumulation 408 of dissolving BSi in subsurface following productive periods inducing a net loss of BSi in late 409 summer ($\int D: \int P = 1.7$) was already identified in the Australian sector of the Southern Ocean (Fripiat et al., 2011b). Since KEOPS-2 took place at the start of the bloom and since there was 410 411 no silica dissolution rate measured from KEOPS-1, such increase of D: P ratio in the 412 Kerguelen area at the end of the blooming season has not been observed but will be discussed 413 in section 4.5.

- 414 Because silica dissolution profiles were not significantly different from each other between all
- 415 the KEOPS-2 bloom stations (Fig. 4b), it can be ruled out as a process explaining the
- 416 variability in D: P ratios. The observed decreasing trend of D: P ratios was actually mainly
- 417 driven by the increase of BSi production rates (from 3.09 ± 0.01 to 47.9 ± 0.4 mmol Si m⁻² d⁻
- 418 ¹) and by the accumulation of living diatoms with high specific Si uptake rates in the euphotic
- 419 layer (table 2).

The fraction of silica production supported by new silicic acid is estimated by 1-D. During 420 421 KEOPS-2 it ranged from -0.58 in the HNLC station, where we observed a net loss of biogenic 422 silica, to a maximum of 0.91 above the plateau, where maximum production rates were 423 recorded (A3-2, table 3). When plotting 1-D vs. gross silica production rate, Brzezinski et 424 al. (2003) found that 8 regional estimates of this parameter representing a large range of ocean 425 environments, fall along an hyperbolic curve and thus it might be possible to predict the 426 strength of the silicon pump in a system based on its mean silica production. To obtain a zero-427 intercept of the curve satisfying the assumption that when the fraction of silica production 428 supported by new silicic acid approaches 0, the production must also be 0, we have plotted the 429 1-[D:]P as a function of the net silica production (instead of the gross production in Brzezinski 430 et al., 2003). Since these two variables are not fully independent, the equation of the model 431 matching all the data follows a rectangular hyperbola (Fig. 6). This fitting has been obtained 432 on KEOPS-2, Brzezinski et al. (2003) and Fripiat et al. (2011b) data.

433 Using figure 6, we can identify several parameters characterizing the distribution of both KEOPS-2 stations and other oceanic regions. The 1-D: P_{max} is centered around 1 as it is not 434 possible to have more than 100 % of silica production supported by new-H₄SiO₄. The K_{oNet} 435 436 $(5.89 \pm 2.24 \text{ mmol m}^{-2} \text{ d}^{-1})$ represents the value of net silica production at which the system shifts from a regenerated to a new biogenic silica production (i.e., $1-\int D : \int P = 0.5$). In KEOPS-2 437 438 stations showing a net silica production below K_{oNet}, the development of diatoms is mainly 439 controlled by recycled sources of silicon, while above this value, the supply of new H₄SiO₄ is 440 the main source of nutrients for biogenic silica production. It is important to keep in mind that 441 this model is mainly governed by the dependency of its two variables and thus, we cannot use 442 it to make predictions about the silicon cycle functioning of stations where only one of the 443 two parameters was estimated (e.g. net silica production estimated from the change in BSi 444 concentrations). It however allows us to sort KEOPS-2 stations into specific groups and to 445 compare them with other oceanic regions. For example, figure 6 is remarkably helpful to 446 differentiate stations with low net production rates (x-axis) that have very variable fraction of 447 new Si production (y-axis). 448 KEOPS-2 stations follow the same trend as that of Brzezinski et al. (2003) and encompass

almost the full range of variability observed in very contrasting oceanic regions (e.g. HNLC,

450 oligotrophic, coastal upwelling, river plume). They can be sorted in 3 functional groups:

- The "low activity stations" group includes the HNLC reference station R-2 and station E-3 451 that showed a net loss of BSi with negative values of $\int \rho Si_{net}$ or close to 0 (Fig. 5; table 2). In 452 453 Fig. 6, the HNLC station falls in the negative part of the hyperbolic curve, close to stations 454 mainly characterized by detrital BSi dominance and where a release of silicon from dissolving 455 BSi takes place following a productive period (e.g. the late summer SAZ-Sense station P2 456 located in the Polar Front Zone; Fripiat et al., 2011b). Despite its low iron concentration, the 457 high [D:]P ratio observed at R-2 suggests that a short development of diatoms could have 458 occurred before our sampling in agreement with Jacquet et al. (2014) and Dehairs et al. (in 459 prep.). This kind of low diatom production in the HNLC area surrounding the Kerguelen 460 Plateau has already been suggested at the end of summer by Mosseri et al (2008). Since net 461 silica dissolution is not sustainable, the values measured at R-2 should necessarily represent 462 conditions that prevail on a short period of time. Production and dissolution rates are indeed 463 snapshot measurements over 24 h or 48 h. Although H₄SiO₄ concentrations were not limiting 464 in surface waters, E-3 was characterized by very low silica production that could be exclusively sustained by recycled silicic acid (1-D) = 0.05 and seemed to have approached 465 466 steady state conditions as siliceous biomass cannot increase in a system supported solely by 467 regenerated silicic acid (Brzezinski and Nelson, 1989). This situation could be the result of a 468 previous attempt to bloom that would have aborted due to the destabilization of the ML.

469 - The "starting-bloom" group is represented by station E-1 that has been visited in the 470 beginning of the KEOPS-2 cruise (early November). Although carbon incubation experiments 471 reveal that the bloom began to grow at this station (Cavagna et al., in prep.), low Si uptake 472 (Fig. 4a) and low net-silica production (Fig. 5) were still observed. A moderate $1-\int D: \int P$ ratio 473 (0.58) indicates that BSi production at E-1 is controlled both by new and regenerated sources 474 of H₄SiO₄.

475 - The "spring-bloom" group includes stations holding a strong capacity for BSi accumulation, 476 i.e. with low dissolution rates and high net silica production rates. Figure 4a allows us to 477 distinguish between stations from the lagrangian study E-4E and E-5, with only moderate surface ρ Si values (respectively $0.62 \pm 0.06 \mu$ mol l⁻¹ d⁻¹ and $0.57 \pm 0.06 \mu$ mol l⁻¹ d⁻¹) and 478 479 [D:]P ratio close to 0.3 (table 3); and stations A3-2, F-L and E-4W showing particularly high surface production rates (> 1 μ mol l⁻¹ d⁻¹) and [D:[P ratio close to 0.1 (table 3). Blooms with 480 such a low D: P ratio have the potential to accumulate a large fraction of BSi production 481 482 and/or export a large amount of BSi into the deep ocean (Quéguiner, 2013; Tréguer and De La 483 Rocha, 2013). Despite their location on both sides of the Polar Front and in different part of

484 the Kerguelen bloom, stations E-4W and F-L fall close to each other along the hyperbolic 485 curve (Fig. 6). Consequently, they should operate in a comparable way in term of silica 486 production dynamic which is quite similar to the average value of PFZ spring bloom 487 conditions measured by Brzezinski et al. (2001). So, even though complex physical settings 488 (Park et al., in prep.) are very different between E-4W (which is not part of PF with high 489 surface DSi concentration of 17 µmol l⁻¹) and F-L (with lower surface DSi concentration of 6 µmol l⁻¹) diatom production regime behave as typical PFZ stations. This was also observed 490 491 with carbon export and mesopelagic remineralization by Jacquet et al. (2014). Compared to F-492 L and E-4W, A3-2 is highly active in term of silica production and can be compared to the 493 Amazon river plume and coastal upwelling systems such as Monterey Bay or Peru. This 494 highlights once again the exceptional character of diatoms-dominated ecosystems sustained 495 by natural iron fertilization in the Southern Ocean.

496 **4.2** Decoupling between Si, C and N cycles in the Kerguelen area

497 In the Kerguelen area, the high NO_3^- concentrations in surface waters compared to H_4SiO_4 498 depletion observed annually at the end of the bloom period suggest a strong decoupling 499 between the seasonal consumption of these two nutrients (Mosseri et al., 2008). This situation 500 could be partly induced by differential recycling processes between Si and N strengthening 501 the silicon pump. Si is thus primarily exported to deeper water through sinking of biogenic 502 silica while PON is mostly recycled in the ML and used as nitrogen source for the 503 development of new phytoplankton organisms including diatoms. Since organic matter is 504 more quickly and efficiently remineralized compared to silica, this decoupling also occurs 505 between Si and C.

