

Si–C interactions during degradation of the diatom Skeletonema marinoi

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1	Si-C interactions during degradation of the diatom <i>Skeletonema</i>
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27 Abstract

28 While a relationship between ballast and carbon in sedimenting particles has been welldocumented, the mechanistic basis of this interaction is still under debate. One hypothesis is that 29 30 mineral ballast protects sinking organic matter from degradation. To test this idea, we undertook a laboratory experiment using the diatom *Skeletonema marinoi* to study in parallel the 31 32 dissolution of one of the most common mineral ballasts, biogenic silica ($bSiO_2$), and the 33 associated degradation of organic matter. Three different models were applied to our results to 34 help elucidate the mechanisms driving bSiO₂ dissolution and organic compound degradation. Results of this modelling exercise suggest that the diatom frustule is made up of two bSiO₂ 35 36 phases that dissolve simultaneously, but at different rates. In our experiments, the first phase was more soluble $(k_{bSiO2} = 0.27 \text{ d}^{-1})$ and made up 31% of the total bSiO₂. In this phase, bSiO₂ was 37 38 mainly associated with membrane lipids and the amino acids glutamic acid, tyrosine, and leucine. The second phase was more refractory ($k_{bSiO2} = 0.016 \,\mathrm{d}^{-1}$), and contained more neutral 39 lipid alcohols and glycine. Until it dissolved, the first bSiO₂ phase effectively protected much of 40 the organic matter from degradation: POC degradation rate constants increased from 0.025 d^{-1} 41 to $0.082 d^{-1}$ after the total dissolution of this phase, and PON degradation rate constants 42 increased from $0.030 d^{-1}$ to $0.094 d^{-1}$. Similar to POC and PON, the THAA degradation rate 43 constant increased from $0.054 d^{-1}$ to $0.139 d^{-1}$ after dissolution of the first bSiO₂ phase. The 44 45 higher THAA degradation rate constant is attributed to a pool of amino acids that was produced during silicification and enclosed between the two silica phases. This pool of amino acids might 46 come from the incorporation of silica deposition vesicles into the diatom wall and might not be 47 48 directly associated with $bSiO_2$. In contrast, most lipid degradation was not prevented by 49 association with the more soluble bSiO₂ phase as the average lipid degradation rate constant decreased from $0.048 d^{-1}$ to $0.010 d^{-1}$ after 17 days of degradation; This suggests that most 50 51 lipids were associated to rather than protected by silica, except pigments that appeared resistant

- 52 to degradation, independantly from silica dissolution. When the only organic compounds
- 53 remaining were associated with the second $bSiO_2$ phase, degradation rate constants decreased
- 54 greatly; concentrations changed only slightly after day 25.
- 55
- Key Words: Biogenic silica, dissolution, carbon, amino acids, lipids, degradation, diatom

56 1. Introduction

57 Organic carbon produced in the ocean's surface layer by phytoplankton is conveyed to depth by particle sedimentation, and fluxes of carbon and minerals (CaCO₃, SiO₂ and aluminosilicates) 58 59 are highly correlated in the deep water column. Based on these observations, Armstrong et al. 60 (2002) highlighted the importance of modeling both carbon and mineral fluxes at the same time. 61 Mineral ballast (CaCO₃ of coccolithophorids; SiO_2 of diatoms; aluminosilicates in dust) provide 62 excess density needed for organic matter to sink; bSiO₂ and carbonate sedimentation are also 63 linked through the ability of phytoplankton to aggregate and through grazing by zooplankton. The combination of these processes strongly increases the sedimentation rate of phytoplankton 64 65 (e.g., Gehlen et al., 2006; Moriceau et al., 2007).

66 The role of mineral ballast in carbon transport is more complex than a simple impact on 67 excess density (Lee et al., 2008), but we are far from fully understanding the processes involved. 68 Lee et al. (2000) and Hedges et al. (2001) hypothesized that mineral ballast could protect organic 69 carbon from degradation; their hypothesis is consistent with the observation of Ingalls et al. 70 (2006) that organic matter was more degraded in areas where diatoms were not the dominant 71 bloom species when compared to sites where diatoms were the main phytoplankton group. Engel 72 et al. (2008) also showed that the presence of the calcite test in coccolithophorids lowers the 73 POC degradation rate during the recycling of these cells. Continuously increasing pressure 74 reduced rates of bSiO₂ dissolution of diatom detritus relative to rates measured under atmospheric pressure conditions (Tamburini et al. 2006). In parallel, naturally collected sinking 75 76 particles, were also less degraded by prokaryotes when pressure was continuously increased to 77 simulate descent from 200 to 1500 (Tamburini et al. 2008). Despite all these findings, few 78 studies (e.g., Ingalls et al., 2003; 2006) have investigated both organic matter degradation and 79 biogenic mineral dissolution in natural settings. The work of Engel et al. (2008) investigated the

80 role of CaCO₃ in carbon degradation, while the present study aims to better understand the role 81 of Si-C interactions during diatom degradation.

82 Diatoms are the dominant species in many ecosystems; they are responsible for up to 35% of 83 the total primary production in oligotrophic oceans and up to 75% in coastal waters and the Southern Ocean (Nelson et al., 1995; Tréguer et al., 1995). Jin et al. (2006) estimated their 84 85 global contribution to net primary production and to carbon export to be 15% and 40%, respectively. The high diatom contributions to primary production and carbon export, could 86 87 potentially explain the empiric relation established by Ragueneau et al. (2002). This relation 88 shows that Si/C ratios decrease with depth and follow the same pattern everywhere in the world 89 ocean. Is there a link between bSiO₂ dissolution and POC degradation such as the one

90 hypothesized in the work cited above?

91 The objective of the present study was to understand how biogenic silica influences the 92 degradation of diatom organic carbon, and conversely the role of organic compounds in $bSiO_2$ 93 dissolution. With this aim, a monospecific culture of the diatom Skeletonema marinoi was 94 incubated in the presence of a natural coastal bacterial community and allowed to degrade over a 95 102-day period. bSiO₂ dissolution and the quantity and composition of organic compounds, 96 including amino acids and lipids, were assessed throughout the incubation period and used to 97 investigate Si-C interactions during decomposition. Three dissolution/degradation models were 98 applied to the experimental data to elucidate the dissolution/degradation pattern of the 99 components of S. marinoi. This modeling experiment yields a better understanding of the 100 structure of the diatom frustule in S. marinoi and of the role of Si-C interactions during diatom 101 recycling.

102

2. Material and methods 103

104 2.1 Biodegradation experiment 105 Skeletonema marinoi (CCAP 1077/3) obtained from IFREMER (Argenton station, France) was grown in f/2 medium (Guillard and Ryther, 1962; Guillard, 1975) under 12/12 dark/light 106 illumination. When cells reached stationary growth phase (6.5 x 10^6 cells ml⁻¹), they were 107 108 transferred into a 4°C chamber and kept in the dark for 5 days. During this period, cells sank to 109 the bottom of the flask, and previous tests showed that diatom viability (number of living cells 110 versus total cells) decreased (unpublished data, method described in Garvey et al. 2007). The 111 supernatant was poured off, and the overlying medium was replaced with natural seawater that 112 had been passed through a 0.7-µm GFF filter to preserve the natural bacteria assemblage. The 113 seawater was collected from a small inlet (Endoume) near Marseille, France, at the end of fall, 114 when the water is naturally poor in silicic acid (dSi~2.5 µM). The mixture of S. marinoi and 115 filtered sea water was then transferred to an incubation flask equipped with a magnetic stirrer 116 and a stopper through which gas exchange could occur via a 0.2-µm Swinnex® filter. The 117 diatoms were incubated for 102 days in the dark at 20°C. Using a peristaltic pump, samples were 118 taken daily for the first 21 days and then at 23, 25, 46, 50, and 102 days; triplicate samples were 119 taken at 0, 5, 11, 46, 50, and 102 days. The sampled solution was well-mixed, allowing the ratio 120 of solid matter to solution to remain constant (Dixit et al., 2001). Ten percent of the liquid volume remained at the end of the experiment. Chemical parameters measured were biogenic 121 122 silica (bSiO₂), silicic acid (dSi), particulate organic carbon and nitrogen (POC, PON), dissolved 123 organic carbon (DOC), total particulate lipids (TLip), and total hydrolyzable amino acids 124 (THAA). In addition total bacterial abundances (diamidinophenylindole: DAPI counts) were 125 counted. Si contamination by dissolution of glassware was measured by analyzing dSi in an incubation bottle with no cells added. We also sampled controls poisoned with 20 mg l^{-1} HgCl₂ 126 127 at 4 times (0, 2, 5, and 11 days) to verify that degradation was due to bacteria and not abiotic 128 factors.

