# A first look at dissolved Ge isotopes in marine sediments

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#### 2 ABSTRACT

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The removal of chemical species from seawater during the precipitation of authigenic minerals is difficult to constrain but may play a major role in the global biogeochemical cycles of some elements, including silicon (Si) and germanium (Ge). Here, we present Ge/Si,  $\delta^{74}$ Ge, and supporting chemical data of pore waters and core incubations at three continental margin sites in California and the Gulf of Mexico. We used these data to partition Ge release and uptake by the various allogenic (delivered via sedimentation) and authigenic (formed *in situ*) phases in these sediments.

About half of the pore water Ge ( $\delta^{74}$ Ge<sub>pw</sub> = 1.3-2.4‰) is supplied by biogenic silica dissolution ( $\delta^{74}$ Ge ~ 3‰), with the other half contributed by lithogenic particulates ( $\delta^{74}$ Ge ~ 0.6‰). The highest Ge/Si (~ 3µmol/mol) and lowest  $\delta^{74}$ Ge (1.3-1.9‰) are observed at the Fe redox horizon, suggesting a supply from detrital Ge-rich Fe oxides. The precipitation of authigenic phases (most likely aluminosilicate clays) in deeper sediments preferentially incorporates Ge over Si, resulting in low pore water Ge/Si (~ 0.3µmol/mol). The lack of corresponding  $\delta^{74}$ Ge<sub>pw</sub> trend indicates negligible Ge isotope fractionation during this process.

Ge fluxes measured via core incubations were variable and appeared strongly controlled by Fe redox behavior near the sediment-water interface. In some cases, reductive Fe oxide dissolution appeared to enhance the benthic Ge flux by over 100% and released fractionated low  $\delta^{74}$ Ge of ~ -0.7‰, resulting in overall benthic  $\delta^{74}$ Ge<sub>inc</sub> between -0.2 and 3.6‰, depending on Fe oxide contribution to Ge flux. We estimate that detrital inputs supply  $19 \pm 15\%$  of total dissolved Ge to continental margin pore fluids globally, resulting in an average dissolved  $\delta^{74}$ Ge of  $2.5 \pm 0.4\%$ . Assuming 10-60% of pore water Ge is captured by the authigenic aluminosilicate sink, the dissolved Ge flux to the ocean derived from terrigenous inputs should be roughly 2.4-6.4 Mmol/y, much higher than previously estimated. Our results imply that authigenic Si burial in continental margins should be in the range of 0.9-8.1 Tmol/y (best estimate 3.1 Tmol/y), sufficient to close the global marine Si budget.

Keywords: germanium, biogenic silica, authigenesis, Fe oxides, isotope fractionation, San Pedro Basin, Santa
 Monica Basin, Gulf of Mexico

#### **1 INTRODUCTION**

Silicon (Si) is a major constituent of Earth's silicate crust that is released to solution during 31 rock weathering and delivered to the ocean by rivers, where it sustains the productivity of 32 marine biosilicifying organisms (Tréguer and De La Rocha, 2013). Oceanic Si concentrations 33 also determine the rate of authigenic clay formation in sediments, which was recently 34 proposed to influence global climate through its effects on the marine carbonate balance 35 (Mackenzie and Garrels, 1966). Silicon isotope ratio ( ${}^{30}$ Si/ ${}^{28}$ Si, expressed as  $\delta^{30}$ Si) and the 36 germanium-to-silicon (Ge/Si) ratio have been developed as proxies tracing the biogeochemical 37 cycling of Si (e.g., Froelich et al., 1985; Murnane and Stallard, 1990; De La Rocha et al., 38 2000; Ziegler et al., 2005). Recently, the stable isotope composition of Ge ( $^{74}$ Ge/ $^{70}$ Ge 39 ratio, expressed as  $\delta^{74}$ Ge) has been measured in the major Earth surface reservoirs and 40 proposed as an additional proxy helping to constrain the coupled biogeochemical cycles of 41 Ge and Si (Rouxel et al., 2006; Escoube et al., 2012, 2015; Rouxel and Luais, 2017; Baronas 42 et al., 2017, 2018). Germanium is a useful tracer of the Si cycle due to its prevalence in 43 silicate rocks and the similar atomic properties and chemical behavior of the two elements 44 (Burton et al., 1959; Froelich and Andreae, 1981; Rouxel and Luais, 2017). The ranges of 45 Ge/Si and especially  $\delta^{74}$ Ge signatures in silicate rocks are relatively narrow (Mortlock and 46 Froelich, 1987; Escoube et al., 2012; Rouxel and Luais, 2017). During continental weathering 47 processes, the precipitation of secondary weathering products such as Al- and Fe-oxides and 48 aluminosilicate clays preferentially incorporates Ge relative to Si (Froelich et al., 1992; Kurtz 49 et al., 2002) and light Ge isotopes preferentially relative to heavy ones, resulting in dissolved 50 river composition that has lower Ge/Si but higher  $\delta^{74}$ Ge relative to silicate rocks (Baronas 51 et al., 2018). 52

The dissolved seawater Ge/Si and  $\delta^{74}$ Ge composition is primarily controlled by a balance of riverine and hydrothermal inputs and biogenic and authigenic outputs (Elderfield and Schultz, 1996; Hammond et al., 2000; Escoube et al., 2015; Baronas et al., 2017). Although biological Ge/Si fractionation is variable, most diatoms appear to discriminate against Ge only when dissolved Si is depleted below ~10 µmol/L (Sutton et al., 2010; Baronas et al., 2016). Ge/Si ratios are relatively invariable in deep seawater and in diatom biogenic silica (bSi) because nearly all dissolved Ge and Si upwelled to the photic zone is exported by diatom growth and settling, and most of this export dissolves congruently in deep water (Froelich et al., 1985,
1992; Ellwood and Maher, 2003; Baronas et al., 2016; Guillermic et al., 2017).

The Ge isotopic composition appears to be unfractionated during Ge incorporation into 62 diatom bSi, although some fractionation may occur during Ge incorporation into the organic 63 cellular material (Mantoura, 2006; Rouxel and Luais, 2017; Guillermic et al., 2017). Because 64 most organic matter is rapidly remineralized in the upper water column, this secondary 65 fractionation should have little effect on sedimentary  $\delta^{74}$ Ge dynamics. In contrast, all siliceous 66 sponge spicules analyzed to date exhibit Ge/Si and  $\delta^{74}$ Ge signatures significantly lower than 67 seawater, indicating strong vital effects (Ellwood et al., 2006, 2010; Rouxel et al., 2006; 68 Guillermic et al., 2017). The magnitude of sponge bSi production and burial is not well known, 69 although it is likely small relative to diatoms (Van Cappellen, 2003), limiting the influence of 70 biological fractionation on global seawater Ge/Si and  $\delta^{74}$ Ge composition. 71

While the burial of diatom bSi has little or no effect on the seawater  $\delta^{74}$ Ge composition, 72 a large portion of bSi reaching the seafloor dissolves at or just below the water-sediment 73 interface, releasing Ge and Si into marine pore waters. A range of complex diagenetic 74 reactions take place in marine sediments, including the precipitation of various authigenic clay 75 minerals (e.g., Aller, 2013). These authigenic minerals incorporate Ge and Si, affecting pore 76 water Ge/Si (Hammond et al., 2000; King et al., 2000; Baronas et al., 2016),  $\delta^{30}$ Si (Ehlert 77 et al., 2016), and possibly  $\delta^{74}$ Ge signatures. Recent studies have suggested that authigenic 78 ("non-opal") burial plays a major role in the global marine cycles of Ge and Si (Baronas et al., 79 2016, 2017; Rahman et al., 2016, 2017). Indeed, variations in authigenic Ge burial fluxes are 80 likely responsible for the large seawater Ge/Si fluctuations over glacial-interglacial cycles 81 (Mortlock et al., 1991; Hammond et al., 2004b; Baronas et al., 2016). A fully constrained 82 global Ge isotope budget and unambiguous interpretation of past  $\delta^{74}$ Ge variations in seawater 83 therefore require knowledge of the chemical and isotopic Ge behavior during marine sediment 84 authigenesis. 85

Here, we present dissolved Ge/Si,  $\delta^{74}$ Ge, and various other solute concentration data at three different continental margin sites. Sediment pore water signatures were used to track the potential isotopic fractionation with progressing sediment diagenesis and authigenesis, while sediment core incubation data were used to constrain the net effect of these processes on the benthic flux. Ultimately, Ge isotopes have allowed us to put preliminary quantitative constraints on Ge partitioning between different sources and sinks in the studied sediments.

#### 2 STUDY SITES

San Pedro and Santa Monica basins are located in the Southern California continental margin, ~20 and ~40 km offshore from Los Angeles, respectively. A detailed description of the geological and oceanographic setting of the Southern California Bight can be found elsewhere (Gorsline, 1992; Hickey, 1992). San Pedro basin has an area of 819 km<sup>2</sup> and is 900 m deep. The Santa Monica basin is 2225 km<sup>2</sup> large and 925 m deep. Both basins are silled

below 725 m depth, restricting the circulation of bottom water, with periodic flushing every 97 few years (Berelson, 1991). Organic matter remineralization renders the restricted bottom 98 water suboxic in both basins, typically  $<9\mu$ M O<sub>2</sub> in San Pedro basin and  $<4\mu$ M O<sub>2</sub> in Santa 99 Monica basin, resulting in reducing sediment pore waters and limited macrofaunal activity in 100 the benthos (Berelson et al., 1987; Leslie et al., 1990; Gorsline, 1992). In San Pedro basin, 101 minor sediment irrigation and bioturbation may occur at times, whereas Santa Monica basin 102 sediments are finely laminated, indicating lack of macrofaunal activity. Material is supplied to 103 both basins primarily through particle infall from the overlying water column and through 104 nepheloid plume transport (e.g., Collins et al., 2011). The sediments are comprised of 1-4%105 bSi, 8-10 % CaCO<sub>3</sub>, and ~4 % organic matter, with the remainder being lithogenic particles 106 (Gorsline, 1992; Cheng et al., 2009). 107

Additional data are presented from the continental Gulf of Mexico shelf, close to the Mississippi River delta. Ge cycling in the area has been described in detail by Baronas et al. (2016), showing that Ge/Si ratios in the region are generally elevated due to contamination from coal ash (Froelich et al., 1985) and that authigenic Ge accounts for about 50% of total Ge burial in the Gulf of Mexico shelf sediments. In this study, new  $\delta^{74}$ Ge data from the two regions are presented and analyzed in the context of previously published pore water concentrations and core incubation results.

## 3 METHODS

#### 115 **3.1 Sample collection**

San Pedro basin (SPB) sediment cores were collected on 2014-09-04 aboard the R/V 116 Yellowfin at the San Pedro Ocean Timeseries (SPOT) study site. SPOT is the site of a 117 multi-year monthly water column sampling campaign (https://dornsife.usc.edu/spot/). Santa 118 Monica basin (SMB) cores were collected on 2012-03-08 aboard the R/V Yellowfin in the 119 central part of the basin (Table 1). The cores were kept on ice before being placed in a 5 120  $^{\circ}$ C cold room within 8 hours from retrieval. For half of the cores, the overlying water was 121 siphoned off while avoiding disturbance of the sediment surface. Then, pore waters were 122 sampled using Rhizons (0.2 µm membrane; Rhizosphere Research Products, The Netherlands). 123 The suction was applied at all depth horizons simultaneously to minimize vertical pore water 124 advection during sampling. The pore waters were collected for up to 24h. All samples were 125 acidified to 0.1 vol% with Teflon-distilled conc. HNO3 inside the sampling syringe. Another 126 set of cores with overlying water were incubated for several days as described below. 127

Additional water column samples were collected at SPB and in the Atlantic Ocean in 2014 and previously (Table 1). The SBP samples were collected in Niskin bottles on a CTD rosette aboard R/V Yellowfin, filtered through a 0.2 µm pore size filter within 10 hours from retrieval, and acidified prior to analyses. The Atlantic Ocean samples were collected at the Bermuda Atlantic Time Series (BATS) station during the June 2008 GEOTRACES intercalibration cruise aboard the R/V Knorr. Further details are given in Baronas et al. (2017). The Gulf of Mexico (GoMex) samples were collected in August 2011 during cruise EN-494 aboard the R/V Endeavor. Sediment cores were collected using a multi-corer and seawater samples using Niskin bottles. Pore waters were collected via sectioning under N<sub>2</sub> atmosphere and centrifugation. All samples were filtered through 0.2  $\mu$ m membrane and acidified prior to analysis. A detailed description of the GoMex methods is given in Baronas et al. (2016).