506 The strength of the silicon pump could be investigated by comparing the Si:C and Si:N 507 uptake-ratios. In this study, we use only the gross uptake ratios ($\int \rho Si: \int \rho N$ and $\int \rho Si: \int \rho C$), 508 calculated respectively as:

509
$$\int \rho \text{Si:} \int \rho \text{N} = \frac{\rho Si}{\rho(\text{NO}_3^- + NH_4^+)}$$
 (9)

510 and
$$\int \rho \text{Si:} \int \rho \text{C} = \frac{\rho Si}{gross \rho C}$$
 (10)

which reflect only the stoichiometry of phytoplankton nutrient uptake. We will not considernet uptake ratios that could be calculated but would be biased by the significant rates of

513 nitrification estimated at all KEOPS-2 stations (see Cavagna et al., in prep. and Dehairs et al.,

- 514 in prep.). Note that both $\int \rho Si: \int \rho C$ and $\int \rho Si: \int \rho N$ uptake ratios are underestimates of actual
- 515 diatom uptake ratios because of the simultaneous C and N uptake by non-siliceous organisms.
- 516 Diatoms growing in nutrient replete conditions present Si:C and Si:N elemental ratios around
- 517 0.13 (from 0.09 to 0.15) and 1 (from 0.8 to 1.2) respectively, with the variability of these
- 518 ratios depending on diatom species, size classes and growth rates (Brzezinski, 1985; Martin-
- 519 Jézéquel et al., 2000).
- 520 Si:C and Si:N uptake ratios are strongly impacted by co-limitations which alter growth rates
- 521 and in most cases increase silicification processes, and thus lead to higher uptake ratios
- 522 (Claquin et al., 2002; Leynaert et al., 2004; Bucciarelli et al., 2010). During the KEOPS-2
- study, $\int \rho Si: \int \rho C$ and $\int \rho Si: \int \rho N$ uptake ratios vary from 0.10 to 0.38 and from 0.32 to 1.51
- 524 respectively (table 3).
- 525 In our study, these variations could not be explained by limitation from macronutrient (such
- 526 as silicic acid) as already proposed by Nelson and Tréguer, (1992) and Quéguiner, (2001).
- 527 Indeed, KEOPS-2 took place at the beginning of the growth period (October-November) and
- 528 a bloom onset was observed above the Plateau (Blain et al., in prep.). It is thus not surprising
- 529 that macronutrient concentrations in the surface layer were not limiting for diatom growth (cf.
- 530 Cavagna et al., in prep. for N uptake). For silicic acid, kinetic experiments conducted during
- 531 the cruise at all sites demonstrated the lack of response of phytoplankton to H_4SiO_4
- 532 enrichment (data not shown). Indeed, at all stations, mixed layer silicic acid concentrations
- 533 were high (from 6.2 to 18.5 μ mol.l⁻¹; Fig. 3) preventing limitation of biogenic silica
- 534 production by H_4SiO_4 availability as V_{Si} at ambient DSi were always similar to V_{max} , the
- 535 maximum uptake rate achievable when Si is not limiting (data not shown).
- 536 Results from a previous cruise in the same area (austral spring 1995) already highlighted the 537 crucial role played by the light-mixing regime on the control of diatom growth in the nutrient 538 replete waters (Blain et al., 2001). Light-limitation also takes part in the decoupling between 539 Si, N and C cycles by decreasing the growth rate and consequently increasing Si:N and Si:C 540 uptake-ratios (Claquin et al., 2002). At all stations the pSi:pC uptake ratios increase slightly 541 with depth and reach a maximum at the bottom of the euphotic layer (data not shown), in 542 agreement with the fact that C assimilation is light-dependent through photosynthesis while 543 silicification processes mainly involves energy coming from respiration (Martin-Jézéquel et
- 544 al., 2000; Claquin et al., 2002). However, because Si uptakes reach very low values at the

base of the euphotic layer (Fig. 4a), our data suggest that BSi production rates were not fully

546 independent of light levels and that there was a close coupling between C and Si assimilation

- 547 processes (see Cavagna et al., in prep. for carbon uptake). This coupling of ρ Si with light was
- also observed in other regions as the Equatorial Pacific and the North Pacific Subtropical gyre
- 549 by Krause et al. (2011). Note however that this is not in contradiction with Claquin et al.
- 550 (2002) given the different time scales of the two studies.

551 Limitation by trace metals (especially iron) also alters the stoichiometry of phytoplankton 552 nutrient uptake and its elemental composition and, eventually, contributes to the decoupling 553 between Si, N and C cycles. From bottle enrichment experiments, it has been argued that 554 diatoms have higher Si:N uptake ratios under Fe stress (Takeda 1998; Hutchins and Bruland 555 1998; Franck et al., 2000). Interestingly, relatively high $\int \rho Si \left[\rho N \right]$ ratios were measured for the 556 Kerguelen spring bloom (table 3), with the highest value above the Plateau (1.5, station A3-2) 557 although this area was naturally Fe-enriched (Sarthou et al., in prep.; Queroue et al., in prep.). 558 In the KEOPS-2 productive stations, diatoms can take up more H_4SiO_4 compared to nitrogen 559 and carbon. Indeed, these organisms are known to store silicic acid in their vacuoles or linked 560 to other intracellular components (Martin-Jézéquel et al., 2000, Hildebrand, 2008), or could 561 be more silicified. By contrast, 3 stations showed low $\int \rho Si \left[\rho N \right]$ uptake ratios (0.44, 0.74 and 562 0.32 for R-2, E-3 and F-L respectively). Lower Si:N and Si:C integrated uptake ratios in these 563 areas might be partly due to changes in phytoplankton composition. By measuring 564 phytoplankton pigment composition in the HNLC station, Lasbleiz et al. (2014) estimated a 565 lower contribution of microphytoplankton due to a higher proportion of nanophytoplankton 566 organisms such as nanoflagellates. Thus, since the phytoplankton community at R may 567 contain a significant fraction of non-siliceous organisms, the C and N uptake ratios were not solely prescribed by diatoms and thus could explain the low $\int \rho Si \cdot \int \rho N$ uptake ratios observed. 568 569 However, the higher concentrations in fucoxanthin over the other pigments at all other 570 stations clearly evidence the dominance of large diatoms in the Kerguelen blooms (Lasbleiz et 571 al., 2014). The very low $\int \rho Si: \int \rho N$ uptake ratios estimated at F-L could not result from a 572 dominance of non-siliceous phytoplanktonic organisms but likely from different diatom 573 communities showing contrasted degree of silicification and adapted to the specific 574 hydrological and biogeochemical conditions occurring north of the Polar Front. Indeed, 575 previous studies in the Southern Ocean have already shown that diatom community 576 composition could explain more differences in silicification than physiological responses to 577 environmental factors such as iron concentration (Baines et al., 2010; Assmy et al., 2013).

578 Thus, at a seasonal scale and above the plateau, the combination of this increased Si uptake 579 and nitrogen regeneration processes including nitrification (see e.g. in Cavagna et al., in prep.; 580 Dehairs et al., in prep.) in the beginning of the bloom, and preferential recycling of organic 581 matter at the bloom offset helps to explain the depletion of most of the Si from surface layer observed between early spring (18.7 μ mol l⁻¹, averaged in the upper 80 m at station A3-2, 582 KEOPS-2) and the end of summer (1.9 μ mol l⁻¹, averaged in the upper 80 m at station A3, 583 KEOPS-1; Mosseri et al., 2008), while nitrate remains abundant (23 µmol l⁻¹, averaged in the 584 585 upper 80 m at station A3, KEOPS-1; Mosseri et al., 2008). Fe-replete diatom assemblages, 586 such as those found at A3-2 (Sarthou et al., in prep.), will deplete silicic acid from the water 587 column before nitrate. Such silicon pump above Kerguelen Plateau would then not be driven 588 solely by Fe-limitation contrary to incubation experiments from coastal upwelling systems 589 (Hutchins and Bruland, 1998). Indeed, Fe enrichments in bottle experiments fertilise on short 590 time scale (days) a diatom community that is not adapted to higher Fe levels. Here, by 591 comparing $\int \rho Si \left[\rho N \right]$ uptake ratios on different natural communities adapted to their specific 592 ambient Fe levels, our results suggest that natural Fe fertilisation might favour diatoms with 593 higher Si:N ratios. Above the Kerguelen Plateau, diatoms seem to maintain a relatively higher 594 degree of silicification until the demise of the bloom, since Mosseri et al. (2008) observed the same range of $\int \rho Si: \int \rho N$ at A3 (1.6 ± 0.5, n=3). 595

596 If confirmed in other naturally fertilized regions, these observations may have great 597 implications in our understanding of the past and future functioning of the Southern Ocean 598 and its role in the regulation of climate. Indeed, the silicon pump occurring in the ACC results 599 in an Antarctic Surface Water (AASW) relatively replete in NO₃⁻ but strongly depleted in 600 H₄SiO₄ as observed by Blain et al. (2007) and Mosseri et al. (2008) in the Kerguelen region. 601 This property is then exported toward lower latitudes by the Antarctic Intermediate Water 602 (AAIW) and Subantarctic Mode Water (SAMW) (Sarmiento et al., 2004), currently favouring 603 non siliceous organisms production in these regions. Thus, any change in Si:N uptake ratios 604 south of the region of AAIW and SAMW formation might in turn modify diatom productivity 605 at low latitude. Matsumoto et al. (2002) proposed that a Silicic Acid Leakage Hypothesis 606 (SALH) could explain the drop of atmospheric pCO₂ during glacial times and would be 607 mainly driven by changes in Si:N ratios of diatoms induced by an increase in iron supply to 608 the Southern Ocean. This hypothesis is based on in vitro and artificial Fe-enrichment 609 experiments showing lower Si:N uptake ratios for diatoms in non Fe-stressed conditions 610 (Takeda, 1998; Hutchins & Bruland, 1998), so that the increase of Fe-deposition in the

611 Southern Ocean during glacial times would drive AASW towards NO_3^- depletion instead of 612 the actual H₄SiO₄ depletion (Brzezinski et al., 2002).

613 Our results could indicate that glacial Fe fertilization may not have necessarily resulted in a 614 decrease in Si:N uptake ratios in AASW contrary to what has been proposed as an explanation 615 for the SALH (Brzezinski et al. 2002). Other processes leading to SALH could be invoked, 616 such as shifts in AAIW and SAMW formation rates (Crosta et al., 2007), variations in 617 phytoplankton assemblages (including relative contribution of non-siliceous organisms) or 618 changes in NO₃⁻ remineralization efficiency. Since this interpretation is based on several 619 incubations for several samplings (during spring and summer) in a single Fe-enriched station 620 (A3, n = 4) and only one station in the HNLC region (R-2), further investigations concerning 621 the seasonality of net silica production and Si:N uptake ratios in both naturally Fe-fertilized 622 and HNLC areas of the Southern Ocean are clearly needed to validate this observation.