129

130 2.2 Analytical Methods

131 Biogenic Silica (bSiO₂) was determined at the beginning and end of the experiment using 132 a variation of the method of Ragueneau and Tréguer (1994). As no lithogenic silica was present 133 in the algal culture, the second digestion step using HF was not necessary. Ten-ml samples were 134 filtered onto 0.2- μ m polycarbonate filters. Filters were analyzed for bSiO₂ and the filtrate for dSi. For bSiO₂, filters were digested in 20 ml of 0.2N NaOH for three hours at 95°C to ensure 135 136 the dissolution of all bSiO₂; dSi concentrations in the solution remained far below the solubility 137 equilibrium of bSiO₂ at all times. After cooling, the solution was acidified with 5 ml of 1N HCl, 138 centrifuged to remove remaining solids, and analyzed for dSi. The precision for triplicate 139 measurements of $bSiO_2$ was < 5%.

140 Silicic acid (dSi) concentrations were determined on 10-ml filtered samples and on 141 digested $bSiO_2$ samples using the molybdate blue spectrophotometric method of Mullin and 142 Riley (1965), as adapted by Tréguer and Le Corre (1975) and modified by Gordon *et al.* (1993) 143 for use in segmented flow colorimetry. We used a Bran and Luebbe Technicon Autoanalyzer 144 (<1% precision).

145 POC and PON concentrations were measured using a Carlo Erba NA 2100 CN analyzer 146 coupled to a Finnigan Delta S mass spectrometer. Five-ml samples were filtered through 0.7- μ m 147 GFF filters. The filters were desiccated overnight in an oven at 50°C and then placed in tin 148 capsules to be introduced into the oven of the analyzer. The precision for triplicate N analyses 149 was \pm 1-6%, and for C analysis \pm 1-5%.

DOC was analyzed after filtration through 0.7-μm GFF filters; 10 ml of each sample was transferred into glass ampoules and sealed after addition of H_3PO_4 as preservative. All glassware was pre-rinsed with 1N HCl and Milli-Q water before being combusted at 465°C; care was taken to minimize contamination during sampling and handling. DOC was measured by hightemperature catalytic oxidation using a Shimadzu TOC 5000 Analyzer (Sempéré *et al.*, 2003; Sohrin and Sempéré, 2005). Samples were acidified to pH 1 with 85% phosphoric acid and bubbled for 10 minutes with CO₂-free air to purge them of inorganic carbon. Three or four 100 157 μ l replicates of each sample were injected into the 680 °C column. The precision of these 158 replicates was $\leq 6\%$.

159 Total particulate lipids were analyzed after filtering 10 ml samples onto 0.7-µm GFF 160 glass fiber filters. Filters were extracted according to Bligh and Dyer (1959). Lipid extracts 161 were separated into classes of compounds and quantified on an Iatroscan model MK-6s (Iatron, Tokyo; H₂ flow 160 ml min⁻¹; air flow 2 l min⁻¹) as described by Goutx *et al.* (2007). The 162 163 elution scheme allows reliable separation and quantification of degradation metabolites from acyl-lipid classes (Striby et al., 1999). Total particulate lipids (TLip) are the sum of the 164 165 separated lipid classes (Table 1). In the present work, the variability within triplicates was < 166 13%.

Amino acids were analyzed on 0.7-µm GFF filters after filtration of 10-ml samples. 167 168 Thawed filters were treated as described in the study of Ingalls et al. (2003). Individual compounds were separated by HPLC using pre-column OPA derivatization after acid hydrolysis 169 170 as described in Lee and Cronin (1982) and Lee et al. (2000). Amino acids were detected by 171 fluorescence and identified by comparison to retention times of standards made from an amino 172 acid mixture (Pierce, Standard H). The non-protein amino acids β -alanine and γ -aminobutyric 173 acid (BALA and GABA) were added individually to the standard mixture. Aspartic acid (ASP) 174 and glutamic acid (GLU) measurements include the hydrolysis products of asparagine and 175 glutamine. THAA is the sum of the 16 characterized amino acids (Table 1). Variation among 176 replicates was generally 15-30%. LYS replicates, however, varied more greatly at times, e.g., 177 50% at day 11.

Total bacterial abundances (DAPI counts): Subsamples for bacterial cell counts were
fixed immediately with buffered formalin (final volume 2%). Cells were collected onto a 25-mm
0.2-μm polycarbonate Nuclepore® membrane and stained with diamidinophenylindole (DAPI;
Porter and Feig, 1980). Slides were stored frozen until counting by epifluorescence microscopy
(Olympus, BH2).

184 **2.3 Kinetics**

Kinetic parameters were calculated over the first 25 days where experimental data are available every 1-2 days. For each compound of interest we tested three degradation/dissolution models. Model 1 is a simple first-order rate equation as described in Greenwood *et al.* (2001) and used in many dissolution studies (e.g. Kamatani and Riley, 1979; Kamatani *et al.*, 1980; Kamatani, 1982):

190
$$\hat{C}(t) = C_0 \exp(-kt)$$
, (1)

where $\hat{C}(t)$ is the concentration (μ M) estimated at time *t* (d), C_0 is the initial concentration, and *k* is the dissolution/degradation rate constant (d^{-1}).

Model 2 assumes simultaneous dissolution/degradation of two phases. The equation used
is similar to that used for carbon degradation in the study of Westrich and Berner (1984):

195
$$C(t) = C_1 \exp(-k_1 t) + C_2 \exp(-k_2 t)$$
 (2)

196 In Model 2, four parameters are estimated: C_1 and C_2 are concentrations (μ M) of phase or pool 1

and 2, and k_1 and k_2 (d⁻¹) are their respective dissolution or degradation rate constants.

As it uses 2 more parameters, Model 2 always gives a better fit than Model 1 except when the initial degradation rate is slower than later rates. In this specific case Model 2 performs no better than Model 1. We therefore developed Model 3, which employs one firstorder equation initially, and a second first-order equation after that. The time at which the dissolution/degradation rate constant changes is called the substitution time (t_s), and is allowed to take on any value >0.

204
$$\hat{C}(t) = C_0 \exp(-k_1 t), \quad 0 < t < t_s;$$

 $\hat{C}(t) = C(t_s) \exp(-k_2 t), \quad t > t_s$ (3b)

206 Model 3 also contains 4 parameters.

(3a)

207 If dissolution/degradation for a given compound is best reconstructed using Model 2, this

208 compound is constituted by at least two phases or pools; on the other hand if Model 3 gives a

209 better description of the dissolution/degradation pattern, either the compound studied is present

210 in 2 phases/pools remineralizing one after the other, or a change in environmental parameters

211 provoked a change in the dissolution/degradation rate constant at time *ts*.

212 In addition, to allow direct comparison of the degradation/dissolution pattern among all

213 compounds, and between this study and previous studies, an initial disappearance rate constant

was calculated for each compound over the first 10 days using Model 1.

215 **2.4 Statistics**

Each fit was optimized by maximizing the likelihood statistic log(L) as described in equation 4 (Armstrong *et al.*, 2002, and references therein). Eq. (4) is based on a Gaussian distribution with a constant variance on a logarithmic scale:

219
$$\log(L) = -\frac{N}{2} \times \log\left(\frac{\sum(\log(\hat{C}_j) - \log(C_j))^2}{N}\right) , \qquad (4)$$

where N is the number of data points, C_j is a measured concentration for data point j, and the \hat{C}_j is the corresponding model prediction. The difference in log(L) (Δ log(L)) between fits of two different models to the same data gives the goodness of fit of one model compared to the other. If one model gives a value for log(L) that is at least 2 points higher per added parameter than another model, it is considered to fit the data better (Hilborn and Mangel, 1997). In the present work, the simplest model (Model 1) was considered to be the best fit unless Model 2 or Model 3 yielded a log(L) more than 4 points better than that of Model 1.

227

228 **3. Results**

230 **3.1** Change in biochemical composition of the diatom *Skeletonema marinoi* over time

231 *3.1.1 General trends*

Changes in bSiO₂, POC, TLip and THAA concentrations over time are shown in
logarithmic scale in Fig. 1. While bSiO₂ concentrations decreased smoothly over the first 25 d,
POC, PON and THAA concentrations decreased until about day 14, when the loss rate increased,
especially for THAA. In contrast, TLip concentration decreased rapidly at the beginning of the
experiment and reached a plateau after day 13.