#### 139 3.2 Core incubations

Core incubations were carried out using the method described by Hammond et al. (2004a). 140 Briefly, sediment cores with 1-1.5 L of overlying water were capped, placed in a 5 °C cold 141 room and the water slowly stirred (20-30 rpm) using a suspended magnetic stir bar. During 142 incubations, 10-20 mL of overlying water was periodically sampled for Ge and Si concentration 143 analyses using a plastic syringe, while the piston was advanced to keep air out. Although care 144 145 was taken to avoid air contact during core retrieval and incubations, in some cases a bubble of air was introduced to the overlying water, either during capping or via leakage around piston 146 o-rings during sampling. The samples were immediately filtered through a 0.2  $\mu$ m membrane 147 and acidified prior to analysis. After the incubation was completed, the remaining overlying 148 water was collected via siphoning, filtered through a  $0.2 \,\mu m$  membrane and acidified prior to 149  $\delta^{74}$ Ge analyses. 150

#### 151 3.3 Solute concentration analyses

Silicic acid and ammonia concentrations were measured using standard colorimetric 152 techniques (Mullin and Riley, 1955; Bower and Holm-Hansen, 1980) with a precision better 153 than 5%. Ammonia analyses were carried out within 24 hours of sample collection to minimize 154 degassing. Iron and manganese concentrations were analyzed by ICP-MS on a Thermo 155 Scientific Element2 with a precision of  $\sim 10\%$ . Sulfate concentrations were measured on 156 a Metrohm Ion Chromatograph. Germanium concentrations were measured using isotope-157 dilution-hydride-generation-ICP-MS on a Thermo Scientific Element2, using the method 158 developed by Mortlock and Froelich (1996) and modified by Baronas et al. (2016). 159

#### 160 3.4 Ge isotope analyses

Ge co-precipitation. Filtered and acidified samples of pore water from similar depths of 161 multiple cores were combined to obtain larger composite samples required for  $\delta^{74}$ Ge analyses. 162 Ge concentrations of individual aliquots were analyzed beforehand to ensure that all cores 163 had similar Ge and Si concentration profiles. Pore water, incubation, and seawater samples 164 165 ranging from 100 mL to 9 L and containing 4-13 ng of Ge were then spiked with a Ge isotope double spike  $(^{73}\text{Ge}/^{70}\text{Ge} \approx 1)$ , previously calibrated and used by Escoube et al. (2012, 2015); 166 Baronas et al. (2018)) in a spike/sample Ge mass ratio of 1-2 and a purified FeCl<sub>3</sub> solution 167 to obtain a Fe concentration of  $\sim 0.2$  mmol/L. The samples were well mixed, and allowed to 168 equilibrate for at least 16h. Next, Fe(OH)<sub>3</sub> flock was precipitated by bubbling pure NH<sub>3</sub> gas 169 through the sample until the solution reached a pH of 8-10. The flock was collected by settling 170 171 and centrifugation, redissolved in 2 mL concentrated Teflon-distilled HNO<sub>3</sub> and diluted to 10

mL with ultrapure (18 M $\Omega$ ) H<sub>2</sub>O. The samples were then dried down, redissolved in 1 mL concentrated Optima-grade HF and diluted to 30 mL with ultrapure H<sub>2</sub>O to obtain a final 1M HF solution. They were then purified through anion exchange columns as described below. The procedural blank was determined by processing spiked ultrapure H<sub>2</sub>O and ranged from 0.01 to 0.3 ng Ge.

Anion-exchange chromatographic separation. A procedure adapted from Rouxel et al. 177 (2006) and described in detail by Guillermic et al. (2017) and Baronas et al. (2018) was used. 178 All reagents used were either in-house Teflon-distilled or Optima-grade. A 10 mL column was 179 loaded with 1.8 mL (wet volume) of BioRad AG1-X8 resin, washed with 10 mL of 3M HNO<sub>3</sub>, 180 0.28M HNO<sub>3</sub>, and ultrapure H<sub>2</sub>O in sequence and conditioned with 5 mL 1M HF. Samples in 181 1M HF solution as prepared above were centrifuged to separate insoluble fluorides and 10-29 182 mL of the solution was carefully added to columns. The presence or the amount of insoluble 183 fluorides at this stage did not appear to affect the final Ge recovery. The remaining matrix 184 was eluted with 5 mL of 1M HF followed by 3 mL of ultrapure H<sub>2</sub>O, leaving fluorinated Ge 185 retained on the column. Ge was then eluted with 10 mL 0.28M HNO<sub>3</sub>. If required, the solution 186 was dried down and redissolved in a smaller volume of 0.28M HNO<sub>3</sub> to obtain the 0.5-10 187 ppb Ge concentration required for isotope measurements. Each column was reused 4-5 times, 188 except when retention of DOC from the previous sample was observed based on the color, in 189 which case the resin was replaced. Ge blanks from reused resin were below detection limit. 190

191 Ge recovery ranged from 20 to 90%, with one sample being as low as 8%. Incomplete 192 recovery was most likely due to variable Ge co-precipitation efficiency with Fe(OH)<sub>3</sub> (resulting 193 from variable precipitation rates, final pH, and variable sample matrices, especially DOC 194 concentrations in pore waters), as well as some loss during co-precipitate recovery from the 195 solution. Importantly, incomplete recovery did not affect the measured  $\delta^{74}$ Ge values, as all 196 samples were double-spiked prior to sample preparation.

Seawater contains relatively high concentrations of methylated Ge, which does not participate in the inorganic Ge cycle (Lewis et al., 1985, 1988, 1989). It is therefore important to separate the inorganic and the methylated species prior to  $\delta^{74}$ Ge analysis. Baronas et al. (2017) achieved this via chromatographic separation of the the methylated and inorganic Ge hydrides. In this study, separation was achieved during both Fe co-precipitation and anion column chromatography, and is confirmed by the agreement of seawater  $\delta^{74}$ Ge determined via both methods (Table 1).

**HG-MC-ICP-MS.** Ge isotope analyses were performed on a Thermo Neptune multi-204 205 collector ICP-MS at Ifremer in Brest, France, using the method of Rouxel et al. (2006) as adapted by Escoube et al. (2015); Guillermic et al. (2017); Baronas et al. (2018). Sample 206 solutions of 0.5-10 ppb natural Ge in 0.28M HNO<sub>3</sub> were introduced into an online hydride 207 generation system (CETAC HGX-200) at a rate of 150  $\mu$ L/min where they were mixed with 208 209 0.25 M NaBH<sub>4</sub> solution (in 1.5 M NaOH) introduced at an equal rate. The dissolved Ge(OH)<sub>4</sub> species were reduced to gaseous GeH<sub>4</sub> and transported into the ICP-MS torch using Ar 210 carrier gas. The Neptune MC-ICP-MS was operated in low mass resolution mode, measuring 211

<sup>70</sup>Ge, <sup>72</sup>Ge, <sup>73</sup>Ge, and <sup>74</sup>Ge in L2, C, H1 and H2 cups, respectively. In addition, L4, L3, 212 L1 and H4 cups were also monitored for <sup>68</sup>Zn (possible interference as <sup>70</sup>Zn), <sup>69</sup>Ga, <sup>71</sup>Ga 213 (possible interferences at m/z 70), and <sup>77</sup>Se (possible interference as <sup>74</sup>Se), respectively. 214 No interferences were detected in any of the runs. The samples were bracketed using a 215 NIST-3120a standard solution that had a total Ge concentration generally within  $\sim 20\%$  of 216 the bracketed sample, and was double-spiked to have a spike/sample ratio within  $\sim 20\%$  of 217 the bracketed sample. Each sample or standard run consisted of 6 measurement blocks each 218 lasting 2 min (30 cycles of 4 s each), and in most cases 4-5 blocks displaying the most 219 stable signal were retained. Therefore, each measurement consisted of 8-10 min of counting 220 statistics at signal intensities ranging from 0.4 to 6 V (4-60 pA) at <sup>74</sup>Ge (depending on Ge 221 concentration in sample solution and instrument tuning). The  $\delta^{74}$ Ge values were calculated 222 for each block using the double-spike data reduction routine of Siebert et al. (2001) and 223 are reported in % as  $^{74}$ Ge/ $^{70}$ Ge sample ratio normalized to the average  $^{74}$ Ge/ $^{70}$ Ge ratio of 224 bracketing measurements of Ge isotope standard NIST 3120a. This method also yields Ge 225 concentration values based on the measured spike/sample ratio. Several different reference 226 materials were analyzed multiple times, interspersed with the samples, and all agreed well with 227 values previously reported by Baronas et al. (2018). The measurement uncertainty is reported 228 as the internal  $2\sigma$  standard error of the used sample blocks, or  $2\sigma$  standard deviation of all 229 NIST 3120a bracketing standard measurements within a given analytical session, whichever is 230 higher. 231

# 4 **RESULTS**

All of the Gulf of Mexico (GoMex) data, with the exception of  $\delta^{74}$ Ge, were previously reported and discussed by Baronas et al. (2016). This section therefore focuses on the newly acquired data from San Pedro and Santa Monica basins.

#### 235 4.1 Seawater

Seawater data are reported in Table 1. Several samples were previously analyzed for  $\delta^{74}$ Ge by Baronas et al. (2017) and re-analyzed in this study, yielding identical values within analytical uncertainty.

The Ge/Si value determined for SPB bottom seawater (885m depth) was 0.74 µmol/mol, 239 close to the global ocean value of 0.76 µmol/mol (Froelich et al., 1985; Sutton et al., 2010). 240 Core top water collected up to 8h after core retrieval exhibited slightly elevated Ge and Si 241 concentrations, with Ge/Si up to 0.87 µmol/mol. These were likely affected by benthic flux and 242 disturbance during transport. SPB bottom seawater  $\delta^{74}$ Ge ranged from 2.9 to 3.2‰, slightly 243 lighter than the 3.4-3.5% determined higher in the water column (Table 1; Baronas et al. 244 (2017)), which is indistinguishable from deep seawater  $\delta^{74}$ Ge values in other oceanic basins 245 (Guillermic et al., 2017). 246

Sample	Station	Date	Lat. (°)	Long. (°)	Depth (m)	Ge (pmol/L)	Si (µmol/L)	Ge/Si (µmol/mol)	δ <sup>74</sup> Ge (‰)	Fe (µmol/L)	Mn (nmol/L)
Gulf of Mexico											
GOM CTD-45-7	Sta. 9	2011-08-15	28.97	-90.40	2.3	43	10	4.28	$2.79 \pm 0.50^{*}$		
GOM CTD-6 (30m)	Sta. 1	2011-08-01	28.59	-90.54	30	45	25	1.84	2.13 ± 0.15		
GOM-CTD-32-3	Sta. G	2011-08-09	26.28	-92.02	2121	17	25	0.70	3.13 ± 0.28		
San Pedro Basin											
SPOT SSW	SPOT	2014-09-10	33.55	-118.40	2	1.6	1.1	1.43			
UP-18	SPOT	2014-04-23	33.55	-118.40	50	7.0	8.0	0.88	3.41 ± 0.25		
SPOT 500m	SPOT		33.55	-118.40	500	55	74	0.74	$3.48 \pm 0.35^{*}$		
SPOT 885m	SPOT	2014-09-10	33.55	-118.40	885	78	105	0.74	2.97 ± 0.20		
MC-2D OLW	SPOT	2014-09-04	33.55	-118.40	885	91	114	0.79	3.06 ± 0.22	1.15	75
MC-3D OLW	SPOT	2014-09-04	33.55	-118.40	885	82	111	0.74	3.23 ± 0.28	0.69	42
MC-5C OLW	SPOT	2014-09-04	33.55	-118.40	885	109	125	0.87	2.90 ± 0.32	0.11	46
MC-1D OLW	SPOT	2014-09-04	33.55	-118.40	885	92	118	0.78		1.71	105
MC-2C OLW	SPOT	2014-09-04	33.55	-118.40	885	103	120	0.85			
MC-2A OLW	SPOT	2014-09-04	33.55	-118.40	885	89	116	0.77			
Atlantic											
GPrI-19	BATS	2011-06-08	31.67	-64.17	1000	12	13	0.90	3.04 ± 0.28		
GDI-30,31 / 32	BATS	2011-06-08	31.67	-64.17	2000	24	17	1.36	3.68 ± 0.28		
GPrl-3	BATS	2011-06-08	31.67	-64.17	3500	27	28	0.97	$3.03 \pm 0.50^{*}$		

**Table 1.** Study site locations and details of seawater samples. Dashes indicate no analyses were done.

\* Previously published in Baronas et al. (2017).