623 **4.3** Seasonality and budget of the silicon cycle above the Kerguelen Plateau

624 In order to investigate the seasonal evolution of Si biogeochemical cycle in the Kerguelen 625 iron-fertilized bloom (station A3), we combined in figure 7 the KEOPS-1 and KEOPS-2 626 silicon fluxes measured using different isotopic approaches (stable and radiogenic isotope tracer incubations: ³⁰Si and ³²Si respectively, and, natural silicon isotopic composition, δ^{30} Si. 627 628 of both diatoms and seawater). Assuming that the whole water mass above the Kerguelen 629 Plateau may have been significantly ventilated with surface waters from the HNLC area at the 630 annual scale, Fripiat et al. (2011a) used the HNLC winter water (WW) characteristics to 631 represent the initial conditions of the winter H₄SiO₄ stock in the fertilized area surface layer (0-100 m). Because the biological activity (Si and C assimilation) took place only in the upper 632 633 80 m during KEOPS-2, Si fluxes, stocks and values estimated by Fripiat et al (2011a) 634 discussed in this section were recalculated from the surface to 80 m (instead of 100 m in 635 Fripiat et al. 2011a). The winter supply of DSi estimated from the seasonal H₄SiO₄ depletion during KEOPS-1 from the Upper Deep Circumpolar Water (UDCW) to the WW is around 2 636 mol m^{-2} , which is not significantly different from the 1.9 mol m^{-2} DSi stock we measured by 637 638 mid-October during KEOPS-2 (A3-1). This confirms that, as suggested by Fripiat et al 639 (2011a), the HNLC winter water is representative of the Si source with initial Si pool conditions prevailing before the bloom onset in the fertilized area. At this time, silica 640 production (which was not measured during this 1st visit) should be very low since only little 641 BSi accumulation was observed in the ML (79.1 mmol m^{-2}), and only 2 - 1.9 = 0.1 mol m^{-2} of 642

- 643 H₄SiO₄ was consumed in the surface water before our sampling compared to the estimated
- 644 WW initial stock. The system was likely exclusively driven by regenerated silica production
- 645 inducing a potentially high $\int D$: $\int P$ ratio (close to 1). Irradiance and mixed layer regime should
- be the more likely dominant factors controlling the bloom development for eukaryotes in
- 647 winter and in early spring (Boyd et al., 2001; Blain et al., 2013). Models have previously
- 648 reported that the interannual variability of mixed layer depth significantly affects both the date
- of the bloom onset and the maximum chlorophyll concentration in the region (Pondaven et al.,
- 1998). Then, because H₄SiO₄ and Fe were not at limiting concentrations in the surface layer
- 651 (Sarthou et al., in prep.), the light-mixing regime that occurred above the Plateau by mid-
- 652 October, should have been still unfavourable to diatom growth.
- 653 As the surface irradiance becomes more favourable with time, biogenic silica production progressively increases and reaches the highest net production value (46.8 mmol $m^{-2} d^{-1}$) 654 measured during our 2nd visit to A3 (mid-November). Between these 2 samplings (i.e. during 655 28 days), the H₄SiO₄ depletion in the ML (1.9 - 1.5 = 0.4 mol m⁻²) yields to an average $\int \rho Net$ 656 of 14.3 mmol m⁻² d⁻¹. Although it was located in different blooms of the Kerguelen region 657 658 with different diatom communities, this value is in good agreement with the net silica 659 production measured at station E-1 which we characterized as a "starting-bloom" dynamic. 660 Consequently, we can predict that simultaneously to the rise of silica production above the Plateau, the JD: P ratio should decrease toward values around 0.5 as measured at E-1, and that 661 662 the silica production could be controlled by both new and regenerated sources of H_4SiO_4 . In this situation, almost all the net BSi production is accumulated in the surface water: 348.5 663 mmol m^{-2} (BSi stock integrated over 80 m, this study) and very low export of biogenic silica 664 is allowed from the ML to the bottom layer: 4.7 mmol $m^{-2} d^{-1}$ (estimated from the difference 665 666 between net production calculated with DSi standing stocks and that calculated using BSi standing stocks, 14.3 and 9.6 mmol $m^{-2} d^{-1}$ respectively). These observations are in good 667 668 agreement with the low carbon export (4 % of the surface primary production) estimated 669 during the first visit at A3 by Jacquet et al. (2014). Indeed, such early spring bloom generally 670 starts by the development of lightly silicified diatoms with potentially low sedimentation rates 671 (Quéguiner, 2013).
- 672 The H_4SiO_4 stock measured during the second visit at A3 (1.5 mol m⁻²) can sustain the strong
- 673 net silica production rates measured there for 32 days. Using natural Si isotopic approach,
- Fripiat et al. (2011a) suggested that diatoms could receive in addition, at least 1.2 mol m^{-2} of
- 675 DSi coming from the WW during the productive period in the upper 80 m of the fertilized

676 area, allowing 26 supplementary days of growth for diatoms with the same high Si uptake 677 rate. Taking into account the sum of such winter and summer vertical Si supply, the high 678 productive period as measured in A3-2 can be maintained during 86 days. Remarkably, this is 679 almost similar to the estimation by Mongin et al. (2008) based on satellite products of 85 days 680 of blooming over the Kerguelen plateau. Using a box-model approach, De Brauwere et al. 681 (2012) suggested that the bloom could also persist over the same duration without considering 682 this summer Si supply from deep-waters. This appears unlikely because the high net silica 683 production we measured at A3-2 could not be sustained for more than 32 days. Even if such 684 high net silica production should probably not be representative of the Si uptake by diatoms 685 over the Plateau during the rest of bloom period, an additional source of Si is needed. The 686 very good accordance of our results with the bloom duration from Mongin et al. (2008) 687 suggests that this summer H₄SiO₄ input in the ML is realistic and could sustain a significant 688 part of the phytoplankton growth above the Kerguelen Plateau. Indeed, high internal wave 689 activity above the plateau is assumed to be a major process for vertical dissolved iron (and 690 other nutrients) supply on the upper waters (Park et al., 2008; Blain et al., 2008).

691 These vertical fluxes allow a 2-month period of highly active blooming, with a system 692 controlled exclusively by new sources of H_4SiO_4 ([D:]P = 0.09, A3-2, table 3), inducing the strong DSi depletion observed in January (DSi stocks at 0.2 mol m⁻², KEOPS-1; Mosseri et 693 al., 2008) and the high BSi accumulation in the ML (348.5 mmol m^{-2} , KEOPS-2; this study) 694 695 which could be exported at the end of summer when the water column stratification becomes weaker. The gross- $\int \rho Si$ measured here is in the upper range of published values in the 696 697 Southern Ocean and the specific Si uptake rate was relatively high (respectively 47.9 mmol m⁻ 2 d⁻¹ and 0.28 d⁻¹ in the euphotic layer; table 2). This could indicate that diatoms have already 698 699 reached their maximum BSi production rate, and that our second visit to A3 could represent 700 the maximum of the bloom dynamic above the Plateau. This is quite consistent with the date 701 of the bloom peak estimated in early December both by modeling and satellite approaches 702 (Mongin et al., 2008 and de Brauwere et al., 2012). The H₄SiO₄ standing stock in the ML would then be depleted by mid-January (0.2 mol m⁻², KEOPS-1; Mosseri et al., 2008). Then 703 the bloom would shift toward a steady state dynamic, almost entirely controlled by 704 regenerated Si with decreasing gross silica production down to 10.7 mmol m⁻² d⁻¹, KEOPS-1 705 (Mosseri et al., 2008) and where H₄SiO₄ concentration in the ML becomes limiting for diatom 706 707 growth.

708 This progression of Si limitation could be associated to a change in the phytoplankton 709 community as observed between the different visits at A3 during KEOPS-1 (Mosseri et al., 710 2008; Armand et al., 2008), and could be related to the selection of species with higher 711 affinities with silicic acid resulting in a better ability to grow at low H₄SiO₄ concentrations. 712 Such change in the community structure in response to physical and biological forcing was 713 also proposed in a conceptual scheme by Quéguiner (2013). In spring, diatoms presenting 714 high growth rates and low degree of silicification dominate the bloom which development is 715 mainly controlled by new nutrients sources. This diatom assemblage will be soon affected by 716 the availability of both silicic acid and iron, and will change for a population showing lower 717 growth rates exclusively sustained by regenerated sources of H₄SiO₄. This second 718 assemblage, beginning to dominate in January, could thus persist at steady state until May, when PAR decreases below the threshold of 1 mol photon $m^{-2} d^{-1}$ (Blain et al., 2013) in waters 719 720 that are quite depleted in H₄SiO₄, as it is composed by small diatoms with high affinities for 721 silicic acid (Mosseri et al., 2008) and a deep silica maximum (DSM) characterized by strongly 722 silicified large-sized diatoms growing at the base of the ML in the nutrient gradient 723 (Quéguiner, 2013). Under such conditions, a very low net silica production and a D: P ratio 724 close to 1 are expected. Unfortunately no silica dissolution measurements are available from 725 KEOPS-1. However, we can try to estimate the net production from the difference between the KEOPS-1 average gross silica production (10.7 mmol $m^{-2} d^{-1}$; Mosseri et al., 2008) and 726 727 average silica dissolution measured at all stations south of the Polar Front during KEOPS-2 $(10.2 \text{ mmol m}^{-2} \text{ d}^{-1})$. The choice of a constant silica dissolution rate throughout the productive 728 729 season may seem surprising at first sight. However, there are several reasons to support this 730 hypothesis: (i) Brzezinski et al. (2001) show that the seasonal variability of silica dissolution rates in the Southern Ocean is very low (6.7 mmol m⁻² d⁻¹ in October/November to 6.6 mmol 731 732 $m^{-2} d^{-1}$ in February/March); (ii) JoDiss measured close to the Polar Front by Fripiat et al. (2011b) in late summer were very close to our spring dissolution values (4.9 to 6.6 mmol m^{-2} 733 734 d^{-1}) (iii) As best seen on Fig. 4b, all KEOPS-2 stations except the HNLC one (R-2) have 735 similar silica dissolution profiles. Thus it is reasonable to assume that the net silica production 736 estimated above the plateau in late summer should be around 0.5 mmol $m^{-2} d^{-1}$.