Bacterial concentrations started at $0.33 \pm 0.03 \times 10^{6}$ cell ml⁻¹ and peaked at day 14 with 237 a concentration of $30 \pm 2.7 \times 10^6$ cell ml⁻¹ (Fig. 2). The bacterial population increased between 238 days 0 and 14 with a rate equal to $0.015 d^{-1}$ (calculated between days 2 and 14) and then 239 generally stabilized reaching a final concentration of $20 \pm 4 \ge 10^6$ cell ml⁻¹ until the end of the 240 experiment. The bacterial growth efficiency (bacterial carbon increase divided by POC decrease) 241 242 between days 0 and 14 was 2%. Bacterial carbon made up a maximum of $2.4 \pm 0.6\%$ of the POC. TOC concentrations over time showed a pattern similar to those of POC (Fig. 3). DOC 243 concentrations increased slightly $(5.6 \pm 0.2 \text{ mg C L}^{-1})$ but much less than POC decreased $(62 \pm 3$ 244 mg C L^{-1}). The bacterial carbon (C_{bact}) is so low compared to the algal organic carbon that the 245 mineralization rate for POC algal (POC+DOC- C_{bact}) is the same as for the total organic carbon 246 (Corg = POC+DOC). This rate calculated as the slope of TOC change over time (Fig. 3) was 247 2.68 mg C $L^{-1} d^{-1}$ during the first 20 days of the experiment and was equivalent to 91% of the 248 249 POC loss.

250

251 3.1.2 Initial biochemical composition

At the beginning of the experiment, the Si/POC ratio was 0.09 ± 0.01 , which is slightly lower than the value of 0.13 measured in fresh diatoms by Brzezinski (1985) but in the range of coastal diatoms measured by Rousseau *et al.* (2002). TLip made up $19 \pm 2\%$ of *S. marinoi* 255 organic carbon which is a little bit higher that Lip/POC ratios measured previously for the same 256 specie (Lavens and Sorgeloos, 1996). TLip mainly included FFA and other degradation 257 metabolites (ALC and MG; see Table 1 for abbreviations) (Fig. 4a). Cellular membrane phospholipids (PE, DPG+PG) and chloroplast membrane glycolipids (MGDG) accounted for 10 258 259 \pm 1% and 13 \pm 1% of TLip, respectively. FFA were the most abundant lipid degradation metabolites (24.8 \pm 0.5%) among those present. PIG included both chlorophyll *a* and its 260 degradation products (Striby *et al.*, 1999); PIG was the largest lipid class (initially $41 \pm 5\%$; Fig. 261 262 4a). ST are involved in membrane rigidity (Parrish, 1988); they were initially minor components 263 of S. marinoi.

264 THAA constituted a larger portion $(36 \pm 9\%)$ of total organic carbon than the lipids, similar to the 45% found in Thalassiosira weissflogii by Cowie and Hedges (1996) and the 25% 265 observed in diatom-rich sediments by Ingalls et al. (2006). THAA include the monomer 266 267 constituents of protein as well as adsorbed amino acids and peptides. Sixteen amino acids were 268 quantified. GLU, ASP and LEU (see Table 1 for abbreviations) together made up one third of 269 the THAA, each more than $11 \pm 3\%$ (Table 2). Other amino acids were between $2.1 \pm 0.6\%$ and 270 $9.3 \pm 2.8\%$ of the THAA except for MET and GABA, which were less than 0.8%. As GABA and MET concentrations were very low throughout the incubation, they are not described in the 271 272 following paragraphs. The initial mole% amino acid compositions we found for S. marinoi 273 (Table 2) were very similar in pattern to those of cultured diatoms reported by Ingalls et al. 274 (2003) and Cowie and Hedges (1996), with highest mole% values for ASP, GLU, and GLY. S. marinoi was higher in mole% LEU than in reports of other diatoms. THAA do not include the 275 276 amino acids incorporated inside the silica frustule, unless part of the frustule is dissolved during 277 acid hydrolysis (see later discussion). We define Si-bound amino acids as Si-THAA as in Ingalls 278 et al. (2003).

279

280 *3.1.3 Change in biochemical composition during degradation*

281	Si/POC decreased from 0.09 ± 0.01 mol of Si/mol of C to 0.06 ± 0.01 mol of Si/mol of C
282	during the first 2 days of incubation and then stabilized until day 15 of the experiment. After
283	increasing to 0.11 ± 0.01 mol of Si/mol of C between days 15 and 21, the Si/POC ratio remained
284	constant until the end of the experiment. TLip/POC varied between 0.13 ± 0.02 mol of C/mol of
285	C and 0.23 ± 0.03 mol of C/mol of C during the first 50 days. Then the ratio decreased to a final
286	value of 0.06 ± 0.01 mol of C/mol of C at day 102. THAA/POC decreased regularly during the
287	whole experiment from 0.36 ± 0.09 mol of C/mol of C initially to 0.10 ± 0.02 mol of C/mol of C
288	after 102 days, except for a sudden increase between day 11 and 15 from 0.28 ± 0.7 mol of
289	C/mol of C to 0.38 ± 0.10 mol of C/mol of C

290 Most of the change in TLip composition (Fig. 4a) occurred between days 0 and 25. Most 291 compounds decreased or remained the same relative to TLip (mol of C/mol of C) except for PIG, 292 which increased from $41 \pm 5\%$ to $80 \pm 7\%$ of the TLip over the course of the experiment. 293 Relative concentrations (mol of C/mol of C) of FFA, the second most abundant lipid class in the 294 algae, decreased regularly from $24.8 \pm 0.5\%$ to $4.0 \pm 0.4\%$. MGDG initially made up $13 \pm 1\%$ of 295 TLip, but totally disappeared by day 4. The MG contribution was constant $(7.0 \pm 0.3\%)$ until day 296 14, when it was completely degraded. The contribution of membrane lipids, the glycolipid 297 MGDG and the phospholipids (PE and DPG+PG) to TLip was low compared to results from 298 Berge et al. (1995) and d'Ippolito et al. (2004). However, lipid composition is highly dependent 299 on culture conditions (d'Ippolito et al., 2004), and in our case the high FFA content probably 300 masked the relative contribution from membrane lipids.

The THAA composition was relatively constant throughout the degradation experiment except between days 15 and 20 where we observed a strong peak of GLY, which increased from $13 \pm 2\%$ to $33 \pm 8\%$ (Fig. 4b). The relative concentrations of HIS (data not shown) and LYS also peaked slightly between days 15 and 20. Relative proportions of other amino acids especially ASP decreased at this time in response to the increases in HIS, GLY and LYS.

307 **3.2 Dissolution and degradation kinetics of** *S. marinoi* constituents

308 *3.2.1 Silica kinetics*

309 The experiment was conducted in glass bottles to eliminate carbon contamination. 310 Controls showed that after 102 days, dSi due to leaching from the glass was a maximum of 5% 311 of the dSi due to dissolution of diatom frustules. From an initial concentration of 680 ± 30 µmol L^{-1} , bSiO₂ decreased rapidly during the first 3 to 5 days of the experiment and then more slowly 312 313 (Fig. 1). After 25 days, 52% of the initial bSiO₂ was dissolved and 76% of the initial was 314 dissolved at the end of the experiment (102 d). The comparison between the log(L) of the three models describing $bSiO_2$ dissolution showed that Model 2 is 4.3 $\times 10^{16}$ times better than Model 1 315 $(\Delta \log(L) = 38.3)$ and almost 4.5 x10⁵ times better than Model 3 ($\Delta \log(L) = 13$). bSiO₂ was the 316 only constituent of the diatom with a dissolution pattern that was best described by Model 2 317 318 (Fig. 5a, Table 4) meaning that the frustule is most likely composed of two phases dissolving 319 simultaneously. The first phase of $bSiO_2$ constituted 31% of the total $bSiO_2$ and was more soluble, with a dissolution rate constant of $0.27 d^{-1}$. The second phase was more refractory with 320 a dissolution rate constant of $0.016 d^{-1}$ (Table 4). For direct comparison with POC and PON, 321 the initial loss rate constant of the total bSiO₂ was calculated using Model 1 over 10 days as 322 $0.049 d^{-1}$. The three dissolution constants are within the range $(0.005 d^{-1} \text{ to } 1.3 d^{-1})$ given in 323 324 the review by Van Cappellen et al. (2002b).