#### 247 4.2 Pore waters

248 The pore water solute concentrations were similar in each individual core. Ge concentrations ranged from 90 to 1200 pmol/L (Supplementary Data) and showed a maximum at 2-3 cm 249 depth, decreasing monotonically below (Fig. 1). In contrast, Si concentrations were lowest at 250 the sediment-water interface and increased from  $\sim 250 \,\mu mol/L$  to  $\sim 400 \,\mu mol/L$  within the top 251 10 cm, with a continued slow increase to  $\sim$ 500  $\mu$ mol/L by 35 cm depth (Fig. 1). Pore water 252 Ge/Si ranged from a high of 2-3 µmol/mol at 2-3 cm depth to a low of 0.2-0.4 µmol/mol at 253 the bottom of the cores.  $\delta^{74}$ Ge values in pore water were lighter than in the water column, 254 ranging from 1.3 to 2.3% in SPB and SMB and from 1.9 to 2.4% in GoMex sediments (Table 255 2). Pore water  $\delta^{74}$ Ge showed little variation with depth. 256

Fe, Mn, NH<sub>3</sub>, and SO<sub>4</sub> concentrations in SPB pore waters are reported in Fig. 1 and 257 Supplementary Data. Fe concentrations ranged from 1 to 300 µmol/L, with a maximum at 258 2-3 cm depth. Mn and NH<sub>3</sub> concentrations ranged from 0 to 600 nmol/L and from 20 to 400 259 µmol/L, respectively, and both increased monotonically with depth. Fe and Mn concentrations 260 reported here are in good agreement with recently published profiles from ancillary cores 261 262 collected during this cruise (Monteverde et al., 2018). Sulfate concentrations were in 24-26 mmol/L range and appeared to slightly decrease with depth in SPB sediments (previously 263 published by Monteverde et al. (2018)). 264



**Figure 1.** Pore water profiles of San Pedro Basin (SPB) sediments collected from five cores. Bottom seawater and initial core top overlying water data is plotted above the sediment-water interface as blue squares. a) Ge isotope composition of composite samples, vertical bars show the depth range of combined samples. Three different composites were measured at the 1-5 cm horizon (see Table 2). Note the reverse x-axis; b) Ge/Si of composite samples as in (a); also shown are individual sample Ge/Si ratios as small gray triangles. Note the log scale of x-axis. The green band shows the estimated  $\delta^{74}$ Ge and the measured Ge/Si composition of bSi (Baronas et al., 2016), equivalent to seawater. The gray bands show the typical composition of lithogenic silicates (Escoube et al., 2012; Rouxel and Luais, 2017); c) Ge and Si concentrations; d) NH<sub>3</sub>, Fe, and Mn concentrations; e) Sulfate concentrations in salinity.

#### 265 4.3 Core incubations

Core incubations were performed with core-top water present for up to six days, to constrain 266 net effect of sediment diagenetic processes on the benthic Ge and Si fluxes and the  $\delta^{74}$ Ge 267 composition of the benthic flux. Throughout the incubations, Ge concentrations increased 268 from 70-80 to 80-120 pmol/L in SPB overlying water and from 90-100 to 120-150 pmol/L in 269 SMB overlying water. Si concentrations increased from ~100 to 130-150 µmol/L in SPB and 270 from ~130 to 150-190 µmol/L in SMB (Fig. 3; Supplementary Data. Table 3 summarizes the 271 chemical and isotopic composition of the post-incubation overlying water and the calculated 272 benthic fluxes. The Ge benthic fluxes were calculated to range from  $\sim 0$  to 1.8 nmol m<sup>-2</sup> d<sup>-1</sup> 273 at SPB and from 0.9 to 1.5 nmol  $m^{-2} d^{-1}$  at SMB. The Si benthic fluxes ranged from 0.7 to 274 1.6 mmol m<sup>-2</sup> d<sup>-1</sup> at SPB and from 0.5 to 1.4 mmol m<sup>-2</sup> d<sup>-1</sup> at SMB. As a result, the benthic 275 flux Ge/Si ratios exhibited a wide range at both sites (0.03-1.1 µmol/mol at SPB and 0.9-1.8 276  $\mu$ mol/mol at SMB). The post-incubation  $\delta^{74}$ Ge was in the 2.9-3.5% range, similar to the 277 initial overlying composition (2.9-3.2‰), with the exception of one SPB incubation (MC-3A; 278  $2.07 \pm 0.71\%$ ) that showed the highest Ge flux. 279

In the Gulf of Mexico, the  $\delta^{74}$ Ge composition of two post-incubation overlying water samples was determined in addition to previous data reported by Baronas et al. (2017) (Table



**Figure 2.** Pore water profiles of Gulf of Mexico (GoMex) sediments collected from three cores at Sta. 2 (Table 2; Baronas et al. (2016)). GoMex seawater data is plotted at an arbitrary depth above the sediment-water interface. a) Ge isotope composition of composite samples, vertical bars show the depth range of combined samples. Note the reverse x-axis; b) Ge/Si of composite samples as in (a). Note the log scale of x-axis. The green band shows the estimated  $\delta^{74}$ Ge and Ge/Si composition of bSi in GoMex shelf area affected by the Mississippi River (pink dotted line; see text). The gray bands show the typical composition of lithogenic silicates (Escoube et al., 2012; Rouxel and Luais, 2017); c) Ge and Si concentrations, including additional Si data from a separate core MC-5; d) NH<sub>3</sub>, Fe, and Mn concentrations in MC-5; e) Sulfate concentrations in MC-5, corrected for salinity. All the data except for  $\delta^{74}$ Ge were previously published in Baronas et al. (2016).

3). At Station 1, which is located close to the Mississippi River delta, post-incubation  $\delta^{74}$ Ge was determined to be 2.4‰ (bottom seawater at this site was 2.1‰), in agreement with data from other cores of Baronas et al. (2017). At Station 2, which was located several hundred kilometers away from the river delta,  $\delta^{74}$ Ge was determined to be 3.4‰, similar to the deep seawater in the Gulf of Mexico and other oceanic basins (Table 1; (Guillermic et al., 2017)).

## **5 DISCUSSION**

#### 287 5.1 Pore waters

Broadly, the  $\delta^{74}$ Ge and Ge/Si composition of fluids within and above sediments is controlled 288 by 1) mixing of various solute sources (dissolution of bSi vs. lithogenic particles, as well as 289 trapped/diffusing bottom water); and 2) solute removal via precipitation of authigenic phases 290 (Fe oxides and authigenic aluminosilicates), and associated elemental or isotopic fractionation. 291 Pore water  $\delta^{74}$ Ge measured in SPB and SMB sediments was 1-2‰ lighter than the expected 292 composition of the dissolving diatom bSi, which is likely to be similar to the overlying 293 seawater (Fig. 1a). Therefore, the low  $\delta^{74}$ Ge<sub>pw</sub> composition must reflect either a significant 294 contribution from an isotopically lighter source, or fractionation during Ge incorporation into 295 precipitating authigenic phases. Previous studies have shown that various lithogenic silicates, 296

Cores	Station	Depth (cm)	Ge (pmol/L)	Si (µmol/L)	Ge/Si (µmol/mol)	δ <sup>74</sup> Ge (‰)
San Pedro Basin						
MC-2A, MC-2C, MC-2D	SPOT	1-5	568	347	1.64	1.33 ± 0.15
MC-1D, MC-3D	SPOT	1-5	595	298	1.99	1.58 ± 0.16
MC-5C (C-1, C-2)	SPOT	1-5	595	460	1.29	1.94 ± 0.31
MC-1D - MC-5C (all cores)	SPOT	6-10	346	391	0.89	$2.12 \pm 0.13$
MC-1D - MC-5C (all cores)	SPOT	11-20	251	430	0.58	$2.00 \pm 0.22$
MC-1D - MC-5C (all cores)	SPOT	21-37	147	455	0.32	2.11 ± 0.21
MC-2A	SPOT	0	531	297	1.78	
MC-2A	SPOT	8	380	356	1.07	
MC-2A	SPOT	15	214	440	0.49	
MC-2A	SPOT	33	111	525	0.21	
<u>Santa Monica Basin</u>						
D1-S2, D1-S4, D3-S1, D3-S3	SMB	4-6	379	366	1.03	$2.30 \pm 0.28$
Gulf of Mexico *						
MC-6C, MC6-D, MC6-E	Sta. 2	0-3	386	313	1.23	$2.29 \pm 0.42$
MC-6C, MC6-D, MC6-E	Sta. 2	3-6	639	462	1.38	$2.44 \pm 0.16$
MC-6C, MC6-D, MC6-E	Sta. 2	6-10	517	521	0.99	$2.12 \pm 0.32$
MC-6C, MC6-D, MC6-E	Sta. 2	10-15	326	520	0.63	$2.03 \pm 0.32$
MC-6C, MC6-D, MC6-E	Sta. 2	15-20	385	435	0.88	1.94 ± 0.32

**Table 2.** Ge and Si chemistry of composite pore waters. Individual high resolution sample measurements are given in Supplementary Data.

\* All Gulf of Mexico data except  $\delta^{74}$ Ge were previously published in Baronas et al. (2016).

including marine sediments, exhibit a narrow  $\delta^{74}$ Ge range of 0.4-0.8‰ (Rouxel and Luais, 2017). Secondary terrestrial weathering products, for example Fe oxides, are known to be enriched in Ge (Kurtz et al., 2002) and to preferentially incorporate light Ge isotopes during formation (Pokrovsky et al., 2014). The  $\delta^{74}$ Ge signature of lithogenic material delivered to the sediments should therefore be close to (potentially slightly lower than) the Bulk Silicate Earth value of 0.58 ± 0.21‰ (Rouxel and Luais, 2017).