737 The BSi standing stock observed in the upper layer in late summer (275.3 mmol m⁻², KEOPS-

1; Mosseri et al., 2008) is lower than that measured in spring (348.5 mmol m^{-2}). This could

indicate that a part of the BSi that was produced before reaching a silica production in steady

state (i.e. sustained almost entirely by regenerating production) was not accumulated in the

surface ML at the end of the productive period. In fact, a part of the summer production can
be stocked in the form a DSM as measured in late summer above the plateau (550.1 mmol m⁻²

- between 100 and 200 m; Mosseri et al., 2008) which was not yet present at KEOPS-2. This
- subsurface biogenic silica accumulation mainly results from the combination of sedimentation
- 745 of living but inactive cells and the occurrence of phytoplankton populations living at depth
- 746 (Uitz et al., 2009; Fripiat et al., 2011a). The part of the summer net silica production which is
- not accumulated below the ML should thus be exported to deeper waters as the seasonal
- stratification breaks down with the intensification of vertical mixing. In term of carbon export,
- this flux could represent 14 to 31 % of the surface primary production (KEOPS-1, station A3;
- 750 Jacquet et al., 2008). As proposed by Quéguiner (2013), a massive export of biogenic silica
- 751 and organic matter (58 days x 46.8 mmol $m^{-2} d^{-1} (275.3 + 550 \text{ mmol } m^{-2}) = 2.2 \text{ mol Si } m^{-2})$
- should occur and could then represent the major annual event of the silicon and biological
- carbon pumps.

754 Although our budget of the silicon biogeochemical cycle above the Kerguelen Plateau is 755 based on different silicon isotopic approaches and is sustained by silicon stocks and fluxes 756 coming from different years, it matches very well with all the previous individual findings. 757 The seasonal variations of [D:]P ratio are in accordance with those observed by Brzezinski et 758 al. (2001) across the Polar Front Zone and are quite well represented by the recent model of 759 Coffineau et al. (2013), who estimate a D:P ratio ranging from 0.64 in winter to 0.19 during 760 the spring bloom. The good accordance of our approach with outcomes from different studies 761 also highlights that, to fully characterize the silicon cycle in a region of interest, we need to 762 measure both silica production and dissolution rates. Indeed, taking only into account the 763 gross silica production in such a synthesis exercise (i.e. without measuring silica dissolution 764 as it is the case in most studies) could lead to misinterpretations of the silicon pump 765 functioning. For instance, we would not be able to identify shifts between "new" and 766 "regenerated" silica production neither to accurately calculate the real bloom duration without 767 considering silica recycling. Without taking into account dissolution, silica production would have been overestimated by 21 % in the mixed layer (58.2 mmol m⁻² d⁻¹ compared to the 47.9 768 mmol $m^{-2} d^{-1}$ of real net silica production), and the bloom duration computation would have 769 yielded 74 days, which is not consistent with the 85 days of KEOPS-1 bloom duration 770 771 observed by Mongin et al. (2008).

773 **5 Conclusions**

Our study addressed the seasonal evolution of the efficiency of the silicon pump and of the

biogenic silica fluxes in the mixed layer under different naturally iron-fertilized bloom

conditions around the Kerguelen region. Integrated Si uptake rates were among the highest

reported so far in the Southern Ocean. They varied from $3.09 \pm 0.01 \text{ mmol m}^{-2} \text{ d}^{-1}$ in the

HNLC area (R-2) to 47.9 ± 0.4 mmol m⁻² d⁻¹ above the plateau (A3-2) and seemed to be

strongly coupled with C uptake over depth. Indeed, C and Si assimilation were very low

below the euphotic layer indicating the occurrence of a subsurface accumulation of living but

781 inactive diatoms. Although significant, silica dissolution rates were generally much lower

than production rates and did not vary between bloom stations nor over depth.

783 We observed a shift from a BSi production regime based on the regeneration of H_4SiO_4 784 during the early stages of bloom onset (with an averaged D: P ratio of 0.98) to a regime 785 based on new production during the bloom development (with an averaged D: P ratio of 786 0.18). This change switched on an active silicon pump which led to the decoupling between Si 787 and N cycles as well as a strong H₄SiO₄ depletion of surface water by late summer, with 788 significant implications for global biogeochemical properties. Indeed, the system 789 progressively shifted toward a stronger silicon pump as Si uptake rates increased and nitrogen 790 became preferentially remineralized when the bloom was well established. This led ultimately 791 to a strong Si limitation and drove the system toward a regenerated silica production regime 792 which allowed the persistence of the bloom in a steady state despite the low concentrations of 793 silicic acid concentrations. Our results confirm and complete the concept of a seasonal 794 transition from a diatom new production to a diatom regenerated production already proposed 795 in the Antarctic Zone by Brzezinski et al. (2003).

796 Moreover, in opposition to previous artificial Fe-enrichment bottle experiments outcomes, 797 Si:N and Si:C uptake ratios during KEOPS-2 were not higher in the HNLC area compared to 798 the fertilized region. This observation could have great implications on our understanding on 799 processes involved in setting atmospheric pCO₂ during glacial-interglacial transitions. Our 800 results suggest that the increase of low latitude diatom production observed during glacial 801 periods should not be controlled primarily by a shift in the nutrient uptake stoichiometry of 802 Antarctic diatoms induced by an enhanced iron supply, as proposed in the silicic acid leakage 803 hypothesis (Matsumoto et al., 2002; Brzezinski et al., 2002), but further investigations are 804 clearly needed to validate this idea.

805 The combination of the results from the two KEOPS cruises (early spring and late summer) 806 and of different isotopic approaches, allowed the first seasonal estimate of a closed silicon 807 biogeochemical budget in the iron-fertilized area above the Kerguelen Plateau based on direct 808 measurements. Our estimates emphasize the interest of combining different tracers and 809 methods with different sensitivities to physical and biological processes to better constrain 810 and quantify all the processes simultaneously. The major outcome of this seasonal budget is that the winter and summer silicon supplies to the mixed layer $(3.2 \text{ mol m}^{-2} \text{ y}^{-1})$ seem to be 811 well balanced by the combination of biogenic silica accumulation (both in the upper layer and 812 813 in the winter waters) and late summer BSi export (respectively $0.3 + 0.6 + 2.2 = 3.1 \text{ mol m}^{-2}$ y^{-1}). This confirms the occurrence of a significant summer Si supply from Winter Water as 814 815 suggested by Fripiat et al. (2011a) sustaining the diatom bloom over the Kerguelen Plateau.

816 Finally, a striking feature of this study is that naturally iron fertilized areas of the Southern 817 Ocean, like the Kerguelen Plateau, could sustain a biogenic silica production regime similar to 818 those observed in coastal upwelling systems or in large river plume. This highlights the 819 exceptional character of diatoms-dominated ecosystems associated to natural iron fertilization 820 in the Southern Ocean, and their significant role in the global Si biogeochemical cycle. Even 821 if the outcomes of this budget are consistent with previous measurements, large uncertainties 822 remain about the seasonal evolution of dissolution rates at the end of the productive period. 823 Indeed, in order to fully characterize the Si-biogeochemical cycle in a region of interest, it is 824 recommended to measure both BSi production and dissolution rates. In combination, the natural silicon isotopic composition (δ^{30} Si) of diatoms and seawater represents a powerful tool 825 826 for identifying silicon sources and silica production over larger temporal and spatial scales 827 (Fripiat et al., 2011a; Tréguer and De la Rocha, 2013), and will be also examined during 828 KEOPS-2. The combination of several isotopic approaches, as well as modeling exercises 829 (such as in Pondaven et al., 1998; De Brauwere et al., 2012; Coffineau et al., 2013), by 830 allowing to better constrain and quantify different physical and biogeochemical processes 831 simultaneously, would strongly improve our understanding of the regional Si biogeochemical 832 cycle and its implications in the global ocean biogeochemistry.

833

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847 **References**

- 848 Assmy P., Smetacek V., Montresor M., Klaas C., Henjes J., Strass V., Arrieta J., Bathmann
- 849 U., Berg G., Breitbarth E., Cisewski B., Friedrichs L., Fuchs N., Herndl G., Jansen S.,
- 850 Krägefsky S., Latasa M., Peeken I., Röttgers R., Scharek R., Schüller S., Steigenberger S.,
- 851 Webb A., Wolf-Gladrow D.: Thick-shelled, grazer-protected diatoms decouple ocean carbon
- and silicon cycles in the iron-limited Antarctic Circumpolar Current. Proceedings of the
- National Academy of Sciences of the United States of America. 110(51):20633–20638, 2013.
- 854 Armand L., Cornet-Barthaux V., Mosseri J., Quéguiner B.: Late summer diatom biomass and
- community structure on and around the naturally iron-fertilised Kerguelen Plateau in the
- 856 Southern Ocean. Deep-Sea Research II. 55:653-676, 2008.
- 857 Baines S., Twining B., Brzezinski M., Nelson D., Fisher N.: Causes and biogeochemical
- 858 implications of regional differences in silicification of marine diatoms. Global
- 859 Biogeochemical Cycles. 24:GB4031. Doi10.1029/2010GB003856, 2010.
- 860 Beucher C., Tréguer P., Hapette A. M., Corvaisier R.: Intense summer Si-recycling in the
- surface Southern Ocean. Geophysical research letters. 31:L09305.
- 862 doi:10.1029/2003GL018998, 2004.
- 863 Bidle K. D., Azam F.: Accelerated dissolution of diatom silica by marine bacterial
- 864 assemblages. Nature. 397:508-512, 1999.