325

326 *3.2.2 POC and PON kinetics*

POC concentration decreased in two steps from the initial value of $7660 \pm 150 \ \mu \text{mol L}^{-1}$ (Fig. 1). Even with more parameters Model 2 did not improve the fit to the data obtained by Model 1 (Fig. 5b); $\log(L)$ calculations for POC loss showed that Model 3 gave the best fit $(\Delta \log(L) = 76.5)$. This suggests that either two pools of POC exist and are degraded successively, or a change in some parameter not directly linked to POC chemistry provoked an

332	increase of the degradation rate at day 10 from 0.025 to 0.082 d^{-1} (Table 4). PON followed the
333	same pattern; Model 3 reproduced the data with more accuracy as shown by the $\Delta \log(L)$ of 37.3
334	compare to Models 1 and 2. Model 3 estimated an increase in PON degradation rate at day 12
335	from 0.03 to $0.094 d^{-1}$ (Table 4). The first POC degradation rate constants are similar to rate
336	constants measured in previous studies $(0.036 d^{-1} \text{ for POC}, \text{ Lee and Fisher}, 1992; 0.035 d^{-1} \text{ for}$
337	POC and 0.047 d ^{-1} for PON, Harvey <i>et al.</i> , 1995). For comparison with bSiO ₂ , we also applied
338	Model 1 over 10 days to calculate the initial loss rate of POC and PON: $0.025 d^{-1}$ for POC and
339	$0.029 \mathrm{d}^{-1}$ for PON.

The POC degradation pattern in the Hg-poisoned controls was similar to that in unpoisoned flasks during the first 5 days but was lower between days 5 and 11. The degradation rate constant measured over the first 11 days using Model 1 was $0.021 d^{-1}$.

343

344 *3.2.3 Lipid kinetics*

From an initial TLip concentration of $1400 \pm 70 \mu mol C L^{-1}$, almost half $(42 \pm 4\%)$ 345 346 degraded in 25 days; 7.0 ± 0.3 % of TLip remained after 102 days. The degradation of 7 of the 8 lipid classes is shown in Fig. 6a. TLip degradation is better described by Model 2 and 3 than by 347 348 Model 1 ($\Delta \log(L) = 15.6$ and 17 respectively). With only 1.5 point of $\log(L)$ difference between 349 Model 2 and 3 but the same degree of complexity (4 parameters each) we chose the best fit from 350 the best likelihood which was given by Model 3 (Table 4). The degradation rate constant for TLip was $0.048 d^{-1}$ during the first 17 days of the experiment and then decreased to $0.01 d^{-1}$. 351 The initial degradation rate constant was $0.046 d^{-1}$, slightly higher than the $0.023 d^{-1}$ measured 352 353 in the study of Harvey et al. (1995).

Using Model 1 we calculated that FFA, PIG, MG and PG+DPG had degradation rate constants of 0.104, 0.012, 0.045 and $0.011 d^{-1}$ over 25 days respectively. For MGDG, ST, PE, and ALC, Model 3 gives the best fit to the data (Table 4).

MGDG turned over slowly $(0.078 d^{-1})$ during the first 2 days, but then much more 357 quickly $(0.52 d^{-1})$; Table 4) and were completely gone after only 4 days. MG were also 358 completely lost very quickly; the initial degradation rate constant of MG was $0.045 d^{-1}$, but the 359 remaining MG was gone after 14 days. This pattern does not fit any of the models used, and 360 suggests an association of these lipids only with the first $bSiO_2$ phase, or no association at all. 361 362 MGDG, ST, and PE degradation rate constants increased by a factor of 8 to 10 during the 363 experiment. They followed the same pattern as POC and PON. ALC degradation rate constant decreased after 19 days. 364

365 *3.2.4 Amino acid kinetics*

From the initial THAA concentration of $3020 \pm 200 \mu mol C L^{-1}$, $86 \pm 17 \%$ of the THAA 366 were degraded after 25 days; 5.0 ± 0.5 % of the THAA still remained after 102 d. As for POC 367 and PON, Model 3 was the best fit for THAA degradation ($\Delta \log(L) = 9.9$ with both Model 1 and 368 369 Model 2). Concentrations of THAA decreased from day 1 to day 13 with an average degradation rate constant of $0.054d^{-1}$. After 13 d THAA turned over with a faster degradation rate constant 370 of $0.139 d^{-1}$ (Table 4), before reaching a period of very low rate constants after day 25; THAA 371 372 concentrations were almost constant until day 102. This last rate was not calculated by models 373 as the modelling exercise was applied only over the first 25 days. The first degradation phase of THAA was similar to that measured on *T. weissflogii* $(0.058 d^{-1})$ by Harvey *et al.* (1995). 374 Initially, the degradation rate constant of the 14 individual amino acids ranged between 0.015 375 and $0.070 d^{-1}$, except for ASP and SER, which turned over more slowly with constants of 0.001 376 and $0.007 d^{-1}$, respectively (Table 4). At day 11 when on average 49 ± 10 % of the THAA had 377

degraded, all concentrations except ARG and SER suddenly increased by 6 to 58 % in 1 to 2

days (Fig. 6b). The largest releases were observed for HIS, GLY and LYS, which increased by

 39 ± 13 %, 58 ± 10 % and 46 ± 12 %, respectively. For every amino acid except TYR,

degradation rate constants increased after this peak. For ASP and SER the increase occurred

- earlier at day 5, and for TYR the degradation rate constant decreased from 0.070 to 0.012 d⁻¹ at
- 383 day 21. Except for TYR, amino acids turned over faster during the second degradation phase,
- and degradation rate constants ranged between 0.110 and 0.182 d^{-1} . The degradation rate
- 385 constant of SER increased even more than the other amino acids reaching $0.897 d^{-1}$ (Table 4).

386 **3.3 Relation between bSiO₂ dissolution and degradation of individual**

387 organic compounds or compound classes

388 3.3.1 Lipid degradation versus bSiO₂ dissolution

There was a strong linear relationship between total bSiO₂ and TLip over the whole range of bSiO₂ concentrations ($r^2 = 0.85$, n = 26) measured. There was no correlation between bSiO₂ and PIG, so that the relationship between bSiO₂ and TLip became even better when pigments were excluded from the other lipids ($r^2 = 0.94$, n = 26). FFA were well correlated with total bSiO₂ concentrations ($r^2 = 0.95$, n = 26); they were completely degraded during the dissolution of the second bSiO₂ phase, when bSiO₂ concentrations eventually reached 260 µmol L⁻¹.

395 Relationships among individual lipid classes and bSiO₂ phases showed three distinct 396 periods (Fig. 7a and b), which were related to $bSiO_2$ dissolution using the $bSiO_2$ model (Figure 397 5a). Period 1 (P1) is the time corresponding to dissolution of 85% of the first bSiO₂ phase and 398 10% of the second $bSiO_2$ phase; Period 2 (P2) is the time corresponding to dissolution of most of 399 the remaining bSiO₂ from the first phase and another 10% of the second bSiO₂ phase; and Period 400 3 (P3) is the time when less than 1.5% of the first phase remained and 60% of the second phase dissolved. At the end of P3 20% of the bSiO₂ from the second phase remained. On average, 26-401 402 34% of the TLip degraded during Period 1; only 14-19% of TLip remained at the beginning of

403 Period 3. The slow decrease of concentrations observed for each lipid class except PIG, during 404 P1 and P3 compared to P2 despite the fact that the three periods lasted the same time (~7 days, 405 Fig. 5a), might show that most TLip except PIG degraded during P1 were associated with the 406 first bSiO₂ phase, and most TLip except PIG degraded during P3 with the second bSiO₂ phase. 407 These specific lipids are denoted as Si(1)-Lip and Si(2)-Lip; their composition is shown Table 3. 408 Of the individual classes, PE, ALC concentrations remained almost constant during P1. 409 In contrast, MGDG was completely degraded within the first 4 days, in P1. Despite the high 410 variability in PE and ST concentration measurements, we determined that ST and MG concentrations decreased only slightly in P1 (>27%). This trend is illustrated by their low 411 degradation constants $(0.016 d^{-1} \text{ and } 0.001 d^{-1}, \text{ respectively})$. 412 413 Degradation of MG, ALC, PE and ST mainly occurred in P2. PE and ST concentrations dropped in P2 and ALC degradation was even faster; Model 3 gave a higher k_1 and a longer ts 414 415 for ALC than for PE and ST. After the precipitous drop, which corresponded to the beginning of 416 P3, 80% of ALC, FFA and PE and 40% of ST were degraded. MG were completely degraded so 417 quickly at the beginning of P2 that the loss is more likely due to sudden release of MG adsorbed 418 onto particles or dissolution (i.e. involving chemical mechanisms) rather than degradation (i.e. 419 biological mechanisms).