In SPB pore waters, the lowest  $\delta^{74}$ Ge<sub>pw</sub> values are found in the 1-5 cm depth horizon, 303 coinciding with the highest Ge/Si ratios and highest Fe concentrations in pore waters (Fig. 304 1). Pore water composition in this zone is therefore most likely controlled by mixing of 305 three sources: dissolution of bSi (Ge/Si = 0.7  $\mu$ mol/mol;  $\delta^{74}$ Ge = 3-3.5‰), dissolution or 306 desorption of Ge from lithogenic particles (Ge/Si  $\ge$  1.5 µmol/mol;  $\delta^{74}$ Ge  $\le$  0.6‰), and the 307 reductive dissolution of authigenic Fe oxides that precipitate at the sediment-water interface 308 (Fig. 1). Germanium is well known to adsorb or co-precipitate with Fe (oxy)hydroxides 309 (e.g., Pokrovsky et al., 2006). Indeed, Fe (oxy)hydroxide (abbreviated as FeOx from here 310 on) co-precipitation is used to pre-concentrate Ge from dissolved samples during sample 311 preparation (see Methods). This scenario might involve non-steady state Fe redox dynamics 312 313 in these sediments, which is possible, given the occasional re-oxygenation of bottom waters (Berelson, 1991) and the possible introduction of oxygen during core retrieval (see Section 314 5.2). For this reason, further pore water discussion focuses on deeper (>10 cm) pore waters, 315

**Table 3.** Summary of core incubation data, showing the final post-incubation composition of the overlying water, as well as the calculated Ge and Si fluxes. Individual time-series data for San Pedro and Santa Monica basins are given in Supplementary Data. Some of the Gulf of Mexico data (except for  $\delta^{74}$ Ge) were previously reported by Baronas et al. (2016). Ge and Si flux uncertainties are 20%.

Coros	Station	Ge	Si	Ge/Si	$\mathbf{x}^{74}$ <b>C</b> $(0())^{\dagger}$	Ge flux	Si flux	Ge/Si flux
Cores	Station	(pmol/L)	(µmol/L)	(µmol/mol)	o Ge (‰)	(nmol m <sup>-2</sup> d <sup>-1</sup> )	(mmol m <sup>-2</sup> d <sup>-1</sup> )	(µmol/mol)
San Pedro Bas	in	fir	nal composi	tion after incu	bation			
MC-3A	SPOT	118	142	0.83	2.07 ± 0.71	1.78	1.59	1.12
MC-4C <sup>#</sup>	SPOT	110	149	0.74	2.93 ± 0.28	0.50	0.75	0.67
MC-5A <sup>#</sup>	SPOT	90	156	0.58	3.53 ± 0.28	0.02	0.73	0.03
MC-5B	SPOT	80	144	0.55	3.14 ± 0.28	0.43	0.77	0.56
MC-5D <sup>#</sup>	SPOT	98	129	0.76	3.38 ± 0.18	0.41	0.98	0.41
<u>Santa Monica E</u>	Basin							
D3-S2	SMB	145	154	0.946	3.26 ± 0.28	0.91	0.50	1.81
D4-S1	SMB	147	165	0.894		1.12	0.75	1.49
D4-S4	SMB	147	170	0.864	3.42 ± 0.28	0.87	0.70	1.24
D5-S1	SMB	151	189	0.799	2.95 ± 0.28	1.48	1.43	1.04
D5-S4	SMB	116	163	0.711		0.90	1.00	0.90
Gulf of Mexico								
MC-2A *	Sta. 1	111	70	1.59	2.71 ± 0.50	2.17	2.09	1.04
MC-2B	Sta. 1	99	58	1.70	2.43 ± 0.28	2.28	1.65	1.38
MC-3A *	Sta. 1	113	83	1.35	2.36 ± 0.50	4.09	2.65	1.54
MC-3B *	Sta. 1	113	89	1.27	2.18 ± 0.50	3.55	2.43	1.46
MC-6A, MC-6B	Sta. 2	58	73	0.79	3.44 ± 0.24		1.97	

\* Previously published in Baronas et al. (2017). Water depth is 30-38 m at Sta. 1 and 22-24 m at Sta. 2.

<sup>†</sup> Uncertainty reported as ± 2σ (sample replicate or bracketing standard reproducibility, whichever is higher).

<sup>#</sup> A notable air bubble was introduced during recovery or incubation for these cores.

which are unaffected by Fe redox dynamics. We will return to the coupling of Ge to Fe redox in the discussion of core incubation data (Section 5.2).

#### 318 5.1.1 Effect of authigenic clay precipitation on Ge isotope composition

319 At about 15 cm depth, pore water Ge/Si values decrease below 0.7 µmol/mol (Fig. 1b), indicating that dissolved Ge is being removed from pore water via the precipitation of 320 authigenic (non-opal) phases, which has previously been observed in SPB (Hammond et al., 321 2000; King et al., 2000; McManus et al., 2003). Although the stoichiometry and the mineralogy 322 of the precipitating phases is poorly constrained, the tight coupling of Ge and Fe concentrations 323 (Fig. 1c-d) suggests it could be Fe-rich aluminosilicate clays. Such clays have been shown to 324 form rapidly in continental margin environments, including GoMex sediments (Michalopoulos 325 and Aller, 1995; Presti and Michalopoulos, 2008; Rahman et al., 2017). The formation of 326 aluminosilicate clays is also indicated by the asymptotic Si concentrations of  $\sim 400-500 \,\mu mol/L$ 327 (Table 2, Figs. 1 and 2), which are significantly lower than the solubility of bSi, the latter 328



**Figure 3.** Incubation results of cores from San Pedro and Santa Monica basins (SPB and SMB, respectively): a & b) The increase of Ge and Si concentrations in the incubated overlying water. These slopes, once corrected for water loss via sampling (see Supplementary Data), were used to determine the benthic Ge and Si fluxes reported in Table 3; c & d) the change in the Ge/Si ratio of the overlying water. The thick dashed line shows the Ge/Si of biogenic silica in the SPB and SMB sediments (0.7  $\mu$ mol/mol; Baronas et al. (2016)); e & f) plots showing Ge vs. Si concentration over the course of the incubations. These slopes were used to calculate the benthic flux Ge/Si ratio (Table 3). The thick dashed line shows Ge/Si of biogenic silica (0.7  $\mu$ mol/mol).

ranging from 600 to 1000 µmol/L, depending on bSi age and the degree of surface passivation
by authigenic coatings (e.g., McManus et al., 1995).

331 Isotopically light Ge is also readily incorporated into sulfide minerals that precipitate

in hydrothermal settings, e.g. sphalerite (ZnS) (Luais, 2007, 2012; Escoube et al., 2012;

Belissont et al., 2014; Escoube et al., 2015; Meng et al., 2015) and possibly during sulfate

reduction in marine sediments (Murnane et al., 1989), although the latter has not been clearly

- 335 demonstrated. Sulfate concentration in SPB pore waters decreases slightly with depth (Fig. 1e),
- indicating likely sulfate reduction. Leslie et al. (1990) previously argued that sulfate reduction

and pyrite precipitation rates decrease with depth in SPB sediments. Therefore, if pyrite 337 precipitation fractionates  $\delta^{74}$ Ge, we would expect to see a strong  $\delta^{74}$ Ge<sub>pw</sub> gradient, which 338 is not observed (Fig. 1a). Overall, authigenic Ge sequestration appears to be independent of 339 sulfide precipitation in this and other continental margins (Hammond et al., 2000; Baronas 340 et al., 2016) and it is therefore unlikely to be a major factor regulating dissolved Ge dynamics 341 in these sediments. We do acknowledge the possibility that the dissolution of several phases 342 (bSi, FeOx) and the precipitation of several others (FeS, FeOx, authigenic silicates) could 343 fractionate  $\delta^{74}$ Ge<sub>pw</sub> in opposite directions, theoretically resulting in the relatively constant 344  $\delta^{74}$ Ge<sub>pw</sub> of about 2‰. Assessing this possibility is, however, outside of the scope of this 345 paper and we therefore argue that the simplest explanation – no  $\delta^{74}$ Ge fractionation during 346 authigenic phase precipitation – is most likely to be the correct one. 347

The minimum estimate of dissolved Ge removed via authigenesis can be calculated from the change in pore water Ge/Si composition. Assuming that all of pore water Ge below 30 cm depth is derived from diatom bSi dissolution (Ge/Si =  $0.7 \mu$ mol/mol), at least ~50-70 % of dissolved Ge has to be precipitated to achieve the measured Ge/Si of 0.2-0.3 µmol/mol, in agreement with previous studies at these sites (Hammond et al., 2000; Baronas et al., 2016). At the same time, the  $\delta^{74}$ Ge<sub>pw</sub> composition remains constant with depth, indicating that any isotopic fractionation during this process is small.

Although the geochemical setting is more complicated and there are less data available, the 355 same conclusion can be drawn from the GoMex pore waters (Fig. 2). Here,  $\delta^{74}$ Ge<sub>pw</sub> are also 356 relatively constant in the 1.9-2.4% range, whereas Ge/Si varies between 1.4 and 0.6 µmol/mol, 357 generally decreasing with depth. The Mississippi River supplies a large amount of Ge and Si 358 to the studied area, partly due to contamination by anthropogenic activity (Mississippi Ge/Si = 359 1.6  $\mu$ mol/mol;  $\delta^{74}$ Ge = 2.0‰; Baronas et al. (2017)). Its discharge and possibly chemical and 360 isotopic composition varies temporally, as a result variably affecting the elemental and isotopic 361 composition of diatom bSi that is supplied to and dissolves in the GoMex shelf sediments. 362 Despite the uncertainty of the bSi end-member, the large decrease in pore water Ge/Si can 363 only be explained by authigenic Ge precipitation (Baronas et al., 2016). Yet, similarly to SPB 364 sediments, no detectable variation of  $\delta^{74}$ Ge<sub>pw</sub> with depth is observed at GoMex (Fig. 2a). 365

Finally, the lack of  $\delta^{74}$ Ge fractionation by authigenic clays is supported by previous 366 measurements of an authigenic glauconite mineral by Rouxel et al. (2006). The iron-rich 367 glauconite clay analyzed (reference material GL-O) was enriched in Ge, with Ge/Si of 7.3 368 µmol/mol, supporting the role of authigenic aluminosilicate clays as the deep non-opal Ge 369 sink in marine sediments. Despite this enrichment, it exhibited  $\delta^{74}$ Ge of 2.44 ± 0.15‰ 370 (re-normalized to NIST 3120a standard), close to the value of  $\delta^{74}$ Ge<sub>auth</sub> calculated here (Table 371 4). It must be noted that the age of this glauconite was estimated to be 95 Ma (Kapusta et al., 372 1997), and  $\delta^{74}$ Ge<sub>sw</sub> at that time may have been significantly different from the current value, 373 potentially invalidating the above interpretation. 374

#### 375 5.2 Core incubations

Benthic flux measurements reflect the composition of solutes diffusing into the overlying water column, integrating the net effect of all sedimentary processes. Core incubation data can therefore be used to more quantitatively assess the various sources of dissolved Ge to pore waters and the potential isotopic fractionation associated with authigenic phase formation.

380 5.2.1 Ge and Si benthic fluxes

The benthic Si fluxes were similar at SPB and SMB sites and individual core experiments agreed well with each other at a given site (relative standard deviation of ~40%; Table 3 and Fig. 3). In contrast, Ge fluxes differed significantly between the two sites and were especially variable within SPB, spanning two orders of magnitude and resulting in a wide range of observed benthic flux Ge/Si ratios. We propose that Ge fluxes in these experiments were affected by variable redox conditions in the sediments that were perturbed during retrieval, transportation, and incubation.

Santa Monica basin bottom waters and sediments are known to be consistently anoxic, exhibiting high benthic Fe flux and dissolved  $Fe^{2+}$  up to (or nearly up to) the sediment-water interface (McManus et al., 1997; Elrod, 2004; Severmann et al., 2010). The Ge/Si flux at SMB was significantly higher than the 0.7 µmol/mol of dissolving bSi in all cases (Fig. 3, Table 3), consistent with the dissolution of Ge-enriched Fe oxides.