- Bidle K. D., Brzezinski M., Long R., Jones J., Azam F.: Diminished efficiency in the oceanic
 silica pump caused by bacteria-mediated silica dissolution. Limnology and Oceanography.
 48(5):1855-1868, 2003.
- 868 Blain S., Tréguer P., Belviso S., Bucciarelli E., Denis M., Desabre S., Fiala M., Martin-
- 869 Jézéquel V., Le Fèvre J., Mayzaud P., Marty J.-C., Razouls S.: A biogeochemical study of the
- 870 island mass effect in the context of the iron hypothesis: Kerguelen Islands, Southern Ocean.
- 871 Deep-Sea Research I. 48:163-187, 2001.
- 872 Blain S., Quéguiner B., Armand L., Belviso S., Bombled B., Bopp L., Bowie A., Brunet C.,
- 873 Brussaard C., Carlotti F., Christaki U., Cordière A., Durand I., Ebersbach F., Fuda J.-L.,
- 874 Garcia N., Gerringa L., Griffiths B., Guigue C., Guillerm C., Jacquet S., Jeandel C., Laan P.,
- 875 Lefèvre D., Lo Monaco C., Malits A., Mosseri J., Obernosterer I., Park Y.-H., Picheral M.,
- 876 Pondaven P., Remenyi T., Sandroni V., Sarthou G., Savoye N., Scouarnec L., Souhaut M.,
- 877 Thuiller D., Timmermans K., Trull T., Uitz J., Van Beek P., Velhuis M., Vincent D., Viollier
- 878 E., Vong L., Wagener T.: Effect of natural iron fertilization on carbon sequestration in the
- 879 Southern Ocean. Nature. 446:1070-1075, 2007.
- 880 Boyd P., Crossley A., diTullio G., Friffiths F., Hutchins D., Queguiner B., Sedwick P., Trull
- T.: Control of phytoplankton growth by iron supply and irradiance in the subantarctic
- 882 Southern Ocean: Experimental results from the SAZ Project. Journal of Geaophysical
- 883 Research. 106:31573-31583, 2007.
- Blain S., Quéguiner B., Trull T.: The natural iron fertilization experiment KEOPS (KErguelen
 Ocean and Plateau compared Study): An overview. Deep-Sea Research II. 55:559-565, 2008.
- 886 Blain S., Renaut S., Xing X., Claustre H., Guinet C.: Instrumented elephant seals reveal the
- seasonality in chloropyll and light-mixing regime in the iron fertilized Southern Ocean.
- 888 Geophysical Research Letters. 40:1-5, 2013.
- 889 Blain S.: KEOPS2: implementation and overview. In preparation for Biogeosciences.
- 890 Boyd P., Crossley A., DiTullio G., Griffiths F., Hutchins D., Quéguiner B., Sedwick ., Trull
- 891 T.: Control of phytoplankton growth by iron supply and irradiance in the subantarctic
- 892 Southern Ocean: Experimental results from the SAZ Project. Journal of Geophysical
- 893 Research. 106(C12):31573-31583, 2001.

- 894 Boyd P., Jickells T., Law C., Blain S., Boyle E., Buesseler K., Coale K., Kullen J., De Baar
- 895 H., Follows M., Harvey M., Lancelot C., Levasseur M., Owens N., Pollard R., Rivkin R.,
- 896 Sarmiento J., Schoemann V., Smetacek V., Takeda S., Tsuda A., Turner S., Watson A.:
- 897 Mesoscale Iron Enrichment Experiments 1993-2005: Synthesis and Future Directions.
- 898 Science. 315:612-617, 2007.
- 899 Brzezinski M.: The Si:C:N ratio of marine diatoms: interspecific variability and the effect of
- some environmental variables. Journal of Phycology. 21:347-357, 1985.
- 901 Brzezinski M., Nelson D.: Seasonal changes in the silicon cycle within a Gulf Stream warm-
- 902 core ring. Deep-Sea Research. 36(7):1009-1030, 1989.
- 903 Brzezinski M., Nelson D., Franck V., Sigmon D.: Silicon dynamics within an intense open-
- ocean diatom bloom in the Pacific sector of the Southern Ocean. Deep-Sea Research II.
- 905 48:3997-4018, 2001.
- 906 Brzezinski M. A., Pride C. J., Frank V. M., Sigman D. M., Sarmiento J. L., Matsumoto K.,
- 907 Gruber N., Rau G. H., Coale K. H.: A switch from Si(OH)4 to NO3- depletion on the glacial
- Southern Ocean. Geophysical Research Letters. 29(12), doi:10.1029/2001GL014349, 2002.
- 909 Brzezinski M., Jones J., Bidle K., Azam F.: The balance between silica production and silica
- 910 dissolution in the sea: Insights from Monterey Bay, California, applied to the global data set.
- 911 Limnology and Oceanography. 48(5):1846-1854, 2003.
- 912 Bucciarelli E., Pondaven P., Sarthou G.: Effects of an iron-light co-limitation on the elemental
- 913 composition (Si, C, N) of the marine diatoms Thalassiosira oceanica and Ditylum brightwellii.
- 914 Biogeosciences. 7:657-669, 2010.
- 915 Buesseler K., Ball L., Andrews J., Cochran J., Hirschberg D., Bacon M., Fleer A., Brzezinski
- 916 M.: Upper ocean export of particulate organic carbon and biogenic silica in the Southern
- 917 Ocean along 170°W. Deep-Sea Research II. 48:4275-4297, 2001.
- 918 Cavagna A. J. Blain S., Cardinal D., Closset I., Dehairs F., Fernandez C., Flores-Leive L.,
- 919 Lasbleiz M., Lefèvre D., Leblanc K., Quéguiner B.: Biological productivity regime and in-situ
- 920 methods comparison around the Kerguelen Island in the Southern Ocean. KEOPS 2 primary
- and community producers regime using various uptake rates and their stoichiometric ratios. In
- 922 preparation for Biogeosciences.

- 923 Claquin P., Martin-Jézéquel V., Kromkamp J. C., Veldhuis M. J. W., Kraay G. W.:
- 924 Uncoupling of silicon compared to carbon and nitrogen metabolisms, and role of the cell
- 925 cycle, in continuous cultures of Thalassiosira pseudonana (Bacillariophyceae) under light,
- nitrogen and phosphorus control. Journal of Phycology. 38:922-930, 2002.
- 927 Coffineau N., De La Rocha C., Pondaven P.: Exploring interacting influences on the silicon
- 928 isotopic composition of the surface ocean: a case study from the Kerguelen Plateau.
- 929 Biogeosciences Discussion. 10:11405-11446, 2013.
- 930 Corvaisier R., Tréguer P., Beucher C., Elskens M.: Determination of the rate of production
- and dissolution of biosilica in marine waters by thermal ionisation mass spectrometry.
- 932 Analytica Chimica Acta. 534:149-155, 2005.
- 933 Crosta X., Beucher C., Pahnke K., Brzezinski M.: Silicic acid leakage from the Southern
- Ocean: Opposing effects of nutrient uptake and oceanic circulation. Geophysical Research
- 935 Letters. 34, L13601, doi:10.1029/2006GL029083, 2007.
- 936 De Baar H., Boyd P., Coale K., Landry M., Tsuda A., Assmy P., Bakker D., Bozec Y., Barber
- 937 R., Brzezinski M., Buesseler K., Boye M., Croot P., Gervais F., Gorbunov M., Harrison P.,
- Hiscock W., LaanP., Lancelot C., Law C., Levasseur M., Marchetti A., Millero F., Nishioka
- 939 J., Nojiri Y., van Oijen T., Riebesell U., Rikenberg M., Saito H., Takeda S., Timmermans K.,
- 940 Veldhuis M., Waite A., Wong C.-S.: Synthesis of iron fertilization experiments: From the Iron
- 941 Age in the Age of Enlightenment. Journal of Geophysical Research. 110:C09S16,
- 942 doi:10.1029/2004JC002601, 2005.
- 943 De Brauwere A., De Ridder F., Elskens M., Schoukens J., Pintelon R., Baeyens W.: Refined
- 944 parameter and uncertainty estimation when both variables are subject to error. Case study:
- 945 estimation of Si consumption and regeneration rates in a marine environment. Journal of
- 946 Marine Systems. 55:205-221, 2005.
- 947 De Brauwere A., Fripiat F., Cardinal D., Cavagna A.-J., De Ridder F., André L., Elskens M.:
- 948 Isotopic model of oceanic silicon cycling: the Kerguelen Plateau case study. Deep-Sea
- 949 Research I. 70:42-59, 2012.
- 950 Dehairs F., Trull T., Fernandez C., Davies D., Cavagna A. J., Planchon F., Fripiat F.: Nitrate
- 951 isotopic composition in the Kerguelen area (Southern Ocean) during KEOPS 2. In preparation
- 952 for Biogeosciences.