420 DPG+PG were not well correlated with $bSiO_2$; generally there was a 36%-degradation 421 during P1, then a fast release of DPG+PG in P2 (40-50%), possibly when the first $bSiO_2$ phase is 422 completely dissolved. Due to the release in P2, 80% of the initial DPG+PG remained at the 423 beginning of P3.

424

425 *3.3.2 Amino acid degradation versus bSiO₂ dissolution*

426 The relationship of THAA with the two $bSiO_2$ phases showed changes at the same times 427 as many of the lipid classes, so we used the same three periods. During Period 1, when the first 428 $bSiO_2$ phase was dissolving, only 3-22 % of the THAA degraded. For the reason described in the

previous section for lipids (3.3.1), this portion of the THAA is referred to as Si(1)-THAA. GLY 429 430 and LYS lost less than 10 % of their initial concentrations; ASP and GLU lost 40 % of their initial concentrations. GLU, ASP, and LEU constituted most of the pool degrading during P1; 431 432 LYS is not correlated to the first $bSiO_2$ phase (Table 2). At the end of period 2, a pool of THAA was released when 95 % of the first $bSiO_2$ phase 433 434 was dissolved (Fig. 8a and b). THAA concentrations decreased by 32 % compared to initial 435 values. Measurement of THAA does not release Si-bound amino acids so that they would not be 436 observed until the bSiO₂ dissolved; the THAA released (Si(2)-THAA) may have been trapped between the 2 phases. The amount of Si(2)-THAA can be calculated from the difference between 437 438 the concentrations of amino acids at and before the peak maximum (Table 2). In total, THAA concentration increased by 11-16 % (in μ mol C L⁻¹); the Si(2)-THAA were composed mainly of 439 440 GLY $(23 \pm 8 \%)$ and LYS $(23 \pm 8 \%)$; see Table 2).

During P3, relative THAA concentrations declined from 34 % to 5 % of the original
THAA. The THAA lost during P3 are called Si(3)-THAA, they had a composition similar to that
of total THAA, except that the contribution of GLY was higher (Table 2).

444

445 **4. Discussion**

446 **4.1 Importance of bacteria in** *S. marinoi* degradation

The very high concentration of algae, when compared to the bacterial concentrations, and the continued degradation in the presence of HgCl₂, suggest that the loss of organic matter was due not only to biological degradation, but also to physical and chemical factors (dissolution). In the presence of HgCl₂, organic matter turned over with a low rate constant ($0.02d^{-1}$, $r^2 = 0.86$, n= 8), which appeared to decrease even more after day 5 ($0.007d^{-1}$, n = 4; 2 replicates). This could be due to initial dissolution of organic matter as the cell begins to fall apart; smaller rate constants after some time would then be due to the absence of bacterial degradation. However, the lack of appropriate samples makes this observation tentative. The slower increase of DOC
concentrations compared to the decrease in POC (Fig. 3) suggests that most of the POC loss may
be due to bacterial degradation despite the similar degradation rate measured in HgCl₂-poisoned
batches. Degradation of particulate matter in the presence of HgCl₂ has been noted before (Liu *et al.*, 2006).

It was not clear why bacteria grew so slowly after day 14 in the unpoisoned experiment. Three factors could have contributed to the decrease and stabilization of bacterial growth. First, O_2 could have been a limiting factor. We did not measure O_2 during the experiment but we calculated average TOC loss, $R_{loss} = 221.6 \mu mol L^{-1}$. Change in oxygen concentrations (C_{O2}) with time (t) can be reconstructed from O_2 consumption rate (R_{loss}) and O_2 diffusion rate (R_{diff}) (eq. 5). The latter is given by the Whitman film model (Gladyshev 2002; eq. 6).

465
$$\frac{dC_{O2}}{dt} = R_{diff} - R_{loss}$$
(5)

466
$$R_{diff} = \frac{D}{\delta} \times \frac{S}{V} \times (Cs_{O2} - C_{O2})$$
(6)

where δ is the thickness of the diffusion layer, and is strongly dependent on stirring; D is the O₂ 467 diffusion coefficient (1.83 cm² d⁻¹, Ploug, 2001); S is the surface of contact between air and 468 water (415.5 cm²); V is the volume of the solution, which changed progressively during 469 470 sampling; and Cs_{O2} is the saturation concentration of the O₂ in seawater (229.9 μ M). Using these equations and parameters, we estimated the maximum δ above which the solution will be anoxic. 471 472 Considering that the risk of consuming all O₂ is greater for a larger volume of solution (with the 473 same surface area), we conservatively used the volume of solution at the beginning of the experiment $V=V_0$ (8100 cm³). 474

475 We seek a value of δ_{max} , at which $\frac{dC_{O2}}{dt}$ will always be positive when C_{O2} approaches 0 so that 476 the system can never go anoxic. This condition is met when

477
$$R_{loss} \le R_{diff} = \frac{D}{\delta} \times \frac{S}{V} \times C_{SO2}$$
(7)

478 or whenever

479
$$\delta_{\max} = \frac{D}{R_{loss}} \times \frac{S}{V_0} \times Cs_{O2}$$
(8)

We calculated that when the volume is at its maximum in the flask (8100 cm^3 at the beginning of the experiment), the depth of the diffusive layer must be less than 0.1 cm for the solution to remain oxic. The thickness of the diffusion layer is 0.27 cm with no stirring and can be as small as 0.0015 cm when the stirring is intense (Gladyshev, 2002). Since our flasks were well mixed, it is very unlikely that O₂ was a limiting factor.

A second explanation is that degradation products built up in the flask and poisoned the bacteria (Westrich and Berner, 1984; Aller and Aller, 1998). We cannot exclude this possibility, but calculating kinetic parameters over only 25 days should alleviate some of this problem. This period of time is a reasonable compromise between the need to follow the degradation of *S*. *marinoi* as long as possible so as to better understand reactions in the water column and the risk of accumulating inhibiting metabolites. Finally, the bacteria might have stopped growing due to viral lysis, grazing or a lack of labile substrate fuelling their growth (Fig. 2).

492

493 **4.2 Importance of Si-C interactions to bSiO₂ dissolution**

494 Previous dissolution studies have assumed that the diatom frustule is composed of a 495 single bSiO₂ phase (see review in Van Cappellen *et al.*, 2002b). Results from our modelling 496 exercise suggest instead that the frustule of *S. marinoi* is composed of two phases of bSiO₂ with 497 different dissolution characteristics. This idea, previously hypothesized by Kamatani and Riley 498 (1979) from dissolution rate measurements and by Gallinari *et al.* (2002) from solubility 499 equilibrium measurements, is consistent with the complexity of the frustule structure. During 500 silicification, polyamines and silaffin proteins catalyze precipitation of organo-silicon particles

501 of different shape and structure that determine the morphology of different diatom species 502 (Kröger et al., 2000; Hildebrand, 2003). As a result of these interactions, diatom frustules have a 503 complex 3-D structure and are shaped like an elliptic or cylindrical box. Each half is composed 504 of a valve and girdle bands that are built at different times in the cell cycle (Hildebrand and 505 Wetherbee, 2003). In our study we distinguish two $bSiO_2$ phases and calculate their dissolution 506 rate constants. Even though we didn't determine a direct relation between these dissolution 507 characteristics and the structure of the frustule, we were able to determine the impact of two-508 phase dissolution on the organic matrix of the cells as discussed below (see section 4.3). 509 Diatom frustules include organic layers that consist mainly of sugars and amino acids 510 (Hildebrand et al., 2006). The major amino acids in this coating are GLY, THR, and SER, 511 suggesting that bonding with silica occurs through SER and THR, using their OH groups (Hecky 512 et al., 1973). GLY enrichment observed during our study in the refractory (second) bSiO₂ phase 513 might suggest a more important role of GLY. Si-C or Si-O-C interactions are thought to protect 514 silica from dissolution until the organic matrix is removed by bacteria (Hecky et al., 1973; 515 Patrick and Holding, 1985; Bidle and Azam, 2001). The different dissolution parameters of the 516 bSiO₂ phases may be due to different associations between silica and organic compounds in 517 different sections of the frustule. Indeed, Abramson et al. (2008) observed changes in the 518 distribution of organic compounds inside the frustule that would support this argument. The very 519 slow dissolution rate constant for the second bSiO₂ phase could be due to protection by the 520 organic matrix. We suggest (1) that only a part of the $bSiO_2$ is protected by the organic coating, 521 and (2) that this protection lasts for a longer time than was previously thought. Low bacterial 522 concentration could also partially explain why the protection of the second bSiO₂ phase lasted so long, while in previous studies the protection was only temporary. The linear correlation between 523 524 FFA and the total bSiO₂ indicates that these compounds are associated with both phases of 525 bSiO₂, even though all FFA were completely degraded while 31% of the bSiO₂ still remained. Few FFA were associated with the intracellular pool of lipids. Because of their amphipathic 526