At SPB, benthic Ge fluxes and Ge/Si ratios varied widely, the latter ranging from ~0.03 to 393 394  $1.12 \,\mu$ mol/mol (Fig. 3, Table 3). Considering that all cores were collected in close proximity (cores MC-5A, -5B, and -5D were collected during a single multi-corer deployment), this 395 variability is unlikely to be caused by spatial heterogeneity in surface sediment composition. 396 The wide range of observed fluxes therefore probably results from the perturbation of bottom 397 redox conditions during core recovery and incubation. It is unavoidable that some oxygen is 398 introduced during sampling. As a result, dissolved  $Fe^{2+}$  in the overlying water and the surficial 399 pore water can be oxidized, capturing a portion (or all, in the case of MC-5A) of the potential 400 Ge benthic flux. Orange, most likely FeOx, flock was observed on sediment surface of most 401 cores. In addition, core MC-5A, which had negligible benthic Ge flux, also had a large burrow 402 at the sediment water interface and seemed the most disturbed during recovery, including air 403 bubbles trapped in the core liner. In support of this hypothesis, the benthic dissolved Fe flux at 404 SPB was previously determined by in-situ incubations and water column measurements to 405 often be 1-2 orders of magnitude lower than that based on pore water Fe gradients, suggesting 406 that a large fraction of dissolved Fe is captured near the water-sediment interface (Elrod, 2004; 407 Severmann et al., 2010; John et al., 2012). 408

In summary, the core incubations performed on SPB and SMB sediments together represent a range of bottom redox conditions. Such sampling-induced perturbation provides an independent test of any  $\delta^{74}$ Ge fractionation potentially associated with Fe oxide precipitation in marine sediments. Below, we perform some simple mass balance modeling to determine benthic flux <sup>413</sup>  $\delta^{74}$ Ge signature of each core and the fraction of Ge released or sequestered by the various <sup>414</sup> solid phases.

415 5.2.2 Modeling Ge isotope mass balance during core incubations

Although the large amount of seawater needed for  $\delta^{74}$ Ge analyses prevents the collection of  $\delta^{74}$ Ge time-series data during core incubations, the final post-incubation  $\delta^{74}$ Ge signature ( $\delta^{74}$ Ge<sub>final</sub>) can be used to assess the isotopic composition of Ge flux that has been affected by non-opal phases. To do this, a simple mass balance model was built, partitioning Ge amounts dissolved and sequestered by the various phases, and their associated Ge isotope signatures (Fig. 4).



**Figure 4.** Schematic representation of the Ge isotope mass balance model used to interpret core incubation data. The main model-calculated parameters (namely  $\delta^{74}$ Ge<sub>inc</sub>, F<sub>auth</sub>, and F<sub>FeOx</sub>) are shown in green. Other values are either measured or calculated as described in text. Back-diffusion of Ge from overlying water into the sediments is assumed to be small and is ignored.

For each incubated sediment core, the amount of Ge  $(n_i, in mol)$  in the overlying postincubation water is the sum of Ge initially present prior to the incubation and the net added via the benthic flux during incubation:

$$n_{\text{final}} = n_{\text{initial}} + n_{\text{inc}} \tag{1}$$

$$n_{\text{initial}} = [Ge]_{\text{initial}} V_{\text{initial}}$$
(2)

$$n_{\rm inc} = F_{\rm inc} t A \tag{3}$$

where  $[Ge]_{initial}$  and  $V_{initial}$  are the measured Ge concentration and volume of the preincubation overlying water.  $F_{inc}$  is the measured Ge incubation flux given in Table 3 (in mol m<sup>-2</sup> time<sup>-1</sup>), t is incubation time and A is the core sediment surface area (in m<sup>2</sup>). The Ge isotope composition in the final post-incubation overlying water is then

$$\delta_{\text{final}}^{74} = \frac{n_{\text{initial}} \,\delta^{74} \text{Ge}_{\text{initial}} + n_{\text{inc}} \,\delta^{74} \text{Ge}_{\text{inc}}}{n_{\text{final}}} \tag{4}$$

Having measured all the other parameters, Eq. 4 can be used to calculate  $\delta^{74}\text{Ge}_{\text{inc}},$  the 429 isotopic composition of Ge released into the overlying water during the incubation. Using a 430 number of additional observations and assumptions (see below), we can quantify the different 431 processes controlling the dissolved Ge isotope mass balance in these sediments: 1) Ge release 432 from bSi and lithogenic particle dissolution (F<sub>lith</sub> and F<sub>bSi</sub>, respectively), followed by 2) the 433 removal or addition of Ge by Fe redox reactions near the sediment-water interface (F<sub>FeOx</sub>), 434 and 3) continued removal of Ge by other authigenic phases in deeper sediments (Fauth), as 435 observed in the pore water profiles (Fig. 4). A Monte Carlo approach (running the model 436 1 million times for each incubated core) was utilized to assess the full range of uncertainty 437 associated with all the input parameters, yielding probability distributions of the calculated 438 values. A Monte Carlo approach (running the model 1 million times for each incubated core) 439 was utilized to assess the full range of uncertainty associated with all the input parameters, 440 yielding probability distributions of the calculated values. 441

A detailed description of model equations and input parameters is given in Section 5.2.2.1. The model results for each individual core are given in Supplementary Materials. Here we provide a summary of the overall results.

The combined model results of all eight incubated SPB and SMB cores are given in Table 4 445 and Fig. 5. Based on the measured benthic Si flux (combined with diatom Ge/Si ratio) and 446 lithogenic FeOx input flux (Leslie et al., 1990) we show that the biogenic and lithogenic Ge 447 input fluxes are roughly similar (Fig. 5e). Using the  $\Delta^{74}$ Ge<sub>FeOx-diss</sub> fractionation factors 448 reported by Pokrovsky et al. (2014) along with the  $\delta^{74}$ Ge<sub>pw</sub> data reported here, we calculate 449 that on average, authigenic Ge sequestration takes up between 8 and 41 % of the total allogenic 450 supply (f<sub>auth</sub> between -0.08 and -0.41; Table 4, Fig. 5c), whereas iron (oxy)hydroxides in 451 most cases sequester a similar proportion of Ge ( $f_{FeOx}$  down to -0.5) but in some cases can 452 also release authigenic Ge during dissimilatory Fe reduction ( $f_{FeOx} > 0$ ). This variability in 453

 $f_{FeOx}$  is consistent with the qualitative discussion of variable Fe redox dynamics and their perturbation during recovery and is discussed in more detail below.

**Table 4.** Summary of core incubation model results, combined for all studied cores. The ranges given are the 25-75th percentiles of all Monte Carlo calculated values. SPB core MC-5A is not included in the reported combined  $\delta^{74}$ Ge<sub>inc</sub> statistics due to very low benthic Ge flux and the resulting very high uncertainty of the calculated  $\delta^{74}$ Ge<sub>inc</sub>.

Parameter	Median	25-75 <sup>th</sup> percentile
f <sub>lith</sub>	0.53	(0.45 – 0.59)
<b>f</b> <sub>bSi</sub>	0.47	(0.41 – 0.55)
<b>f</b> <sub>auth</sub>	-0.20	(-0.410.08)
f <sub>FeOx</sub>	-0.20	(-0.50 – 0.10)
<b>f</b> <sub>released</sub>	0.54	(0.31 – 0.85)
δ <sup>74</sup> Ge <sub>supply</sub> (‰)	1.88	(1.69 – 2.11)
δ <sup>74</sup> Ge <sub>auth</sub> (‰)	2.13	(1.92 – 2.34)
δ <sup>74</sup> Ge <sub>FeOx</sub> (‰)	-0.66	(-1.32 – -0.18)
δ <sup>74</sup> Ge <sub>inc</sub> (‰)	1.59	(0.80 – 2.39)

For SMB cores, the fraction of allogenic Ge input that is captured by authigenic 456 phases is relatively constant, ranging from 14 to 29% (median f<sub>released</sub> between 0.71-0.86; 457 Supplementary Materials). Our modeling indicates that authigenic phases play a relatively 458 minor role in SMB sediments ( $-0.19 < f_{auth} < -0.1$  and  $-0.17 < f_{FeOx} < 0.07$ ). While SPB 459 sediments exhibited similar Ge sequestration by the deep (possibly clay) authigenic phase 460  $(-0.23 < f_{auth} < -0.16$  with the exception of MC-3A where median  $f_{auth} = -0.81$ ), dissolved 461 Ge uptake or release by authigenic Fe oxides was much more variable ( $-0.75 < f_{FeOx} < 0.80$ ) 462 at SPB, likely caused by perturbations and variable oxygen introduction during core retrieval 463 and incubation (Table 3). As a result, the net fraction of dissolved Ge released back into 464 overlying waters, varied widely among SPB core incubations ( $f_{released} = 0.02 - 0.98$ ). Below 465 we give a detailed description of how these different estimates were calculated. 466

#### 467 5.2.2.1 Detailed core incubation model description

Having measured the pre- and post-incubation overlying water  $\delta^{74}$ Ge, Eq. 4 allows the calculation of  $\delta^{74}$ Ge<sub>inc</sub>, or the Ge isotope composition of the incubation flux. Here we describe the subsequent calculations used to partition this flux between the different Ge sources and sinks in the incubated sediments. All input parameter values and their uncertainty ranges are given in Supplementary Material.

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473 First, the Ge benthic flux expected from bSi dissolution was calculated simply as

$$F_{bSi} = F_{inc}^{Si} Ge/Si_{bSi}$$
(5)



**Figure 5.** Probability distributions of the fraction of Ge released or consumed by different phases (relative to the allogenic supply from bSi and lithogenic inputs) during San Pedro and Santa Monica basin core incubation experiments, and their isotopic compositions: a)  $f_{released}$  values <1 indicate net Ge sequestration in sediments, and values >1 indicate additional Ge release from authigenic Fe oxide reduction; b) the combined probability distribution of  $\delta^{74}$ Ge released during all eight core incubations; c) fraction of Ge consumed (<0) or released (>0) from the deep authigenic (possibly clay) phase ( $f_{auth}$ ) and authigenic Fe oxides ( $f_{FeOx}$ ); d) the calculated Ge isotope composition of the two authigenic phases; e) the calculated fraction of allogenic Ge supply by lithogenic particles and bSi; f) the assigned Ge isotope composition of lithogenic and bSi phases, and the calculated composition of total allogenic Ge input ( $\delta^{74}$ Ge<sub>supply</sub>.)

which assumes that the amount of Si captured by sediment authigenesis is negligible (e.g. less than 10%). This assumption is validated by the good agreement between the average annual measured  $F_{inc}^{Si}$  flux of  $1.2 \pm 0.3$  mmol m<sup>-2</sup> d<sup>-1</sup> during 2004-2006 (Hammond et al., 477 unpub. data) and the bSi rain to the seafloor of  $1.3 \pm 0.2 \text{ mmol m}^{-2} \text{ d}^{-1}$  during this time, 478 measured using sediment traps (Collins et al., 2011). Ge/Si<sub>bSi</sub> is taken to be equivalent to 479 deep Ge/Si<sub>sw</sub> in the California Margin, as confirmed by a bSi-targeting weak alkaline leach of 480 the trap material of Collins et al. (2011) (Supp. Material of Baronas et al. (2016)).