- 953 Dugdale R., Wilkerson F., Minas H.: The role of a silicate pump in driving new production.
- 954 Deep-Sea Research I. 5(42):697-719, 1995.
- 955 Elskens M., de Brauwere A., Beucher C., Corvaisier R., Savoye N., Tréguer P., Baeyens W.:
- 956 Statistical process control in assessing production and dissolution rates of biogenic silica in
- marine environments. Marine Chemistry. 106:272-286, 2007.
- 958 Franck V., Brzezinski M., Coale K., Nelson D.: Iron and silicic acid concentrations regulate
- 959 Si uptake north and south of the Polar Frontal Zone in the Pacific Sector of the Southern
- 960 Ocean. Deep-Sea Research II. 47:3315-3338, 2000.
- 961 Fripiat F., Corvaisier R., Navez J., Elskens M., Schoemann V., Leblanc K., André L.,
- 962 Cardinal D.: Measuring production-dissolution rates of marine biogenic silica by ³⁰Si-isotope
- 963 dilution using a high-resolution sector field inductively coupled plasma mass spectrometer.
- Limnology and Oceanography: Methods. 7:470-478, 2009.
- 965 Fripiat F.: Isotopic approaches of the silicon cycle: The Southern Ocean case study (Ph. D.
 966 dissertation). Université Libre de Bruxelles, Brussels. 266p. 2010.
- 967 Fripiat F., Cavagna A. J., Savoye N., Dehairs F., André L., Cardinal D.: Isotopic constraints
- 968 on the Si-biogeochemical cycle of the Antarctic Zone in the Kerguelen area (KEOPS). Marine
 969 Chemistry. 123:11-22, 2011a.
- 970 Fripiat F., Leblanc K., Elskens M., Cavagna A. J., Armand L., André L., Dehairs F., Cardinal
- 971 D.: Efficient silicon recycling in summer in both the Polar Frontal and Subantarctic Zones of
- the Southern Ocean. Marine Ecology Progress Series. 435:47-61, 2011b.
- Hildebrand M.: Diatoms, Biomineralization Processes, and Genomics. Chemical Review.
 108(11):4855-4874, 2008.
- 975 Hutchins D., Bruland K.: Iron-limited diatom growth and Si:N uptake ratios in a coastal
- 976 upwelling regime. Nature. 393:561-564, 1998.
- 977 Jacquet S. H. M., Dehairs F., Savoye N., Obernosterer I., Christaki U., Monnin C., Cardinal
- 978 D.: Mesopelagic organic carbon remineralization in the Kerguelen Plateau region tracked by
- biogenic particulate Ba. Deep-Sea Research II. 55:868-879, 2008.

- Jacquet S. H. M., Dehairs F., Cavagna A. J., Planchon F., Monin L., André L., Closset I.,
- 981 Cardinal D.: Early season mesopelagic carbon remineralization and transfer efficiency in the
- naturally iron-fertilized Kerguenlen area. Biogeosciences Discussion. 11:9035-9069, 2014.
- 983 Kamatani A.: Dissolution rates of silica from diatoms decomposing at various temperatures.
- 984 Marine Biology. 68:91-96, 1982.
- 985 Karl D., Tien G.: MAGIC: A sensitive and precise method for measuring dissolved
- 986 phosphorus in aquatic environment. Limnology and Oceanography. 37(1):105-116, 1992.
- 987 Krause J. W., Nelson D. M., Brzezinski M. A.: Biogenic silica production and the diatom
- 988 contribution to primary production and nitrate uptake in the eastern equatorial Pacific Ocean.
- 989 Deep-Sea Research II. 58:434-448, 2011
- 990 Lasbleiz M., Leblanc K., Blain S., Ras J., Cornet-Barthaux V., Hélias Nunige S., Quéguiner
- 991 B.: Pigments, elemental composition (C, N, P, Si) and stoichiometry of particulate matter, in
- the naturally iron fertilized region of Kerguelen in the Southern Ocean. Biogeosciences
- 993 Discussion. 11:8259-8324.
- 994 Leynaert A., Bucciarelli E., Claquin P., Dugdale R. C., Martin-Jézéquel V., Pondaven P.,
- Ragueneau O.: Effect of iron deficiency on diatom cell size and silicic acid uptake kinetics.
 Limnology and Oceanography. 49(4):1134-1143, 2004.
- 997 Matsumoto K., Sarmiento J. L., Brzezinski M. A.: Silicic acid leakage from the Southern
- 998 Ocean: A possible explanation for glacial atmospheric pCO2. Global Biogeochemical Cycles.
- 999 16(3), doi:10.1029/2001GB001442, 2002
- 1000 Martin-Jézéquel V., Hildebrand M., Brzezinski M.: Silicon metabolism in diatoms:
- 1001 implications for growth. Journal of Phycology. 36:821-840, 2000.
- 1002 Mongin M., Molina E., Trull T.: Seasonality and scale of the Kerguelen plateau
- 1003 phytoplankton bloom: A remote sensing and modeling analysis of the influence of natural iron
- 1004 fertilization in the Southern Ocean. Deep-Sea Research II. 55:880-892, 2008.
- 1005 Mosseri J., Quéguiner B., Armand L., Cornet-Barthaux V.: Impact of iron on silicon
- 1006 utilization by diatoms in the Southern Ocean: A case study of Si/N cycle decoupling in a
- 1007 naturally iron-enriched area. Deep-Sea Research II. 55:810-819, 2008.

- 1008 Nelson D. and Goering J. J.: A Stable Isotope Tracer Method to Measure Silicic Acid Uptake
- 1009 by Marine Phytoplankton. Analytical Biogeochemistry. 78:139-147, 1977a.
- 1010 Nelson D. and Goering J. J.: Near-surface silica dissolution in the upwelling region off
- 1011 northwest Africa. Deep –Sea Research. 24:65-73, 1977b.
- 1012 Nelson D., Tréguer P.: Role of silicon as a limiting nutrient to Antarctic diatoms: evidence
- 1013 from kinetic studies in the Ross Sea ice-edge zone. Marine ecology progress series. 80:255-1014 264, 1992.
- 1015 Park Y.-H., Charriaud E., Ruiz Pino D., Jeandel C.: Seasonal and interannual variability of the
- 1016 mixed layer properties and steric height at station KERFIX, southwest of Kerguelen. Journal
- 1017 of Marine Systems. 17:571-586, 1998.
- 1018 Park Y.-H., Roquet F., Durand I., Fuda J.-L.: Large-scale circulation over and around the
- 1019 Northern Kerguelen Plateau. Deep-Sea Research II. 55:566-581, 2008.
- 1020 Park Y.-H.: Water masses and circulation in the Polar Front region east of the Kerguelen
- 1021 Islands. In preparation for Biogeosciences.
- Pollard R., Lucas M., Read J.: Physical controls on biogeochemical zonation in the Southern
 Ocean. Deep-Sea Research II. 49:3289-3305, 2002.
- 1024 Pondaven P., Fravalo C., Ruiz-PinoD., Tréguer P., Quéguiner B., Jeandel C.: Modelling the
- 1025 silica pump in the Permanently Open Ocean Zone of the Southern Ocean. Journal of Marine
- 1026 Systems. 17:587-619, 1998.
- Pondaven P., Ragueneau O., Tréguer P., Hauvespre A., Dézileau L., Reyss J.L.: Resolving the
 "opal paradox" in the Southern Ocean. Nature. 405(6783):168-172, 2000.
- 1029 Quéguiner B.: Biogenic silica production in the Australian sector of the Subantarctic Zone of
- 1030 the Southern Ocean in late summer 1998. Journal of Geophysical Research. 106(C12):31627-
- 1031 31636, 2001.
- 1032 Quéguiner B.: Iron fertilization and the structure of planktonic communities in high nutrient
- 1033 regions of the Southern Ocean. Deep-Sea Research II. 90:43-54, 2013.

- 1034 Quéguiner B., Brzezinski M.: Biogenic silica production rates and particulate organic matter
- distribution in the Atlantic sector of the Southern Ocean during austral spring 1992. Deep-SeaResearch II. 49:1765-1786, 2002.
- 1037 Quéroué F.: Dissolved iron in the vicinity of the Kerguelen plateau (KEOPS-2 experiment). In1038 preparation for Biogeosciences.
- 1039 Ragueneau O., Treguer P., Leynaert A., Anderson R. F., Brzezinski M. A., DeMaster D. J.,
- 1040 Dugdale R. C., Dymont J., Fisher G., François R., Heinze C., Maier-Reimer E., Martin-
- 1041 Jézéquel V., Nelson D. M., Quéguiner B.: A review of the Si cycle in the modern ocean:
- 1042 recent progress and missing gaps in the application of biogenic opal as a paleoproductivity
- 1043 proxy. Global Planet Change. 26:317-365, 2000.
- 1044 Ragueneau O., Savoye N., Del Amo Y., Cotten J., Tardiveau B., Leynaert A.: A new method
- 1045 for the measurement of biogenic silica in suspended matter of coastal waters: using Si:Al
- 1046 ratios to correct for the mineral interference. Continental Shelf Research. 25:697-710, 2005.
- 1047 Reynolds B., Frank M., Halliday A.: Silicon isotope fractionation during nutrient utilization in
 1048 the North Pacific. Earth and Planetary Science Letters. 244:431-443, 2006.
- 1049 Roquet F., Park Y.-H. Guinet C., Bailleul F., Charrassin J.-B.: Observations of the Fawn
- 1050 Trough Current over the Kerguelen Plateau from instrumented elephant seals. Journal of
- 1051 Marine Systems. 78:377-393, 2009.
- 1052 Sarmiento J., Gruber N., Brzezinski M., Dunne J.: High-latitude controls of thermocline
- 1053 nutrients and now latitude biological productivity. Nature. 427:56-60, 2004.
- 1054 Sarthou G., Chever F., Quéroué F., Bowie A., Van der Merwe P., Cheize M., Sirois M.,
- 1055 Bucciarelli E.: Fe-Cu impact in incubation experiments of natural plankton communities and
- 1056 Fe- and Cu-binding ligand production at the vicinity of the Kerguelen Island, Southern Ocean.
- 1057 In preparation for Biogeosciences.
- Strickland J., Parsons T.: A pratical handbook of sea water analysis. Fisheries research boardof Canada. 167:65-70, 1972.