527 properties due to the carboxyl group bonded to the long carbon chain, FFA probably play a role 528 in the organization of the organic matrix involved in building the frustule (Chevallard and 529 Guenoun, 2006). The relationship between FFA degradation and dissolution of the two bSiO₂ 530 phases, and the fact that FFA were completely degraded before the total dissolution of the bSiO₂, 531 might also indicate another type of interaction, possibly adsorption of FFA on the silica surface. 532 In the present study, lipid classes and individual amino acids showed a general 533 correlation with bSiO₂ concentration (Figs. 7 and 8). The modelling experiment showed that 534 bSiO₂ and carbon pool degradation followed different patterns; they are represented in our model by different sets of equations. Moreover, in our in vitro experiment, external parameters like 535 536 temperature were constant and can not be responsible for the relation observed in Figures 7 and 8. Thus we can safely suppose that a causal correlation exists between bSiO₂ dissolution and the 537 538 amino acids or lipids degradation. The turnover of the portion of these organic compounds that is 539 correlated with dissolution of either the first or second bSiO₂ phase (P1 and P3) was very slow 540 compared to the degradation of the remaining pool of these compounds. For example, for the 541 amino acids, the turnover of Si(1)-THAA and Si(3)-THAA was slow compared to the loss of 542 THAA in general. This correlation suggests that there may exist a direct association between 543 each bSiO₂ phase and the corresponding organic compounds (Si(1)-THAA, Si(3)-THAA, Si(1)-Lip and Si(2)-Lip). Since the amounts of Si-THAA and Si-TLip related to each phase were 544 545 similar ($\sim 10-30$ %), differences between bSiO₂ dissolution rate constants stem mainly from the 546 compositions of the pools. Some membrane lipids mainly MGDG were mostly associated with 547 the first bSiO₂ phase. DPG+PG still had high concentrations at the end of P2 when most 548 degradation occurred. The second bSiO₂ phase was more strongly correlated with neutral lipid 549 alcohols (ST and ALC) but also with membrane lipids DPG+PG and PE. In the first bSiO₂ 550 phase, GLU, ASP, and LEU constituted most of the Si(1)-THAA pool and this phase contains no 551 LYS at all. In contrast in the second $bSiO_2$ phase, Si(3)-THAA had a composition similar to the diatom's intracellular THAA, except for an increase of the GLY composition. 552

553 It is not clear whether differences in organic carbon content and/or different associations 554 between $bSiO_2$ and organic carbon in each of the two $bSiO_2$ phases explains the differences between bSiO₂ dissolution rates and also between solubility equilibria. The organic matter that 555 556 makes up part of the diatom frustule helps strengthen the structure, thus increasing its resistance 557 to mechanical forces like those from grazers (Hamm et al., 2003). The role of organic 558 compounds in silica dissolution is, however, more difficult to determine. In addition to the 559 bonds between silica and the OH moiety of SER or THR (Hecky et al., 1973), silica particles are 560 linked to organic compounds by nitrogen bonds (Sumper and Kröger, 2004). Indeed, Gendron-Badou et al. (2003) noted that Si-C and Si-N bonds are present in fresh diatoms while only Si-O-561 562 Si and Si-O-R bonds are visible in fossilized diatoms. Different associations between silica and 563 organic compounds resist degradation and dissolution differently and may explain the 564 differences we observed between dissolution rate constants and solubility equilibria of the two 565 bSiO₂ phases.

566 The different dissolution rates of the two bSiO₂ phases might also be due to different 567 physico-chemical properties in the silica structure itself. In the presence of some sillafins, silica 568 precipitation of porous blocks has been observed in vitro; in contrast, when silica precipitation is 569 catalyzed by polyamines, spherical silica particles are formed (Sumper and Kröger, 2004). If the 570 first silica phase is more porous due to the presence of organic matter or due to the compounds 571 that catalyzed silica formation, the dissolution rate constant of this bSiO₂ will be higher (Van 572 Cappellen et al., 2002a). Also, when the silica structure is chemically more organized (in mineral 573 form, as opposed to amorphous, like opal), dissolution rate and solubility are lower. Gendron-574 Badou *et al.* (2003) determined that the structure of $bSiO_2$ from fossilized diatoms from 575 sediments is more organized than that of fresh diatom frustules due to condensation processes 576 that continue after they are deposited. It is also possible that fresh diatoms already have a more 577 organized phase, as suggested by the low dissolution rate (in the present study) and the low solubility equilibrium (Gallinari et al., 2002) of the second bSiO₂ phase. During dissolution, the 578

two different rate constants will cause an increase of the ratio of the more organized phase to theamorphous phase.

The difference between dissolution rate constants and between solubility equilibria could be due to chemical bonds between silica and organic matter or to structural characteristics of the 583 bSiO₂. In any case it may be closely linked to the presence of organic compounds either inside the frustule itself or during the silicification process. This conclusion emphasizes the need to link studies of carbon and silica production and recycling if we are to better understand both C and Si cycles. The following section will accentuate this conclusion by showing the reverse: the importance of bSiO₂ to carbon degradation.

588

589 **4.3 Si-C interactions and their role on carbon degradation**

590 bSiO₂ dissolution was best described by Model 2, while the turnover of each organic 591 compound investigated here was best fit by Model 1 or 3 (Fig. 5). The use of Model 3 to 592 describe compound turnover means that either (1) most organic compounds were present as two 593 pools of matter degrading one after the other or (2) degradation rate constants increased at some 594 point due to a change in environmental factors. In most cases the loss rate constant increased (POC, PON, THAA, PE, and MGDG) or decreased (other lipid classes) when the dissolution of 595 596 the first bSiO₂ phase was almost complete, from which we conclude that the first bSiO₂ phase 597 must have influenced loss of organic matter.

Most of the ALC, PE and ST present in the intracellular pool of organic compounds seems to have been protected by the first $bSiO_2$ phase, but due to the high variability of lipid class behaviour the pattern for TLip is less clear than the one for THAA. THAA degradation is consistent with the idea that Si(2)-THAA is encased within diatom frustules, and is released as soon as the first $bSiO_2$ phase is completely removed. While several amino acids increased in concentration between day 11 and day 20 (Fig. 6b), bacterial numbers peaked at day 13, which corresponds to the THAA maximum (Fig. 1 and Fig. 2). However, bacterial carbon accounted

605 for only 0.2 to 2.4 ± 0.3 % of total carbon (Fig. 3). Bacterial biomass cannot account for the 606 increase of THAA, but the increase of bacterial number could be explained by this input of labile 607 organic carbon. Ingalls et al. (2003, 2006) measured Si-THAA (THAA bound and/or within 608 bSiO₂) obtained after complete dissolution of the bSiO₂ using successive treatments with 6N HCl 609 and HF. They found that Si-THAA made up 0.7-7 % of the total THAA in diatoms from 610 plankton tows and sediment trap samples. In our study, Si-THAA was a larger portion of total 611 THAA than in the studies of Ingalls et al. (2003, 2006). We define three pools of THAA: Si(1)-612 THAA degradation is correlated to the dissolution of the first $bSiO_2$ phase; Si(2)-THAA is the 613 pool of THAA that are enclosed between the two $bSiO_2$ phases. Both of these pools were 614 protected by the first $bSiO_2$ phase. The third pool, Si(3)-THAA, is attached to the second phase 615 of the frustule; due to the low dissolution rate constant of this phase, it was protected for longer 616 time.