The detrital Ge flux can be estimated from the external Fe (oxy)hydroxide input to SPB 481 sediments, which was previously calculated by Leslie et al. (1990) as the flux required to 482 sustain the pyrite burial flux, equal to 26  $\mu$ mol m<sup>-2</sup> d<sup>-1</sup>. Making the simplifying assumption 483 that reducible Fe oxides are the dominant lithogenic source of dissolved Ge in these sediments, 484 the detrital Ge flux can be calculated from the Ge/Fe ratio of these oxides. Given that both 485 Ge and Fe are highly immobile and mostly retained in the solid phases during weathering 486 (Baronas et al., 2018), detrital Fe oxides should have Ge/Fe that is similar to continental crust, 487 i.e. around 27  $\mu$ mol/mol (Rudnick and Gao, 2014). Using Ge/Fe = 27 ± 10  $\mu$ mol/mol yields a 488 lithogenic-derived Ge contribution to pore waters (F<sub>lith</sub>) of  $0.70 \pm 0.26$  nmol m<sup>-2</sup> d<sup>-1</sup>. The 489 isotopic composition of this detrital flux should be close to the crustal value, given that more 490 than 95% of river-delivered Ge is retained in the solid phase and that riverine suspended 491 sediment  $\delta^{74}\mbox{Ge}$  composition is for the most part indistinguishable from primary igneous rock 492 values (Baronas et al., 2018). 493

494 The allogenic supply of dissolved Ge to the sediments is therefore:

$$F_{supply} = F_{bSi} + F_{lith}$$
(6)

and its isotopic composition:

$$\delta^{74} \text{Ge}_{\text{supply}} = \frac{F_{\text{bSi}} \,\delta^{74} \text{Ge}_{\text{bSi}} + F_{\text{lith}} \,\delta^{74} \text{Ge}_{\text{lith}}}{F_{\text{supply}}} \tag{7}$$

Finally, the measured incubation flux ( $F_{inc}$ ) is affected by the sequestration or release of Ge by authigenic phases, which we here separate into iron oxides (*FeOx*; note that these are authigenic oxides that precipitate and dissolve within sediments, distinct from the detrital rain-delivered Fe oxides denoted above as *lith*) and other authigenic phases, such as authigenic clays (*auth*):

$$F_{inc} = F_{supply} + F_{FeOx} + F_{auth}$$
(8)

$$\delta^{74} Ge_{inc} F_{inc} = \delta^{74} Ge_{supply} F_{supply} + \delta^{74} Ge_{FeOx} F_{FeOx} + \delta^{74} Ge_{auth} F_{auth}$$
(9)

where positive F values indicate Ge release into dissolved phase, and negative F valuesindicate Ge sequestration into solid phase. The isotopic composition of FeOx is simply

$$\delta^{74} \text{Ge}_{\text{FeOx}} = \delta^{74} \text{Ge}_{\text{supply}} + \Delta^{74} \text{Ge}_{\text{FeOx-diss}}$$
(10)

where  $\Delta^{74}\text{Ge}_{\text{FeOx-diss}}$  is assigned the experimentally determined range between -1.6 and -4.6% (Pokrovsky et al., 2014). In the case of downcore authigenic phases, no experimental or theoretical  $\Delta^{74}\text{Ge}_{\text{auth-diss}}$  values are available. However, the lack of pore water  $\delta^{74}\text{Ge}_{pw}$ gradient observed at SPB (Fig. 1) limits  $\Delta^{74}\text{Ge}_{\text{auth-diss}}$  to  $0 \pm 0.3\%$ . The isotopic composition of this downcore authigenic phase is then

$$\delta^{74} \text{Ge}_{\text{auth}} = \delta^{74} \text{Ge}_{\text{pw}} + \Delta^{74} \text{Ge}_{\text{auth-diss}}$$
(11)

Using  $\delta^{74}$ Ge<sub>pw</sub> instead of  $\delta^{74}$ Ge<sub>supply</sub> for the initial value is necessary in this case, to allow for the potential influence of isotopically distinct Ge released from reductive FeOx dissolution higher in the sediment column.

#### Equations 8-11 can be combined to solve for $F_{FeOx}$ and $F_{auth}$ .

The fractions of Ge supplied or consumed by different phases are defined as fractions of the allogenic supply:

$$f_{lith} = F_{lith} / F_{supply}$$
(12)

$$f_{bSi} = F_{bSi}/F_{supply}$$
(13)

$$f_{FeOx} = F_{FeOx} / F_{supply}$$
(14)

$$f_{auth} = F_{auth} / F_{supply}$$
(15)

$$f_{\text{released}} = F_{\text{inc}} / F_{\text{supply}}$$
(16)

Given that  $n_{inc}$  for the incubated cores ranged between 1 and 35% of  $n_{final}$ , and the 514 similar values between  $\delta^{74}$ Ge<sub>initial</sub> and  $\delta^{74}$ Ge<sub>final</sub>, the calculated  $\delta^{74}$ Ge<sub>inc</sub> can be sensitive to 515 analytical and experimental uncertainty. We used a Monte Carlo approach to fully assess this 516 uncertainty and to further deconvolve the various factors influencing  $\delta^{74}$ Ge<sub>inc</sub>. The above set 517 of calculations is performed a large number of times (n = 1,000,000), each time randomly 518 selecting from within the uncertainty range of each given parameter (see Supplementary 519 520 Material). Finally, the following boundary conditions are applied to remove physically impossible results and very long probability distribution tails that can arise from certain 521 combinations of input parameter values: 522

 $F_{bSi}, F_{lith}, F_{supply} > 0$   $F_{auth} < 0$   $f_{auth} > -10$   $f_{FeOx} < 10$ 

523 The model results for each individual core are given in Supplementary Material.

#### 524 5.3 Implications for the global Ge isotope budget

Across all incubated cores, the probability distribution of  $\delta^{74}$ Ge<sub>inc</sub> is centered around 525 1.6‰ (25-75th percentile range 0.8-2.4‰; Table 4, Fig. 5b), possibly slightly lighter than the 526 allogenic Ge supply  $(1.9 \pm 0.6\%)$ ; a mixture of isotopically light lithogenic- and isotopically 527 heavy biogenic-sourced Ge) but essentially indistinguishable within uncertainty. In long-term 528 mass balance terms, the isotopic composition of Ge released to pore waters (input) is expected 529 to equal the output, i.e. the combined composition of authigenic phases and the benthic flux. 530 Given that there is no detectable Ge isotope fractionation during authigenic Ge uptake at depth 531 (i.e.  $\Delta^{74}\text{Ge}_{\text{auth}} \approx 0\%$ ; see discussion above), the allogenic input and the benthic flux should 532 be isotopically indistinguishable. Any natural Fe redox-induced perturbations, much like 533 the ones observed during our incubation experiments, reflect a non-steady state process and 534 ultimately should have little effect on Ge benthic flux or authigenic composition. The latter 535 should therefore primarily depend on the ratio of Ge supplied from terrigenous vs. biogenic 536 inputs. 537

Assuming that the lack of authigenic clay  $\delta^{74}$ Ge fractionation is applicable to continental margin sediments in general and making a number of additional simplifying assumptions, we can calculate a rough estimate of authigenic and benthic flux  $\delta^{74}$ Ge in continental margin sediments on a global scale.

All the input parameters used in the following calculations and their range of uncertainties are summarized in Supplementary Material. The biogenic silica flux to shelf sediments was previously estimated as 16-87 Tmol/y, with about 3 Tmol/y buried and the rest dissolving (Tréguer and De La Rocha, 2013). Using a Ge/Si<sub>bSi</sub> ratio of 0.4-0.7  $\mu$ mol/mol (to account for potential inputs of low-Ge/Si bSi from sponges and radiolarians, e.g., Rouxel and Luais (2017)) yields a biogenic Ge flux of 26 ± 12 Mmol/y to the shelf sediments (Eq. 17).

$$FGe_{shelf}^{bSi} = FSi_{shelf}^{bSi} \times Ge/Si_{bSi}$$
(17)

To estimate the detrital Ge flux to continental margin pore waters, we first assume that this Ge must come from reducible secondary weathering products, such as amorphous Fe (oxy)hydroxides. Second, we make the simplifying assumption that all continental secondary

Ge is contained in such Fe (oxy)hydroxides. Given that a significant (but unknown) portion of 551 secondary Ge is associated with aluminosilicate clays, and that not all of the global riverine 552 particulate matter will settle on continental shelves, this calculation will yield a maximum 553 estimate of lithogenic Ge flux to shelf pore waters. Baronas et al. (2018) calculated that for 554 various global rivers, the fraction of dissolved Ge released during weathering that remains 555 in solution  $(f_{diss}^{Ge})$  ranges between 1-10% and therefore Ge uptake into secondary phases is 556 between 90-99%. Using a global riverine dissolved Ge flux (FGe<sup>riv</sup><sub>diss</sub>) of  $3.2 \pm 1.2$  Mmol/y 557 yields a maximum lithogenic Ge flux to continental margin sediments (FGe<sup>shelf</sup><sub>lith-total</sub>) of  $79 \pm 62$ 558 Mmol/y (Eq.18). 559

$$FGe_{lith-total}^{shelf} = FGe_{diss}^{riv}(\frac{1}{f_{diss}^{Ge}} - 1)$$
(18)

Next, we use our data from the San Pedro Basin to determine the fraction of lithogenic particulate Ge that gets released into pore waters (Eq. 21) and, in the absence of similar data from other sites, apply this value globally. First, the total lithogenic Ge flux to SPB sediments is calculated:

$$FGe_{lith-total}^{SPB} = F_{SPB}^{detrital} \times [Ge]_{UCC}/A_{W}$$
(19)

where  $F_{SPB}^{detrital} = 350 \pm 30 \text{ mg m}^{-2} \text{ d}^{-1}$  is the detrital flux measured via sediment traps (Collins et al., 2011), [Ge]<sub>UCC</sub> = 1.4 ± 0.2 ppm is Ge concentration of average upper continental crust (Rudnick and Gao, 2014), and A<sub>w</sub> = 72.6 g/mol is the atomic mass of germanium, yielding FGe\_{lith-total}^{SPB} of 6.8 nmol m^{-2} d^{-1}.

The net flux of Ge *released* to pore waters from Fe (oxy)hydroxides (FGe<sup>SPB</sup><sub>lith-net</sub>) can be estimated from the amount of Fe required to sustain pyrite burial in SPB sediments ( $F_{SPB}^{FeOx}$ ), calculated to be 26 µmol m<sup>-2</sup> d<sup>-1</sup> Leslie et al. (1990):

$$FGe_{lith-net}^{SPB} = F_{SPB}^{FeOx} \times Ge/Fe_{UCC}$$
(20)

where Ge/Fe<sub>UCC</sub> = 27  $\mu$ mol/mol (Rudnick and Gao, 2014). Equation 20 yields FGe<sup>SPB</sup><sub>lith-net</sub> = 0.70  $\pm$  0.26 nmol m<sup>-2</sup> d<sup>-1</sup>, which is the value used in the core incubation model described above.

The fraction of lithogenic Ge (including all silicate and oxide detrital phases) that dissolves, i.e. is released to the pore waters is thus about 10%:

$$f_{lith-released}^{Ge} = \frac{FGe_{lith-net}^{SPB}}{FGe_{lith-total}^{SPB}}$$
(21)

576 Applying this value to the global shelf lithogenic Ge input yields 6.3 Mmol/y released into 577 pore waters:

$$FGe_{lith-net}^{shelf} = FGe_{lith-total}^{shelf} \times f_{lith-released}^{Ge}$$
(22)

578 Finally, assuming that the lithogenic and the biogenic dissolved Ge inputs fully mix in 579 the sediments, the isotopic composition of the average global shelf sediment pore water is 580 calculated as

$$\delta^{74} \text{Ge}_{\text{shelf}-\text{PW}} = \frac{\text{FGe}_{\text{bSi}}^{\text{shelf}} \times \delta^{74} \text{Ge}_{\text{bSi}} + \text{FGe}_{\text{lith-net}}^{\text{shelf}} \times \delta^{74} \text{Ge}_{\text{lith}}}{\text{FGe}_{\text{bSi}}^{\text{shelf}} + \text{FGe}_{\text{lith-net}}^{\text{shelf}}}$$
(23)

Using  $\delta^{74}$ Ge<sub>lith</sub> = 0.58 ± 0.21‰ and  $\delta^{74}$ Ge<sub>bSi</sub> of 2.5 - 3.5‰ (to account for the potential contribution of isotopically lighter sponges; (Guillermic et al., 2017)), Eq. 23 yields a  $\delta^{74}$ Ge<sub>shelf-PW</sub> value of 2.47±0.42‰ (median±1 $\sigma$  of 1 million repeat Monte Carlo calculations). This value is similar to the pore water composition reported for the Southern California and Gulf of Mexico margins here. In summary, continental margin sediments are expected to exhibit lower pore water  $\delta^{74}$ Ge relative to open ocean sediments, due to an estimated 19±15% contribution of Ge from isotopically light lithogenic particles.

588 Importantly, our observations at San Pedro Basin imply that the shelf pore water composition 589 is translated to the deep authigenic Ge sink (likely, aluminosilicate clays) without significant fractionation. Therefore, the long-term benthic flux out of shelf sediments should also equal 590 approximately  $2.47 \pm 0.42\%$ . However, the magnitude of this flux (and its effect on seawater 591  $\delta^{74}$ Ge composition) will depend on the average efficiency of the authigenic Ge sink in shelf 592 sediments. Using values between 10 and 60% for dissolved Ge sequestration efficiency yields 593 a total shelf authigenic Ge burial flux of 7.0-17.7 Mmol/y. The uncertainty of this value is now 594 significantly lower compared to the previous 3.4-27 Mmol/y estimate of Baronas et al. (2016), 595 596 demonstrating the power of isotopic mass balance constraints in refining global elemental budgets. Importantly, if 10-60% of lithogenic-derived Ge is captured in shelf sediments, the 597 remaining 40-90% escapes back into the water column, contributing 2.4-6.4 Mmol/y Ge to 598 seawater. This flux is significantly higher than the previous Si budget-based estimate of detrital 599 Ge input to the ocean,  $1.6 \pm 1.5$  Mmol/y (Baronas et al., 2017). 600

Finally, we can use the refined authigenic Ge burial flux value to estimate the equivalent burial flux of Si. Rahman et al. (2017) showed that anywhere from 50 to 75% of biogenic Si is typically converted to authigenic clays in continental margin settings. Using previously measured bSi burial rates in San Pedro Basin sediments (Hammond et al., 2000; McManus et al., 2003; Baronas et al., 2016) and assuming that authigenic clay Si burial is three times higher, we can estimate authigenic Si burial rates of about 0.16 mmol m<sup>-2</sup> d<sup>-1</sup>. Combining with independent estimates of authigenic Ge burial (Baronas et al., 2016) yields authigenic clay Ge/Si values between 2.2-7.6  $\mu$ mol/mol. Assuming these Ge/Si values are applicable to margin sediments globally, the authigenic Si burial in continental margins should be in the range of 0.9-8.1 Tmol/y (best estimate of 3.1 Tmol/y), in good agreement with recent <sup>32</sup>Si-based estimates of 4.5-4.9 Tmol/y (Rahman et al., 2017), and sufficient to close the global marine Si budget.

# **6 CONCLUSIONS**

We have presented Ge/Si,  $\delta^{74}$ Ge, and supporting chemical data from seawater, pore waters, 613 and core incubations at three continental margin sites. During core incubations the flux 614 of dissolved Ge from sediments was highly variable, likely due to variable oxygenation of 615 the cores, perturbing the Fe redox conditions in shallow sediments. The incubation results 616 demonstrate the strong coupling between Ge and Fe in reducing continental margin sediments. 617 Below the very shallow Fe redox boundary, pore water  $\delta^{74}$ Ge is a mixture Ge released via 618 dissolution of isotopically heavier biogenic silica and isotopically lighter lithogenic particles 619 620 (possibly reducible Fe oxides). With depth, the precipitation of a Ge/Si-enriched authigenic 621 phase (possibly aluminosilicate clays) results in up to 50% depletion of dissolved Ge. Pore water  $\delta^{74}$ Ge signatures remain constant with depth, suggesting negligible fractionation during 622 this process. Therefore, the pore waters, the authigenic clays, and the long-term benthic 623 flux should all have identical  $\delta^{74}$ Ge signatures within uncertainty. Using global estimates 624 of biogenic and lithogenic Ge input to global continental shelf sediments, we calculate an 625 average lithogenic Ge contribution of  $19 \pm 15\%$ , with the resulting average dissolved  $\delta^{74}$ Ge of 626  $2.5 \pm 0.4\%$  in the continental margin. 627

# **CONFLICT OF INTEREST STATEMENT**

The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

# **AUTHOR CONTRIBUTIONS**

JJB and DEH designed the study. JJB and DM collected the samples. JJB and OR performed
Ge isotope analyses. JJB performed the experiments and modeling and wrote the article, with
input from all co-authors.

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#### DATA AVAILABILITY STATEMENT

All data discussed in this study is supplied in the main text and supplementary tables.

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#### SUPPLEMENTARY MATERIAL

**Table S1.** Si and trace metal concentrations in high resolution pore water samples. No measurements were made where data are not available.

Cara	Depth,	Ge,	Si,	Ge/Si,	Fe,	Mn,
Core	cm	pmol/L	µmol/L	µmol/mol	µmol/L	nmol/L
MC-1D	1	380	249	1.53	63	57
MC-1D	3	696	324	2.15	106	90
MC-1D	3	707	308	2.30	128	100
MC-1D	5	471	352	1.34	118	124
MC-1D	7	356	367	0.97	40	161
MC-1D	9	379	391	0.97	87	183
MC-1D	13	203	390	0.52	55	202
MC-1D	13	249	422	0.59	60	228
MC-1D	16	262	452	0.58	59	265
MC-1D	24	199	462	0.43	53	375
MC-1D	29	180	492	0.37	39	321
MC-1D	33	181	517	0.35	39	398
	0	504	0.07	4 70	00	00
MC 2A	0	720	297	1.78	00	30
MC 2A	1	130	340	2.15	150	54 75
MC-ZA	2	1177	363	3.24	159	75
MC-2A	4	432	353	1.22	118	226
MC-ZA	8	380	356	1.07	83	123
MC-2A	15	214	440	0.49	12	217
MC-2A	29	117	517	0.23	43	354
MC-2A	33	111	525	0.21		
MC-2D	0	336	230	1.46	18	3
MC-2D	3	614	338	1.82	137	72
MC-2D	5	351	370	0.95	113	113
MC-2D	10	286	395	0.72	84	174
MC-2D	16	266	468	0.57	94	223
MC-2D	20	171	457	0.37	63	266
MC-3D	1	859	273	3.15	128	66
MC-3D	2	695	320	2.17	117	83
MC-3D	2	556	282	1.97	97	102
MC-3D	15	540	451	1.20	77	259
MC-3D	22	196	447	0.44	38	250
MC-3D	30		464		46	619
MC-3D	35	139	508	0.27	29	374
MC-5C-1	1	518	309	1.67	1	0
MC-5C-1	2	531	514	1.03	261	175
MC-5C-1	4	522	546	0.96	272	352
MC-5C-1	8	446	463	0.96	306	321
MC-5C-1	13	315	426	0.74	79	218
MC-5C-1	19	252	447	0.56	67	258
MC-5C-1	23	177	460	0.38	51	290
MC-5C-1	27	120	439	0.27	30	278
MC-5C-2	1.5	413				
MC-5C-2	6.5	535				
MC-5C-2	14.5	241				
MC-5C-2	25.5	130				
MC-5C-2	34.5	89				

**Table S2.** Ammonia concentrations in high resolution pore water samples and overlying water. Cores MC-1B, MC-2B, and MC-4B are distinct from cores analyzed for Ge and Si, but were collected alongside the cores discussed above.

Coro	Depth,	NH <sub>3</sub> ,	Coro	Depth,	NH <sub>3</sub> ,
Core	cm	µmol/L	Cole	cm	µmol/L
MC-1B	0.5	24	MC-1B	OLW	8
MC-1B	2.5	51	MC-2B	OLW	8
MC-1B	4.5	78	MC-4B	OLW	24
MC-1B	6.5	93			
MC-1B	8.5	120	MC-2A	0	44
MC-1B	12.5	158	MC-2A	1	37
MC-1B	16	187	MC-2A	2	33
MC-1B	20.5	217	MC-2A	3	59
MC-1B	27	276	MC-2A	4	62
			MC-2A	5	83
MC-2B	0.5	29	MC-2A	7	102
MC-2B	3.5	63	MC-2A	8	97
MC-2B	5.5	116	MC-2A	9	127
MC-2B	10.5	136	MC-2A	10	123
MC-2B	15.5	176	MC-2A	11	160
MC-2B	20.5	203	MC-2A	12	137
MC-2B	25.5	238	MC-2A	14	169
MC-2B	30.5	263	MC-2A	15	172
MC-2B	36	302	MC-2A	17	209
MC-2B	42.5	337	MC-2A	20	234
			MC-2A	24	261
MC-4B	0.5	17	MC-2A	27	264
MC-4B	3.5	57	MC-2A	29	274
MC-4B	6.5	86	MC-2A	30	294
MC-4B	9.5	139	MC-2A	33	297
MC-4B	15.5	201			
MC-4B	18.5	222	MC-4B	34	339
MC-4B	22.5	273	MC-4B	40	379
MC-4B	28.5	299	MC-4B	43	430

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Samplo	Depth,	SO <sub>4</sub> ,
Sample	m	mmol/L
SPOT SSW	0	27.4
SPOT 885m	885	27.2
MC-5B-1 (Core inc.)	885	26.4
5B-Final (Core inc.)	885	26.5
MC-2C OLW	885	29.6
MC-2D OLW	885	27.2
MC-5C OLW	885	27.8
MC-1D OLW	885	26.5
MC-2A OLW	885	26.1
Coro	Depth,	SO <sub>4</sub> ,
COIE	cm	mmol/L
MC-1D	1	26.5
MC-1D	3	24.4
MC-1D	5	26.1
MC-1D	9	26.3
MC-1D	16	26.0
MC-1D	24	24.5
MC-1D	29	24.9
MC-1D	33	23.5
MC-2A	0	26.1
MC-2A	4	25.6
MC-2A	12	25.4
MC-2A	20	24.7
MC-2A	29	25.4

**Table S3.** Sulfate concentrations in seawater, high resolution pore water samples, and overlying water. Measurement uncertainty is 4 %.

Commis	Time	Corr. time,	Ge,	Si,	Ge/Si,
Sample	Time, n	d/m *	pmol/L	µmol/L	µmol/mol
MC-3A					
1	0	0	73	103	0.71
2	8.0	2.8	83	112	0.74
3	17.5	6.2	86	125	0.69
4	40.3	14.5	104	130	0.80
5	64.3	23.5	116	145	0.81
Final	65.8	24.1	118	142	0.83
<u>MC-5D</u>					
1	0	0	83	105	0.79
2	6.0	1.8	86	108	0.79
3	23.4	7.2	88	115	0.76
4	45.4	14.2	87	120	0.73
5	68.4	21.7	94	127	0.74
Final	68.4	21.7	98	129	0.76
<u>MC-4C</u>					
1	0	0	83	106	0.78
2	3.3	1.0	83	110	0.75
3	3.5	1.1	85	109	0.77
4	23.0	7.3	88	120	0.74
5	46.0	14.9	87	122	0.71
6	69.8	22.9	92	130	0.71
7	92.8	30.8	101	136	0.74
8	118.5	39.8	102	143	0.72
9	140.5	47.6	109	147	0.74
<u>MC-5A</u>					
1	0	0	81	105	0.77
2	5.5	1.9	82	112	0.73
3	23.0	8.1	84	120	0.70
4	45.0	16.1	84	125	0.67
5	70.0	25.3	87	134	0.65
6	93.0	34.0	83	140	0.60
7	117.5	43.4	86	147	0.58
8	139.5	52.1	85	155	0.55
<u>MC-5B</u>					
1	0	0	74	104	0.71
2	5.5	1.5	74	108	0.69
3	23.0	6.4	69	114	0.61
4	45.0	12.6	78	120	0.66
5	70.0	19.9	78	126	0.62
6	93.0	26.8	86	131	0.65
7	117.5	34.2	84	139	0.61
8	139.0	40.8	87	142	0.61

**Table S4.** Ge and Si concentrations during San Pedro Basin core incubations.

\* Calculated as sum of incubation time in days divided by height of the overlying incubated water column in m at a given time. The height of the water column decreases throughout the incubation due to water removal by sampling. This calculation corrects for the effect of decreasing water volume, and is used to calculate the Ge and Si fluxes reported in Table 3 of main text.

Sample	Time, h	Corr. time, d/m *	Ge, pmol/L	Si, umol/L	Ge/Si, umol/mol
D3-S2			p		
1	11.5	4.9	96	126	0.76
2	24.6	10.6	104	131	0.79
3	31.2	13.6	107	133	0.80
4	49.7	22.2	114	140	0.81
5	119.2	56.0	145	154	0.94
D4-S1					
1	9.8	3.5	95	129	0.74
2	22.7	8.3		134	
3	29.6	10.9	105	137	0.76
4	48.5	18.3		143	
5	118.4	46.4	147	165	0.89
<u>D4-S4</u>					
1	9.8	4.4	99	129	0.76
2	22.8	10.5		135	
3	29.6	13.9	102	138	0.74
4	48.5	23.4		146	
5	118.2	59.7	147	170	0.86
<u>D5-S1</u>					
1	8.5	2.9	91	131	0.70
2	21.6	7.4		136	
3	28.3	9.8	96	138	0.69
4	46.8	16.6		139	
5	116.9	42.9	151	189	0.80
<u>D5-S4</u>					
1	8.6	2.4	88	131	0.67
2	23.0	6.4		134	
3	28.3	8.0	90	136	0.66
4	46.9	13.5		145	
5	116.8	34.6	116	163	0.71

**Table S5.** Ge and Si concentrations during Santa Monica Basin core incubations. No measurements were made where data are not available.

\* Calculated as sum of incubation time in days divided by height of the overlying incubated water column in m at a given time. The height of the water column decreases throughout the incubation due to water removal by sampling. This calculation corrects for the effect of decreasing water volume, and is used to calculate the Ge and Si fluxes reported in Table 3 of main text.

Parameter	Range	PDF	Source
F <sub>lith</sub> (nmol m <sup>-2</sup> d <sup>-1</sup> )	0.70 ± 0.26	normal	1
Ge/Si <sub>bSi</sub> (µmol/mol)	0.70 ± 0.10	normal	2
δ <sup>74</sup> Ge <sub>initial</sub> (‰)	3.06 ± 0.17	normal	3
δ <sup>74</sup> Ge <sub>bSi</sub> (‰)	3.28 ± 0.52	normal	4
δ <sup>74</sup> Ge <sub>lith</sub> (‰)	0.58 ± 0.21	normal	5
δ <sup>74</sup> Ge <sub>pw</sub> (‰)	2.13 ± 0.25	normal	6
$\Delta^{74}\text{Ge}_{\text{FeOx-diss}}$ (‰)	-4.61.6	uniform	7
$\Delta^{74}\text{Ge}_{\text{auth-diss}}$ (‰)	-0.3 - 0.3	uniform	8

Table S6. Core incubation model input parameters.

(1) Calculated from FeOx flux reported by Leslie et al. (1990) and UCC Ge/Fe ratio (Rudnick & Gao, 2014) – see supp. text; (2) Baronas et al. (2016); (3) Mean of all measured pre-incubation overlying water values (Table 1); (4) Mean of meaured seawater values (Table 1); (5) Rouxel & Luais (2017); (6) Mean of all pore water values below 5 cm depth (incl. the single SMB measurement); (7) Pokrovsky et al. (2014); (8) Based on the negligible  $\delta^{74}Ge_{pw}$  gradient in SPB sediments (Fig. 1)

Table S7.	Core incubation model results for San Pedro and Santa Monica Basin cores	. Values are reported as median with 25-75th
percentiles	in parentheses.	

			San Pedro Basin				Santa Monica Basin	
Core	MC-3A	MC-5D	MC-4C	MC-5A	MC-5B	D3S2	D4S4	D5S1
n <sub>initial</sub> (nmol)	62 (59 - 66)	81 (77 - 85)	82 (78 - 86)	69 (65 - 73)	80 (77 - 84)	67 (63 - 72)	60 (55 - 65)	73 (69 - 78)
n <sub>inc</sub> (nmol)	34 (32 - 36)	8.1 (7.5 - 8.6)	20 (19 - 22)	1.01 (0.94 - 1.07)	17 (16 – 18)	18 (17 – 19)	17 (16 – 18)	30 (28 - 32)
f <sub>lith</sub>	0.39 (0.35 - 0.42)	0.49 (0.45 - 0.53)	0.56 (0.52 - 0.6)	0.57 (0.53 - 0.6)	0.55 (0.51 – 0.59)	0.65 (0.62 - 0.69)	0.57 (0.53 – 0.61)	0.4 (0.37 - 0.44)
f <sub>bSi</sub>	0.61 (0.58 - 0.65)	0.51 (0.47 - 0.55)	0.44 (0.4 - 0.48)	0.43 (0.4 - 0.47)	0.45 (0.41 - 0.49)	0.35 (0.31 – 0.38)	0.43 (0.39 - 0.47)	0.6 (0.56 - 0.63)
f <sub>auth</sub>	-0.81 (-1.320.43)	-0.22 (-0.440.09)	-0.2 (-0.370.08)	-0.23 (-0.430.1)	-0.16 (-0.310.07)	-0.11 (-0.220.05)	-0.1 (-0.20.04)	-0.19 (-0.360.08)
f <sub>FeOx</sub>	0.8 (0.41 - 1.32)	-0.48 (-0.610.26)	-0.38 (-0.50.2)	-0.75 (-0.880.55)	-0.48 (-0.580.33)	0 (-0.12 - 0.16)	-0.17 (-0.270.04)	0.07 (-0.07 - 0.25)
f <sub>released</sub>	0.98 (0.89 - 1.08)	0.29 (0.26 - 0.32)	0.41 (0.37 - 0.46)	0.02 (0.02 - 0.02)	0.34 (0.31 - 0.38)	0.86 (0.77 - 0.97)	0.71 (0.63 – 0.79)	0.85 (0.77 - 0.93)
F <sub>bSi</sub> (nmol m⁻² d⁻¹)	1.11 (1.02 - 1.21)	0.71 (0.65 - 0.76)	0.53 (0.49 - 0.57)	0.52 (0.48 - 0.57)	0.55 (0.51 - 0.6)	0.35 (0.33 - 0.38)	0.51 (0.47 - 0.55)	1.02 (0.94 - 1.1)
F <sub>supply</sub> (nmol m <sup>-2</sup> d <sup>-1</sup> )	1.81 (1.69 - 1.94)	1.39 (1.29 - 1.5)	1.21 (1.11 - 1.31)	1.2 (1.11 - 1.3)	1.23 (1.13 – 1.33)	1.03 (0.94 - 1.12)	1.19 (1.09 – 1.29)	1.71 (1.6 – 1.84)
F <sub>auth</sub> (nmol m⁻² d⁻¹)	-1.46 (-2.380.78)	-0.31 (-0.610.13)	-0.24 (-0.450.1)	-0.28 (-0.520.12)	-0.2 (-0.390.09)	-0.12 (-0.230.05)	-0.12 (-0.240.05)	-0.33 (-0.620.14)
F <sub>FeOx</sub> (nmol m <sup>-2</sup> d <sup>-1</sup> )	1.43 (0.74 - 2.35)	-0.65 (-0.850.35)	-0.45 (-0.620.24)	-0.88 (-1.060.64)	-0.58 (-0.730.39)	0 (-0.13 – 0.15)	-0.2 (-0.330.04)	0.11 (-0.12 - 0.41)
F <sub>inc</sub> (nmol m⁻² d⁻¹)	1.77 (1.65 – 1.89)	0.41 (0.38 - 0.43)	0.5 (0.46 - 0.53)	0.02 (0.02 - 0.03)	0.42 (0.4 - 0.45)	0.89 (0.83 - 0.95)	0.84 (0.78 – 0.9)	1.45 (1.36 – 1.55)
δ <sup>74</sup> Ge <sub>supply</sub> (‰)	2.24 (2.09 - 2.39)	1.98 (1.85 – 2.13)	1.79 (1.66 - 1.92)	1.78 (1.65 – 1.92)	1.83 (1.7 - 1.97)	1.54 (1.42 - 1.66)	1.79 (1.66 - 1.92)	2.21 (2.07 - 2.36)
δ <sup>74</sup> Ge <sub>FeOx</sub> (‰)	-0.85 (-1.60.11)	-0.42 (-1.02 - 0.01)	-0.73 (-1.350.26)	-0.47 (-0.860.15)	-0.47 (-0.920.12)	-1.29 (-2.080.64)	-0.69 (-1.390.2)	-0.72 (-1.490.03)
Δ <sup>74</sup> Ge <sub>FeOx-diss</sub> (‰)	-3.1 (-3.852.35)	-2.39 (-3.011.95)	-2.51 (-3.152.02)	-2.23 (-2.651.9)	-2.27 (-2.761.91)	-2.83 (-3.642.17)	-2.46 (-3.21.97)	-2.94 (-3.722.24)
δ <sup>74</sup> Ge <sub>inc</sub> (‰)	-0.16 (-1.41 - 0.92)	3.08 (1.37 - 4.67)	1.54 (0.66 - 2.38)	27.2 (13 - 41.46) *	2.76 (1.98 - 3.52)	1.09 (0.67 - 1.4)	1.7 (1.27 - 2.09)	1.78 (1.33 – 2.16)
* Value highly uncertain and	likely inacurrate due to th	e extremely low Ge incl	ubation flux (F <sub>inc</sub> ) for this	s core. This value is there	fore excluded from the sur	nmary table and figures in	the main text.	

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**Table S8.** Input parameters used to calculate  $\delta^{74}$ Ge<sub>shelf-PW</sub>, the average global isotope composition of dissolved Ge in continental margin pore waters.

Parameter	Symbol	Value	Units	Source
Ge concentration in riverine sediments	[Ge] <sub>UCC</sub>	1.4 ± 0.2	ppm	Rudnick & Gao (2014)
Dissolved riverine Ge flux	$FGe^{riv}_{diss}$	3.2 ± 1.2	Mmol/y	Baronas et al. (2017)
Fraction Ge released during continental weathering taken up into secondary phases	1- f <sub>diss</sub> Ge	90-99%		Baronas et al. (2018)
Ge isotope composition of riverine sediments	$\delta^{74}Ge_{\text{lith}}$	0.58 ± 0.21‰		Rouxel & Luais (2017)
bSi dissolving in global continental margin sediments	FSi <sup>bSi</sup> shelf	12.7-83.7	Tmol/y	Treguer & De La Rocha (2013)
Detrital rain to the seafloor (San Pedro Basin)	F <sup>detrital</sup> SPB	350 ± 30	mg/(m <sup>2</sup> d)	Collins et al. (2011)
Reducible Fe(OH) <sub>3</sub> flux (San Pedro Basin)	F <sup>FeOx</sup> SPB	26	µmol/(m <sup>2</sup> d)	Leslie et al. (1990)
Ge/Si ratio of biogenic silica	Ge/Si <sub>bSi</sub>	0.5-0.7	µmol/mol	Rouxel & Luais (2017)
Ge isotope composition of biogenic silica	$\delta^{74}Ge_{bSi}$	2.5 - 3.5‰		Guillermic et al. (2017)



Figure S1. Summary of core incubation model results, showing the probability distributions of calculated values for each core.