- 1060 Tagliabue A., Mtshali T., Aumont O., Bowie A., Klunder M., Roychoudhury A., Swart S.: A
- global compilation of dissolved iron measurements: focus on distributions and processes inthe Southern Ocean. Biogeosciences. 9:2333-2349, 2012.
- 1063 Takahashi T., Sutherland S., Wanninkhof R., Sweeney C., Feely R., Chipman D., Hales B.,
- 1064 Friederich G., Chavez F., Sabine C., Watson A., Bakker E., Schuster U., Metzl N.,
- 1065 Yoshikawa-Inoue H., Ishii M., Midorikawa T., Nojiri Y., Körtzinger A., Steinhoff T.,
- 1066 Hoppema M., Olafsson J., Arnarson T., Tilbrook B., Johannessen T., Olsen A., Bellerby R.,
- 1067 Wong C.S., Delille B., Bates N.R., debar H.: Climatological mean and decadal change in
- 1068 surface ocean pCO2, and net sea-air CO2 flux over the global oceans. Deep-sea Research II.
- 1069 56:554-577, 2009.
- 1070 Takeda S.: Influence of iron availability on nutrient consumption ratio of diatoms in oceanic1071 waters. Nature. 393:774-777, 1998.
- 1072 Tréguer P., De La Rocha C. : The World Ocean Silica Cycle. Annual Review of Marine
 1073 Science. 5:477-501, 2013.
- 1074 Trull T., Rintoul S., Hadfield M., Abraham E.: Circulation and seasonal evolution of polar
- 1075 waters south of Australia: Implications for iron fertilization of the Southern Ocean. Deep-Sea1076 Research II. 48:2439-2466, 2001.
- 1077 Uitz J., Claustre H., Griffiths B., Ras J., Garcia N., Sandroni V.: A phytoplankton class-
- 1078 specific primary production model applied to the Kerguelen Islands region (Southern Ocean).
- 1079 Deep-Sea Research I. 56:541-560, 2009.
- 1080 Zhou M., Zhu Y., d'Ovidio F., Park Y.-H., Durand I., Kestenare E., Sanial V., Van-Beek P.,
- 1081 Quéguiner B., Carlotti F., Blain S.: Surface currents and upwelling in Kerguelen Plateau
- 1082 regions. Biogeosciences Discussion. 11:68-6876, 2014.
- 1083

1084 Tables

- 1085 Table 1: Characteristics of the stations sampled for Si-fluxes measurements during KEOPS-2.
- 1086 Ze represents the bottom of the euphotic zone (1% of surface Photosynthetically Active
- 1087 Radiation), MLD is the depth of the Mixed Layer (from Park et al., in prep.).

Station	Zone	Position		Date	7 0 (m)	
Station		Latitude	Longitude	Date	Ze (m)	MLD (m)
R-2	HNLC	50°21.55 S	66°43.0 E	26-Oct	92	124
E-1	Meander	48°27.4 S	72°11.3 E	30-Oct	64	69
E-3	Meander	48°42.1 S	71°58.0 E	4-Nov	68	35
F-L	Polar front	48°31.2 S	74°39.5 E	7-Nov	29	47
E-4W	Plume	48°45.9 S	71°25.5 E	12-Nov	31	55
E-4E	Meander	48°42.9 S	72°33.8 E	14-Nov	34	80
A3-2	Plateau	50°37.5 S	72°34.9 E	17-Nov	38	123
E-5	Meander	48°24.7 S	71°54.0 E	19-Nov	54	41

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1090	Table 2: Biogenic silica	concentration (∫[BSi]),	, Si-uptake (∫ρSi)	, biogenic silica dissolution
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($\int \rho Diss$), silica net production ($\int \rho Net$) integrated over the euphotic layer (1% of surface 1091

1092 Photosynthetically Active Radiation), and integrated specific rates of Si-uptake and silica

dissolution (calculated as $VSi = \int \rho Si / [BSi]$ and $Diss = \int \rho Diss / [BSi]$ respectively). 1093

	Zone	∫[BSi]	∫pSi	∫ρDiss	∫pNet	Specific rates	
Station		mmol m ⁻²	mmol $m^{-2} d^{-1}$	mmol $m^{-2} d^{-1}$	mmol $m^{-2} d^{-1}$	∫VSi (d ⁻¹)	∫VDiss (d ⁻¹)
R-2	HNLC	33.28 ± 0.1	3.09 ± 0.01	4.88 ± 0.01	-1.78 ± 0.02	0.09	0.15
E-1	Meander	96.1 ± 0.2	16.8 ± 0.1	7.11 ± 0.02	9.6 ± 0.1	0.17	0.07
E-3	Meander	83.6 ± 0.2	10.5 ± 0.1	9.99 ± 0.03	0.5 ± 0.1	0.13	0.12
F-L	Polar front	97.8 ± 0.5	27.5 ± 0.3	3.79 ± 0.03	23.8 ± 0.3	0.28	0.04
E-4W	Plume	142.0 ± 0.7	31.8 ± 0.3	3.97 ± 0.03	27.9 ± 0.3	0.22	0.03
E-4E	Meander	104.3 ± 0.5	21.0 ± 0.2	$5.89 \pm 0.03 *$	$15.1 \pm 0.2*$	0.20	0.06*
A3-2	Plateau	173.6 ± 0.7	47.9 ± 0.4	4.50 ± 0.03	43.4 ± 0.4	0.28	0.03
E-5	Meander	159.5 ± 0.4	27.5 ± 0.2	6.97 ± 0.03	20.5 ± 0.2	0.17	0.04

1094 1095

* Since no dissolution rates were measured at E4E, these values do not correspond to calculations from direct 1096 measurements but only to estimations. Dissolution was calculated as the average of all KEOPS-2 integrated 1097 dissolution rates.

1098

Table 3.: Dissolution to production ratio ([D:]P), fraction of the silica production supported by 1099

new silicic acid (1-D:P), silicon to carbon (C) and nitrogen (N) uptake ratios (C and N 1100

1101 assimilation were measured by Cavagna et al., in prep.). pN represents both nitrate and

1102 ammonium uptake. All these values are integrated over the euphotic layer (1% of surface

1103 Photosynthetically Active Radiation).

Statio	on Zone	∫D:∫P	1-∫D:∫P	gross-∫ρSi:∫ρC	gross-∫ρSi:∫ρN
R-2	HNLC	1.58	-0.58	0.28	0.44
E-1	Meander	0.42	0.58	0.38	1.27
E-3	Meander	0.95	0.05	0.18	0.74
F-L	Polar front	0.14	0.86	0.10	0.32
E-4V	V Plume	0.12	0.88	0.15	0.93
E-4F	E Meander	0.28	0.72	0.27	1.26
A3-2	2 Plateau	0.09	0.91	0.30	1.51
E-5	Meander	0.25	0.75	0.35	1.41

* Since no dissolution rates were measured at E4E, these values do not correspond to calculations from direct
measurements but only to estimations. Dissolution was calculated as the average of all KEOPS-2 integrated

- 1107 dissolution rates.
- 1108

1104

1109 **Figures captions**

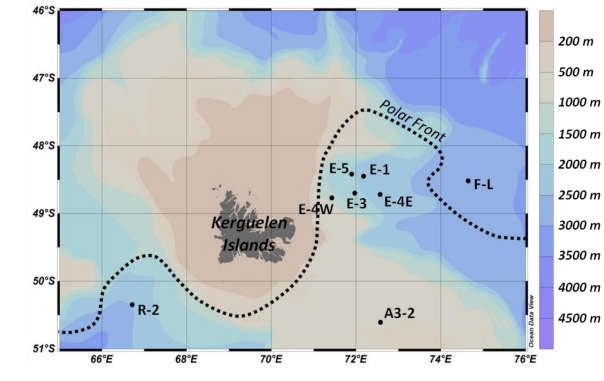
1110 Figure 1: Map of the KEOPS-2 cruise area (Indian sector of the Southern Ocean) showing the

- 1111 location of stations discussed in this study. Dotted line represents the position of the Polar1112 Front from Park et al. (in prep.).
- 1112 Front from Fark et al. (in prop.).
- 1113 Figure 2. Comparison of changes in ³⁰Si-abundance of seawater for each incubation (symbols)
- 1114 with detection limit of the ³⁰Si-isotopic dilution method (plain line) estimated from the
- 1115 reproductibility of an internal standard (0.43%, n = 40). The dotted line represents the
- 1116 detection limit obtained from the average reproductibility of all dissolution duplicates (0.32%,
- 1117 n = 35).
- 1118 Figure 3: Vertical distribution of chlorophyll *a* (continuous-black line; estimated from CTD
- 1119 fluorescence), biogenic silica concentration ([BSi], light dots) and H₄SiO₄ concentration
- 1120 ([DSi], dark dots). Dotted lines show the bottom of the euphotic layer (1% of
- 1121 Photosynthetically Active Radiation, Ze) for each station. Dark dashed lines represent the
- 1122 Mixed Layer Depth (MLD; estimated by Park et al., in prep.) and grey dashed lines
- 1123 correspond to a 2nd density gradient identify from the density CTD-profile.
- 1124 Figure 4: Vertical distribution of Si-uptake (pSi, panel a.) and biogenic silica dissolution
- 1125 (pDiss, panel b.) in KEOPS-2 stations. Open symbols represent the depth at the bottom of the
- 1126 euphotic layer (1% of Photosynthetically Active Radiation) for each station.

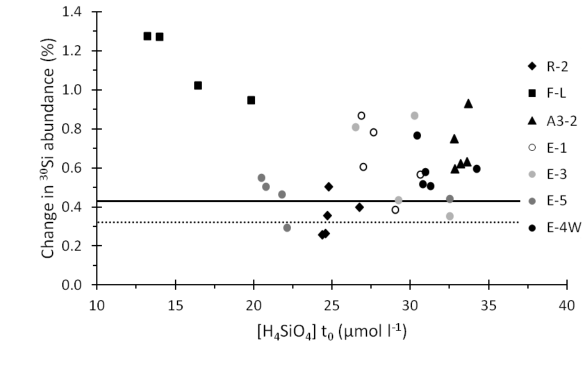
- 1127 Figure 5: Si-uptake (black), biogenic silica dissolution (white) and net silica production (grey)
- 1128 integrated over the euphotic layer (1% of Photosynthetically Active Radiation). Italic values 1129 correspond to the integrated dissolution to production ratio ($\int D: \int P$).
- 1130 Figure 6: Fraction of biogenic silica production supported by new silicic acid (1-[D:[P) as a function of the integrated net silica production rate ($\int \rho_{net}$) during KEOPS-2 (filled black 1131 1132 diamonds) compared with different regions of the global ocean (Open circles; Brzezinski et 1133 al., 2003 and references therein; and open diamonds; Fripiat et al., 2011b). Triangles and 1134 squares show the mean values for different growth seasons for SAZ and PFZ, respectively 1135 (Fripiat et al., 2011b). The plain line is a rectangular hyperbola to fit all data points (grey 1136 symbols were excluded from the model since they represent either negative 1-JD:JP value which are not allowed by the model, or average while all other symbols refer to single 1137 stations). The equation of the curve is $1-\int D : \int P = 1.01 \text{ x} \int \rho_{net} / (5.89 + \int \rho_{net})$. Dotted lines 1138
- 1139 correspond to \pm 1sd of the 1-JD:JP_{max}.

1140 Figure 7: Schematic view of the seasonal silicon cycle in the mixed layer above the Kerguelen 1141 Plateau as estimated from natural and enriched Si isotopic measurements. Blue silicon fluxes 1142 correspond to estimated values while dark fluxes correspond to direct measurements. The 3 1143 main water masses are represented by dark-grey for Upper Circumpolar Deep Water 1144 (UCDW), medium-grey for the Winter Water (WW) and light-grey for the Mixed Layer 1145 (ML). Variation of biogenic silica and H₄SiO₄ standing stocks integrated over 80m 1146 (respectively [BSi] and [DSi]) are shown in the upper panel. Vertical continuous arrows 1147 represent DSi supplies from deep water to the ML, and dotted arrows correspond to 1148 particulate silica fluxes. Integrated silica production rates are calculated from the surface to 1149 80m. Horizontal white arrows represent the state of the bloom (indicated by the D:P ratio) 1150 through time. Winter consumption has been estimated from the difference of winter mixing 1151 supply (Fripiat et al., 2011a) and the standing stock measured at the first A3 visit (KEOPS-2). 1152 The difference in standing stocks between the two visits at A3 (28 days, KEOPS-2) yields to the net silica production of 14.3 mmol.m⁻².d⁻¹. The DSi standing stock at the second visit of 1153 1154 A3 can sustain the net silica production measured for 32 days (this study) while the summer 1155 DSi supply estimated by Fripiat et al. (2011a) can sustain the same net production by an extra 1156 26 days. The fall production measurements and standing stocks are from Mosseri et al. (2008, 1157 KEOPS-1). Since no silica dissolution is available from KEOPS-1, fall net production has 1158 been estimated from the difference between average gross silica production during KEOPS-1

- 1159 (Mosseri et al., 2008) and average silica dissolution measured at all stations south of the Polar
- 1160 Front (KEOPS-2, this study). Deep BSi accumulation is calculated by integrating over the
- 1161 100-200m depth layer data from Mosseri et al. 2008 (see text for further details).
- 1162
- 1163 Figures



1165 Figure 1.



1166 1167 Figure 2.

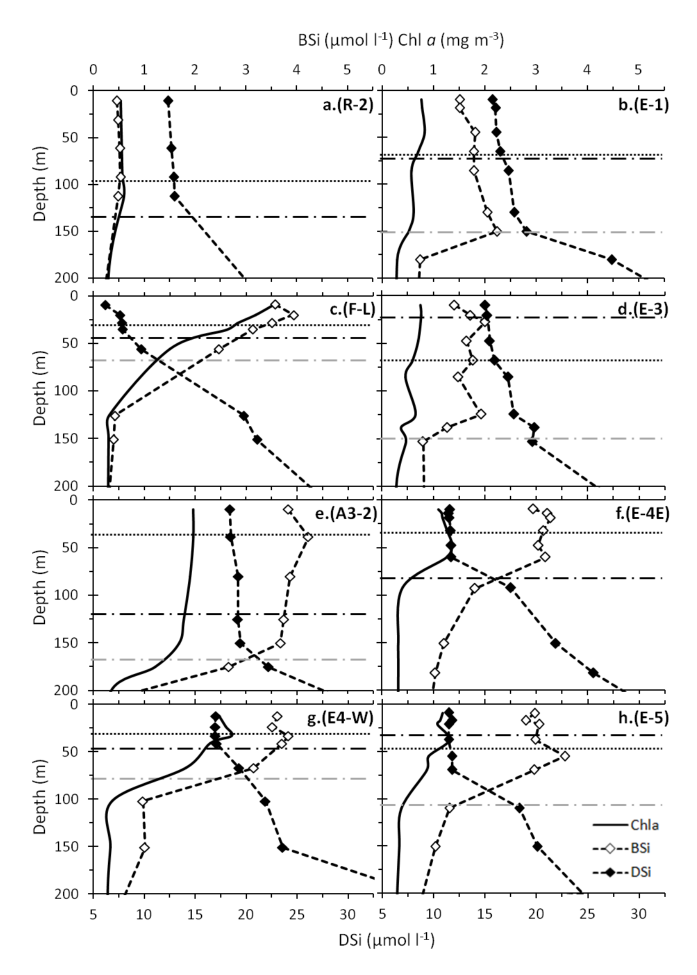
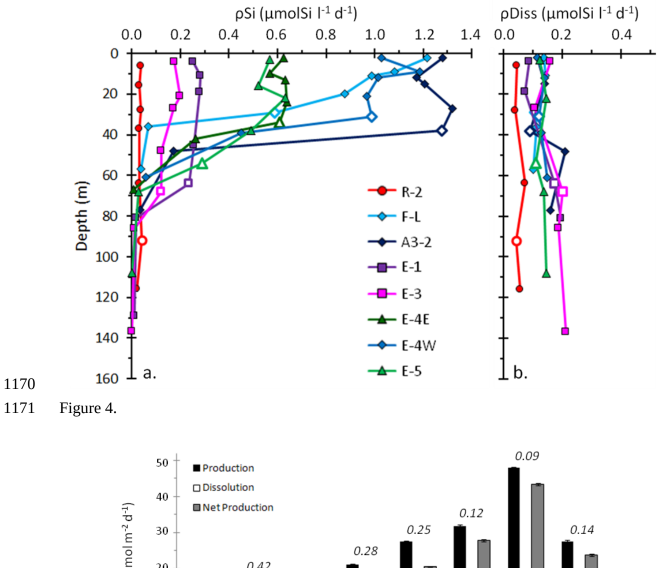
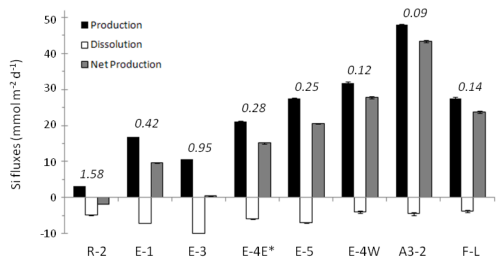


Figure 3. 1169



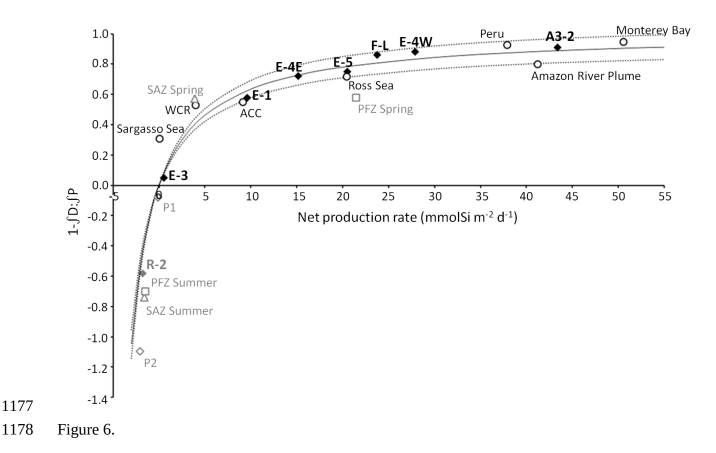


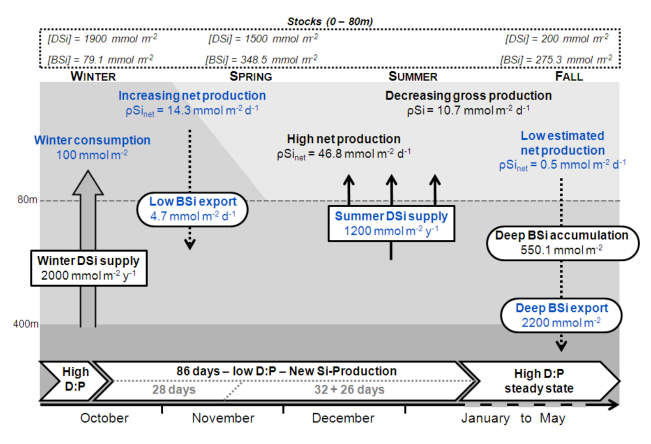


1173 * Since no dissolution rates were measured at E4E, dissolution and net production do not correspond to

1174 calculations from direct measurements but only to estimations. Dissolution was calculated as the average of all

- 1175 KEOPS-2 integrated dissolution rates.
- 1176 Figure 5.





1179

1180 Figure 7.