617 The composition of Si(2)-THAA is dominated by GLY and LYS, which are major 618 components of silaffins (Table 2). As part of the silicification process (Hildebrand, 2003; 619 Sumper and Kröger, 2004), these proteins are present in the silica deposition vesicles (SDV) that 620 become part of the diatom wall at the end of frustule formation (Martin-Jézéquel et al., 2000). 621 Thus the Si(2)-THAA pool may be assembled during the silicification process, and may result 622 from the integration of the SDV's into the frustule. Similarity between most of the substitution 623 times ts listed in Table 4 and the release time of Si(2)-THAA suggests that k_2 represents the 624 degradation rate constant of this portion of THAA. After being released, Si(2)-THAA turned over very fast (k_2 in Table 4). Most of these amino acids were completely dissolved or degraded 625 626 shortly after the total dissolution of the first bSiO₂ phase, which suggests that this pool of THAA 627 is not directly bound to $bSiO_2$. The high loss rate constant of Si(2)-THAA may suggest that this 628 pool of THAA was dissolved (chemical mechanism) rather than degraded (biological 629 mechanism) as soon as it was exposed. Another possible explanation to this fast turn over rate is 630 a change in degradation mechanism. During degradation, amino acids are released from protein

631 by enzymatic cleavage at the end of the proteins (exopeptidase) or in the middle of the 632 polypeptide chain (endopeptidase). If the proteins are opened up during the silicification process, exopeptidases could act at both ends of the protein and degradation would be faster. 633 634 Due to their high turnover rate Si(2)-THAA may have been dissolved during the strong HCl treatment or degraded before analysis in the study of Ingalls et al. (2003; 2006). 635 636 Accordingly, the quantification of Si-THAA made by Ingalls et al. (2003) might have only 637 targeted THAA bound to the second $bSiO_2$ phase of the frustule (Si(3)-THAA). The 638 composition of Si-THAA in Ingalls et al. (2003) is similar to our Si(3)-THAA except that LYS is 639 more abundant in Si(3)-THAA and GLY slightly less abundant (Table 2). The low solubility of 640 the second bSiO₂ phase is consistent with the increase in Si-THAA/THAA with depth in their 641 study.

642 The bSiO₂ protected from degradation the organic matter that was directly associated 643 with the frustule (Si(1)-THAA, Si(3)-THAA and Si-Lip). The first bSiO₂ phase also protected 644 the lipids (some DPG+PG) and amino acids (Si(2)-THAA) trapped inside the frustule possibly 645 between the 2 phases, as shown by the correlation curve (Figs. 7 and 8). Moreover, the sudden 646 increase in POC degradation rate (Fig. 1) after release of the trapped material is not associated with an increase in bacterial concentration (Figs. 2 and 3) but with the end of the dissolution of 647 648 the first $bSiO_2$ phase. It is hardly a coincidence that the end of dissolution of the first $bSiO_2$ 649 phase occurred exactly when the degradation of the POC, PON and THAA increased. The 650 mechanisms behind this observation are not clear yet; the dissolution of this phase probably gave 651 bacteria better access to the internal carbon of the cell, possibly because the integrity of the 652 frustule can not be maintained without the presence of the first bSiO₂ phase. This could happen 653 through increasing pores size as sometimes shown by pictures of diatom frustules during 654 dissolution, or because the box-shaped frustule opens at the end of the first bSiO₂ phase 655 dissolution as observed after sexual phases (Crawford 1995), both triggering cell lysis. In any 656 case we can safely conclude that the first bSiO₂ phase of the diatom frustule also protects most of

the intracellular carbon; at the end of the dissolution of the first $bSiO_2$ phase 69% of the POC was still not degraded. Due to the very low dissolution rate of the second $bSiO_2$ phase, the associated organic compounds (Si(3)-THAA and Si(2)-Lip) might be protected for a long period of time; they would even be preserved in the sediment with the $bSiO_2$.

The presence of organic compounds inside the frustule and/or during silicification
determines the solubility of the different parts of the frustule. Moreover, intracellular carbon and
Si-bound organic compounds may be protected by at least some part of the frustule. These
reverse interactions prove that carbon and silica production and recycling must be studied in
parallel if we want to improve our understanding of mechanisms driving both POC and bSiO₂
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860 Figure legends

Fig. 1. Changes in the relative concentrations of POC (closed black diamonds), bSiO₂
(open diamonds), THAA (closed grey squares) and TLip (closed grey triangles) during the
degradation of *S. marinoi* in the dark at 20°C during the first 25 days of the experiment. The
concentrations relative to initial values are on a logarithmic scale.

- Fig. 2. Change in total bacterial concentration over time during the 102-day degradation
 experiment.
- 867 Fig. 3. Change in algal TOC (open circles), DOC (open diamonds), POC (closed squares)
 868 and bacterial carbon (closed circles) with time during the degradation experiment.

Fig. 4. Change in the concentration of (a) 6 of the 8 lipid classes in μ mol C L⁻¹ relative to TLip concentrations in μ mol C L⁻¹ (b) 9 of the 14 individual amino acids in μ mol AA L⁻¹

relative to THAA in units of μ mol AA L⁻¹, over time during the first 25 days of the 102-days

872 degradation experiment. Note that due to the low number of C atoms in GLY, the GLY peak in

873 μ mol AA L⁻¹ is more visible than if using μ mol C L⁻¹.

Fig. 5. Model comparisons (a) Dissolution of $bSiO_2$. In this experiment, $bSiO_2$ is the only

875 compound whose loss is best represented by Model 2. $\Delta \log(L)$ between Model 2 and Model 3

is 13. Period 1 is the period of time corresponding to the dissolution of 85% of the first $bSiO_2$

phase and 10% of the second $bSiO_2$ phase. During period 2, the last 15% of the first $bSiO_2$

dissolved and 10% more of the second bSiO₂ phase dissolved. In Period 3 only the second

 $bSiO_2$ phase dissolved as less than 1.5% of the first $bSiO_2$ phase remained. (b) The curve depicts

the loss of POC (or any organic compound) with a dissolution or degradation rate constant that

increases with the *ts*. Model 1 fits the curve using $C_0 = 8297 \mu \text{mol C L}^{-1}$ and $k = 0.047 \text{ d}^{-1}$ with a

likelihood log(L) = 98.7. Model 2 fits the model with the same likelihood using the same

883 parameters, i.e. $C_1 + C_2 = 8297 \text{ } \mu \text{mol C } \text{L}^{-1}$, $k_1 = k_2 = 0.047 \text{ } \text{d}^{-1}$. Model 3 give the best fit

884 $(\log(L) = 174.7)$ using $C_0 = 7614 \ \mu \text{mol C L}^{-1}$, $ts = 10 \ d$, $k_1 = 0.025 \ d^{-1}$ and $k_2 = 0.082 \ d^{-1}$. This 885 example clearly shows that only Model 3 can depict accurately the loss when the rate constant 886 increases at some point in the experiment. Moreover, Model 2 never gives a better likelihood 887 than Model 1 under these conditions.

Fig. 6. Change in organic compound concentrations with time during the degradation of *S. marinoi*. For clarity, 7 of the 8 lipid class concentrations in μ mol Cl_{ip} L⁻¹ over time are shown (5a) and only 10 of the 14 amino acids in μ mol AA L⁻¹ over time (5b). Results depicted are only for the first 25 days. Note that THAA concentrations are in μ mol AA L⁻¹.

892 Fig. 7. Si-TLip interactions during the degradation of the S. marinoi. (a) Correlation 893 between the concentrations of the dissolved bSiO₂ from the first phase relative to its initial 894 concentration (estimated by the model) and each lipid class relative to its initial concentration. 895 (b) Correlation between concentration of the dissolved $bSiO_2$ from the second phase relative to 896 its initial concentration (estimated by the model) and each lipid class relative to its initial 897 concentration. Period 1 is the period of time corresponding to the dissolution of 85% of the first 898 $bSiO_2$ phase and 10% of the second $bSiO_2$ phase. During period 2, the last 15% of the first 899 bSiO₂ dissolved, and 10% more of the second bSiO₂ phase dissolved. In Period 3 only the 900 second $bSiO_2$ phase dissolved as less than 1.5% of the first $bSiO_2$ phase remained.

901Fig. 8. Si-THAA interactions during the degradation of S. marinoi. (a) Correlation902between the concentration of the dissolved $bSiO_2$ from the first phase relative to its initial903concentration (estimated by the model) and individual THAA concentrations relative to their904initial concentrations. (b) Correlation between concentration of the dissolved $bSiO_2$ from the905second phase relative to its initial concentration (estimated by the model) and individual THAA906concentrations relative to their initial concentrations. The periods shown are the same as in Fig.9077.

Table 1: List of abbreviations used in the text to refer to organic and inorganic compounds

910	measured	during	the degr	adation	experimen	t.
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Biogenic Silica	bSiO ₂	Total Organic Carbon	TOC
Silicic acid	dSi	Particulate organic carbon	POC
Particulate organic nitrogen	PON	Dissolved Organic carbon	DOC
Total Lipid classes	TLip	Total Hydrolyzed Amino Acids	THAA
Alcohols	ALC	Alanine	ALA
Di- and monophosphatidyl	DPG+PG	Arginine	ARG
glycerides		Aspartic acid	ASP
Free fatty acids	FFA	Glutamic acid	GLU
Monogalactosyldiglycerides	MGDG	Glycine	GLY
Monoglycerides	MG	Histidine	HIS
Phosphatidylethanolamines	PE	Isoleucine	ILE
Pigments	PIG	Leucine	LEU
Sterols	ST	Lysine	LYS
		Methionine	MET
		Phenylalanine	PHE
		Serine	SER
		Threonine	THR
		Tyrosine	TYR
		Valine	VAL
		γ-Aminobutyric acid	GABA

912 Table 2: Concentration in μ mol C L ⁻¹	¹ and composition in mole% of the different pools of THAA
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913 in S. marinoi. The THAA row shows the initial concentration and composition of THAA before

- 914 dissolution began and does not include Si-THAA. Si(1)-THAA is the pool associated with the
- 915 first bSiO₂ phase, Si(2)-THAA is the pool of THAA enclosed between the bSiO₂ phases and
- 916 Si(3)-THAA is the pool associated with the second $bSiO_2$ phase.

THAA	ALA	ARG	ASP	GLU	GLY	HIS	ILE	LEU	LYS	PHE	SER	THR	TYR	VAL	TOT
THAA	105	223	318	379	139	59	174	317	187	260	128	136	170	174	2769
% THAA / THAAtot	4%	8%	11%	14%	5%	2%	6%	11%	7%	9%	5%	5%	6%	6%	
Si(1)-THAA	20	31	122	143	12	12	48	92	0	63	16	29	58	45	691
% Si(1)-THAA/ Si(1)-THAAtot	3%	5%	19%	22%	2%	2%	7%	14%	0%	10%	2%	5%	9%	7%	
Si(2)-THAA	9	0	52	37	92	17	21	19	96	21	0	12	18	21	332
% Si(2)-THAA/ Si(2)-THAAtot	2%	0%	13%	9%	23%	4%	5%	5%	24%	5%	0%	3%	5%	5%	
Si(3)-THAA	20	28	51	64	72	20	26	43	53	35	29	30	26	26	523
% Si(3)-THAA/ Si(3)-THAAtot	4%	5%	10%	12%	14%	4%	5%	8%	10%	7%	5%	6%	5%	5%	

- 918 Table 3: Concentration in μ mol C L⁻¹ and composition in mole% of the different lipid class in S.
- 919 *marinoi*. The lipid row shows the initial concentration and composition of lipid before
- 920 dissolution began and does not include Si-lipid. Si(1)-Lip is the pool associated with the first
- 921 $bSiO_2$ phase, Si(2)-Lip is the pool of THAA associated with the second $bSiO_2$ phase.

Lipid class	ALC	DPG+PG	FFA	MG	MGDG	PE	PIG	ST	total
Lipids	30	98	366	104	189	44	604	40	1475
lipid class/ total lipid	2%	7%	25%	7%	13%	3%	41%	3%	
Si(1)-Lip	0	0-36	180-201	15-25	0	1 - 19	NC	1-11	386 - 469
Si(1)-Lip/ tot Si(1)-Lip	0%	0-8%	43-47%	4-5%	41-49%	0-4%		0-2%	
Si(2)-Lip	0-4	40-69	3	0	0	7-13	NC	9-20	125-175
Si(2)-Lip/ tot Si(2)-Lip	0-2%	32-39%	39-55%	0%	0%	6-7%		7-11%	

922 NC: No correlation

924	Table 4: Kinetic parameters and likelihood $(log(L))$ calculated by the 3 models. C_0 is the initial
925	concentration of the compound. C_1 and C_2 are the initial concentrations of the two phases (for
926	$bSiO_2$) or the two pools (organic compounds). k is the degradation/dissolution rate constant
927	calculated with Model 1. k_1 and k_2 are degradation/dissolution rate constants of C_1 and C_2 in
928	Model 2, respectively, or used before and after the substitution time <i>ts</i> in Model 3, respectively.
929	The last column indicates which model has been chosen in this study to determine the
930	degradation/dissolution rate constant and the initial concentration of each compound. For

	Model 1			Model 2					Model 3					Best
	Co	k	log(L)	C_1	C_2	<i>k</i> ₁	<i>k</i> 2	$\log(L)$	C ₀	ts	<i>k</i> ₁	<i>k</i> 2	$\log(L)$	fit
PON	1499	0.050	<i>98.5</i>	26	1473	0.051	0.051	98.5	1364	12	0.030	0.094	135.6	III
POC	8297	0.047	<i>98.7</i>	6884	1412	0.047	0.047	<i>98.7</i>	7614	10	0.025	0.082	174.7	III
bSiO ₂	591	0.030	70.7	209	462	0.268	0.016	109.0	658	6	0.063	0.019	97.7	II
TLip	1232	0.033	27.9	704	724	0.014	0.100	43.5	1420	17	0.048	0.010	44.9	III
ALC	44	0.080	9.2	46	5	0.122	0.001	10.1	52	19	0.099	0.001	13.5	III
FFA	297	0.104	21.8	139	207	0.068	0.214	23.1	330	15	0.125	0.077	25.0	Ι
MG	97	0.045	24.5	21	85	0.497	0.033	25.0	107	5	0.094	0.036	25.5	Ι
MGDG	199	0.145	9.0	93	107	0.145	0.145	9.0	189	2	0.078	0.520	19.6	III
PE	54	0.097	22.3	51	3	0.097	0.097	22.3	42	6	0.016	0.119	26.6	III
PG+DPG	69	0.011	17.9	55	13	0.011	0.011	17.9	65	16	0.001	0.054	19.0	Ι
PIG	641	0.012	29.7	32	610	0.012	0.012	29.7	627	5	0.000	0.016	30.4	Ι
ST	41	0.031	30.0	11	30	0.031	0.031	30.0	35	13	0.001	0.080	38.3	III
THAA	3599	0.080	32.9	1903	1696	0.080	0.080	32.9	3137	13	0.054	0.139	42.8	III
ALA	132	0.070	42.3	31	102	0.073	0.073	42.3	117	12	0.048	0.120	54.1	III
ARG	339	0.113	26.6	39	300	0.113	0.113	26.6	243	9	0.024	0.175	54.3	III
ASP	419	0.100	24.5	312	107	0.100	0.100	24.5	324	5	0.001	0.121	29.1	III
GLU	454	0.080	39.2	23	431	0.080	0.080	<i>39.2</i>	405	13	0.065	0.147	50.5	III
GLY	172	0.045	24.9	67	105	0.045	0.045	24.9	147	16	0.015	0.182	39.3	III
HIS	73	0.063	28.9	44	29	0.063	0.063	28.9	65	16	0.039	0.167	38.6	III
ILE	223	0.082	34.4	210	13	0.082	0.082	34.4	192	12	0.050	0.137	44.8	III
LEU	413	0.090	34.0	117	299	0.091	0.091	34.0	348	12	0.050	0.150	48.2	III
LYS	268	0.068	18.1	181	88	0.068	0.068	18.1	223	14	0.031	0.170	26.1	III
PHE	336	0.083	31.6	332	4	0.083	0.083	31.6	280	12	0.040	0.148	45.8	III
SER	162	0.075	42.6	21	140	0.075	0.075	42.6	135	5	0.007	0.090	<i>49.7</i>	III
THR	165	0.064	43.1	17	149	0.064	0.064	43.1	151	13	0.045	0.110	51.1	III
TYR	185	0.068	<i>48.3</i>	157	28	0.068	0.068	<i>48.3</i>	187	21	0.070	0.012	49.3	Ι
VAL	219	0.076	37.9	179	39	0.076	0.076	37.9	194	12	0.050	0.124	46.5	III
932														

931 compounds, see abbreviations table (Table 1).









945 Fig. 3





950 Figure 5:













- 962 Fig. 8:

