

Anthropogenic perturbations of the silicon cycle at the global scale: Key role of the land-ocean transition

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Received 19 May 2008; revised 23 June 2009; accepted 13 August 2009; published 31 December 2009.

[1] Silicon (Si), in the form of dissolved silicate (DSi), is a key nutrient in marine and continental ecosystems. DSi is taken up by organisms to produce structural elements (e.g., shells and phytoliths) composed of amorphous biogenic silica (bSiO₂). A global mass balance model of the biologically active part of the modern Si cycle is derived on the basis of a systematic review of existing data regarding terrestrial and oceanic production fluxes, reservoir sizes, and residence times for DSi and bSiO₂. The model demonstrates the high sensitivity of biogeochemical Si cycling in the coastal zone to anthropogenic pressures, such as river damming and global temperature rise. As a result, further significant changes in the production and recycling of bSiO₂ in the coastal zone are to be expected over the course of this century.

Citation: Laruelle, G. G., et al. (2009), Anthropogenic perturbations of the silicon cycle at the global scale: Key role of the land-ocean transition, *Global Biogeochem. Cycles*, 23, GB4031, doi:10.1029/2008GB003267.

1. Introduction

[2] Silicon (Si) is the second most abundant element in the Earth's crust after oxygen. Most Si, however, is bound in the form of quartz and silicate minerals, and is therefore unavailable for uptake by organisms. Thus, despite its abundance, Si is a major limiting element in many aquatic systems [Conley and Malone, 1992; Egge and Aksnes, 1992; Paasche, 1980; Leynaert et al., 2001], and is also an essential nutrient for the growth of many terrestrial plants [Epstein, 1999; Datnoff et al., 2001]. Key aspects of the global biogeochemical silicon cycle remain poorly understood, such as the biological cycling of Si on the continents [Conley, 2002a], the role of the coastal zones in regulating the transfer of reactive Si from land to the open ocean [Conley, 1997; DeMaster, 2002], the fate of biogenic silica

produced in oceanic surface waters and its decoupling from carbon during sinking [Nelson et al., 1995; Ragueneau et al., 2002, 2006a], and ongoing changes to the Si cycle by human activities [Chauvaud et al., 2000; Conley et al., 1993; Friedl and Wüest, 2002; Friedl et al., 2004; Humborg et al., 2000, 2006; Ragueneau et al., 2005, 2006b, 2006c; Conley et al., 2008].

[3] Global-scale studies of the biogeochemical Si cycle have focused mainly on the marine aspect. An important landmark in the assessment of Si fluxes in the world ocean is the work of Tréguer et al. [1995]. These authors, however, provide no estimates of the amounts of biogenic silica stored in the oceans and underlying sediments. Furthermore, only a crude representation of the land-ocean interface is included in their global Si budget. In this respect, the current state of knowledge and modeling of

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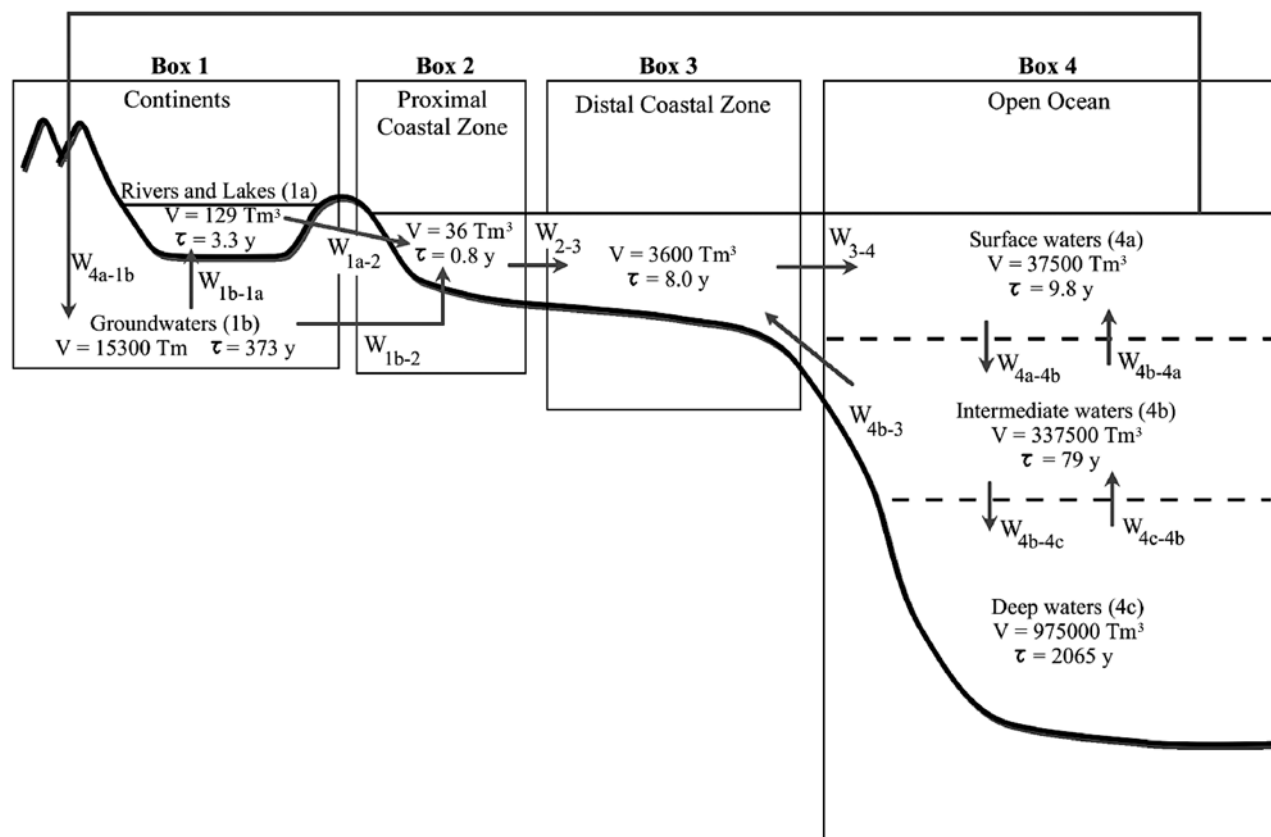


Figure 1. Global Water Cycle with water masses in Tm^3 and water fluxes (W) in $\text{Tm}^3 \text{yr}^{-1}$. V , volume; τ , residence time.

global carbon, nitrogen, phosphorus and sulfur cycles [Mackenzie *et al.*, 1993, 1998; Ver, 1998; Rabouille *et al.*, 2001] are significantly more advanced than for silicon.

[4] In this study, we provide global-scale estimates of reservoir sizes and fluxes of reactive Si on the continents in the ocean and at the continent-to-ocean transition. Emphasis is placed on the biogeochemical dynamics of Si at the Earth's surface, from the recent past to the end of the 21st century. We therefore do not explicitly represent the long-term endogenic Si cycling, but rather include the Earth's lithosphere as the ultimate source and sink of reactive Si. The two forms of reactive Si considered are dissolved silicate (DSi) and biogenic silica (bSiO_2). The main transformation processes in the global biogeochemical Si cycle are uptake of DSi followed by biomineralization as bSiO_2 in terrestrial plants and aquatic organisms, and the dissolution of bSiO_2 into DSi.

[5] The resulting mass balance model is used to explore the sensitivity of the global Si cycle and gain insight into its function. Special attention is paid to the role of the coastal zone and continental shelves on the coupling of terrestrial and oceanic Si dynamics. In addition, the response of the global biogeochemical Si cycle to two anthropogenically driven forcings is analyzed: global temperature rise and river damming. These forcings are selected because both siliceous production and bSiO_2 dissolution are sensitive to temperature [Wollast, 1974; Cossins and Bowler, 1987; Rickert, 2000; Van Cappellen *et al.*, 2002], while increased

river damming, especially since the 1950s, has considerably modified the reactive Si delivery to the oceans [Conley, 2002b; Humborg *et al.*, 2006].

2. Global Biogeochemical Si Model

2.1. Water Cycle

[6] The Earth's surface environment is divided into four compartments (Figure 1): continents (box 1), proximal (box 2) and distal (box 3) coastal zones, and the open ocean (box 4). The proximal and distal coastal zones are those proposed by Rabouille *et al.* [2001]. As shown by these authors, this division of the global coastal zone provides a more realistic representation of the role of continent to ocean transition in the biogeochemical cycling of carbon and nutrients. The proximal zone consists of large bays, the open water parts of estuaries, inner deltas, inland seas and coastal marshes [Woodwell *et al.*, 1973]. The distal zone comprises the rest of the continental margins up to the shelf break.

[7] These compartments are linked to one another via the water cycle (Figure 1). Water on the continents is subdivided into an aquatic reservoir, which comprises exorheic rivers and lakes including their floodplains (box 1a), and a groundwater reservoir (box 1b). The open ocean is by far the largest compartment, with a mean water depth of 3600 m and covering 92% of the world ocean [Tréguer *et al.*, 1995]. Three vertical subcompartments of the water column

Table 1. Calculations and References Used to Constrain the Water Fluxes in the Model

Flux	Value (Tm ³ yr ⁻¹)	Calculation	Reference ^a
W _{1b-1a}	39	-	1
W _{1a-2}	39	W _{4a-1} - W _{1b-2}	-
W _{1b-2}	2	-	2
W ₂₋₃	41	W _{1a-2} + W _{1b-2}	-
W ₃₋₄	441	W ₂₋₃ - W ₄₋₃	-
W ₄₋₃	400	-	3, 4, 5
W _{4a-4b}	3800	W _{4b-4a} + W ₃₋₄ - W _{4a-1}	-
W _{4b-4a}	3400	-	6
W _{4b-4c}	472	-	6
W _{4c-4b}	472	W _{4b-4c}	-
W _{4a-1b}	41	-	7

^aReferences are as follows: 1, *Fekete et al.* [2002]; 2, *Slomp and Van Cappellen* [2004]; 3, *Brink et al.* [1995]; 4, *Ver* [1998]; 5, *Rabouille et al.* [2001]; 6, Deduced from residence time by *Broecker and Peng* [1982]; 7, *Anikouchine and Sternberg* [1981].

are considered: a 100 m thick surface layer where photosynthesis takes place (box 4a), mesopelagic oceanic waters (100–1000 m depth) (box 4b) and deep waters (box 4c).

[8] Water fluxes (W) considered in the baseline scenario are based on the simplified steady state water cycle summarized in Table 1. It should be noted that a contribution from subsurface groundwater discharge to the proximal coastal zone is explicitly considered (W_{1b-2}) [*Slomp and Van Cappellen*, 2004]. The combination of water reservoir

sizes with water fluxes yields water residence times that agree well with previous studies [*Garrels and Mackenzie*, 1971; *Broecker and Peng*, 1982; *Macdonald*, 1998].

2.2. Reactive Si Reservoirs

[9] The DSi and bSiO₂ reservoir masses are time-dependent variables of the Si cycle model (Figure 2). DSi mainly consists of undissociated monomeric silicic acid, Si(OH)₄, and represents the main form under which silicon can be assimilated by organisms [*Del Amo and Brzezinski*, 1999]. Organisms use DSi to build structural elements made of amorphous, hydrated silica, part of which is preserved after the death of the organisms [*Simpson and Volcani*, 1981; *Conley and Schelske*, 2001]. Here, bSiO₂ includes the amorphous silica in both living biomass and biogenic detritus in open waters, soils and sediments. It should be noted that bSiO₂ may undergo significant chemical and mineralogical changes [*Van Cappellen et al.*, 2002], even including a complete diagenetic transformation of the opaline silica into aluminosilicate minerals [*Michalopoulos et al.*, 2000].

[10] In marine environments, the major producers of bSiO₂ are diatoms, although other organisms, such as radiolarians, sponges, and chrysophytes, may locally be important sources of bSiO₂ [*Simpson and Volcani*, 1981]. On land, large quantities of DSi are fixed by higher plants and deposited as amorphous silica in so-called phytoliths

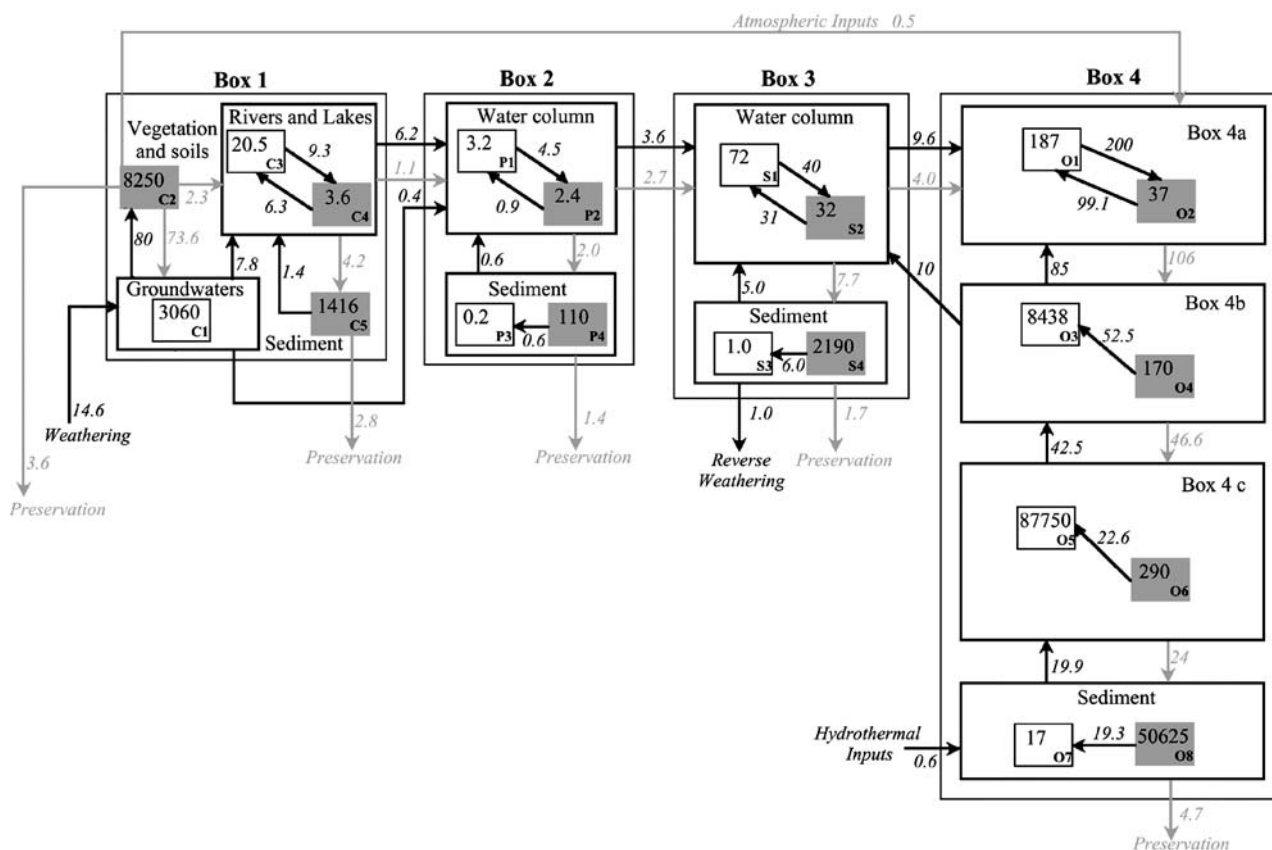


Figure 2. Steady state biogeochemical cycle of silicon, with reservoirs in Tmol Si and fluxes in Tmol Si yr⁻¹. Shaded squares represent bSiO₂ reservoirs, open squares represent DSi reservoirs. The steady state cycle of Si is used as initial condition (nominally 1950) in the perturbation simulations.

Table 2. Reservoir Sizes of Dissolved Silicate (DSi) and Biogenic Silica (bSiO₂) in the Model^a

	Reservoir	Size (Tmol Si)	Calculation	Reference ^b
		<i>Continents</i>		
C1	Terrestrial DSi	3060	200 $\mu\text{mol Si L}^{-1}$ ^(a) \times V(1b) ^(b) , where a is average DSi concentration in groundwaters and b is volume of groundwaters	1, 2
C2	Terrestrial bSiO ₂	8250	0.42g Si ^(a) \times 0.005 g g ^(b) \times (75.10 ¹⁸ m ² ^(c) \times 1m ^(d) \times 2.65 g.cm ⁻³ ^(e) \times 0.55 ^(f))/M(Si), where a is mass of Si per gram of phytoliths, b is average mass of phytoliths per gram of soil, c is global surface area covered by soils, d is assumed global average soil depth, e is average dry density of soils, and f is solid fraction of soils	2, 3, 4, 5
C3	Aquatic bSiO ₂	3.6	-	6
C4	Aquatic DSi	20.5	(9.3 mg Si L ⁻¹ ^(a) \times V(1b) ^(b))/M(Si), where a is average DSi concentration in rivers and b is global volume of lakes and floodplains	7
C5	bSiO ₂ in river and lake sediments	1416	50 mg SiO ₂ g sediment ⁻¹ ^(a) /M(SiO ₂) \times VS(1) ^(b) \times (1 - ϕ) \times ρ , where a is average bSiO ₂ content of river sediments and b is volume of lake plus river sediments	8
		<i>Proximal Coastal Zone</i>		
P1	DSi in the water column	3.2	F _{P1S1} \times V(2)/W ₂₋₃	-
P2	bSiO ₂ in the water column	2.4	F _{P2S2} \times V(2)/W ₂₋₃	-
P3	pore water DSi in sediments	0.2	300 $\mu\text{mol Si L}^{-1}$ \times VS(2) \times ϕ	9
P4	bSiO ₂ in sediments	110	0.02 g Si g ⁻¹ ^(a) \times VS(2) \times (1 - ϕ) \times ρ /M(Si), where a is mass of Si as bSiO ₂ in proximal coastal sediments (2 wt%)	10
		<i>Distal Coastal Zone</i>		
S1	DSi in the water column	72	F _{S1O1} \times V(3)/W _{3-4a}	-
S2	bSiO ₂ in the water column	32	F _{S2O2} \times V(3)/W _{3-4a}	-
S3	pore water DSi in sediments	1	300 $\mu\text{mol Si L}^{-1}$ \times VS(3) \times ϕ	9
S4	bSiO ₂ in sediments	2190	0.03 g Si g ⁻¹ ^(a) \times VS(3) \times (1 - ϕ) \times ρ /M(Si), where a is mass of Si as bSiO ₂ in distal coastal sediments (3 wt%)	9
		<i>Open Ocean</i>		
O1	DSi in the surface ocean	187	5 $\mu\text{mol L}^{-1}$ \times V(4a)	11, 12, 13
O2	bSiO ₂ in the surface ocean	37	1 $\mu\text{mol L}^{-1}$ \times V(4a)	-
O3	DSi in intermediate waters	8438	25 $\mu\text{mol L}^{-1}$ \times V(4b)	10, 14, 15
O4	bSiO ₂ in intermediate waters	170	0.5 $\mu\text{mol L}^{-1}$ \times V(4b)	-
O5	DSi in deep sea	87750	100 $\mu\text{mol L}^{-1}$ \times V(4c)	10, 14, 15
O6	bSiO ₂ in deep sea	290	0.1 $\mu\text{mol L}^{-1}$ \times V(4c)	-
O7	pore water DSi in sediments	17	300 $\mu\text{mol Si L}^{-1}$ \times VS(4) \times ϕ	9
O8	bSiO ₂ in sediments	50625	0.05 g Si g ⁻¹ ^(a) \times VS(4) \times (1 - ϕ) \times ρ /M(Si), where a is mass of Si as bSiO ₂ in deep sea sediments (5 wt%)	16

^aV(x) and VS(x) refer to the volumes of water and sediment reservoirs of box x as described in Figure 1. The porosity ($\phi = 0.7$) and the density ($\rho = 2.5$ g cm⁻³) of the sediments are taken from *Maher et al.* [2004]. M(Si) and M(SiO₂) stand for the atomic mass of Si and molar weight of SiO₂, respectively.

^bReferences are as follows: 1, J.-D. Meunier (personal communication); 2, *Berner and Berner* [1996]; 3, *Conley* [2002b]; 4, *Food and Agriculture Organization/U.N. Educational, Scientific and Cultural Organization (FAO/UNESCO)* [1986]; 5, *Schroeder* [1978]; 6, *Conley* [1997]; 7, *Dürr et al.* [2009]; 8, *Conley* [1988]; 9, *Jahnke et al.* [1982]; 10, *Ledford-Hoffman et al.* [1986]; 11, *Nelson et al.* [1995]; 12, *Yool and Tyrrell* [2003]; 13, *Conkright et al.* [1994]; 14, *Dittmar and Birkicht* [2001]; 15, *Hill and Wheeler* [2002]; 16, *Bonn* [1995].

[Piperno, 1988]. The significant contribution of phytolith production and dissolution in the global Si cycle has only recently been highlighted [Bartoli, 1983; Meunier et al., 1999; Conley, 2002a].

[11] Estimations of DSi and bSiO₂ reservoir sizes in the four earth surface compartments are summarized in Table 2. Reservoir masses are mostly derived from estimates of the average DSi or bSiO₂ concentrations and the volumes of the

Table 3. Silicon Fluxes in the Model^a

Flux	Flux (Tmol yr ⁻¹)	Calculation or Explanation	Reference ^b	
Continents				
F _w	Weathering	14.6	$F_{C1C2} - F_{C2C1} + F_{C1C3} + F_{C1P1}$	
F _{C2-burial}	Terrestrial bSiO ₂ burial	3.6	25% of F _w	1
F _{C1C2}	Terrestrial DSi uptake	80	reported range: 60 to 200 Tmol Si yr ⁻¹	2
F _{C1C3}	DSi export from box 1b to box 1a	7.8		
F _{C1P1}	DSi export from box 1b to box 2	0.4	$W_{1b-2} \times 200 \mu\text{mol Si L}^{-1}$ in groundwaters	3
F _{C2C1}	Terrestrial bSiO ₂ dissolution	73.6	92% of F _{C1C2}	1
F _{C2O2}	Eolian silica export	0.5		4
F _{C2C4}	bSiO ₂ export to rivers	2.3		
F _{C3C4}	DSi uptake in rivers	9.3	61% ^(a) of carbon primary production \times 17.10 ¹² m ² ^(b) \times 13.5g C m ⁻² yr ⁻¹ ^(c) \times (Si/C) ^(d) /M(C), where a is fraction of primary production due to diatoms in fresh waters, b is surface of rivers and lakes, c is average riverine primary production in carbon, and d is molar Si to C ratio of 0.79 \pm 0.43.	5, 6, 7, 8
F _{C3P1}	DSi export from box 1a	6.2		9
F _{C4C3}	bSiO ₂ dissolution	6.3		7
F _{C4C5}	bSiO ₂ Sedimentation in rivers	4.2	$F_{C2C4} + F_{C3C4} - F_{C4P1} - F_{C4C3}$	
F _{C4P2}	bSiO ₂ export from box 1a	1.1		2
F _{C5C3}	Sediment bSiO ₂ dissolution	1.4		
F _{C5-burial}	Aquatic bSiO ₂ burial	2.8	$F_{C4C5} - F_{C5C3}$	
Proximal Coastal Zone				
F _{P1P2}	DSi uptake	4.5	75% ^(a) of carbon primary production (40 Tmol C yr ⁻¹ ^(b)) \times (Si/C) ^(c) , where a is fraction of primary production due to diatoms in the coastal zone, b is primary production in the proximal coastal zone in carbon, and c is molar Si to C ratio of 0.15	10, 11
F _{P1S1}	DSi export from box 2	3.6	$P1 \times W_{2-3}$	
F _{P2P1}	bSiO ₂ dissolution	0.9	$F_{P1P2} + F_{C4P2} - F_{P2S2} - F_{P2P4}$	
F _{P2P4}	Sedimentation	2	$F_{P4P3} + F_{P4-burial}$	
F _{P2S2}	bSiO ₂ export from box 2	2.7	$F_{P1P2} - F_{P2P4} - F_{P2P1}$	
F _{P3P1}	DSi efflux	0.6	Within the range of 0.8 \pm 0.4 Tmol Si yr ⁻¹ given by 3.6 Tm ² \times 22 \pm 11.10 ⁻² mol Si yr ⁻¹ m ⁻²	12
F _{P4P3}	Sediment bSiO ₂ dissolution	0.6	F_{P3P1}	
F _{P4-burial}	bSiO ₂ Burial	1.4	$P4 \times 0.13 \text{ cm yr}^{-1}$ ^(a) /10 cm ^(b) (and so that F _{P4-burial} + F _{S4-burial} = 3.1), where a is sediment accumulation rate in the proximal coastal zone and b is depth of the active sediment layer in the proximal coastal zone	11, 13, 14
Distal Coastal Zone				
F _{S1S2}	DSi uptake	40	[18% ^(a) of global siliceous primary production (240 Tmol Si yr ⁻¹ ^(b)]-F _{P1P2} , where a is fraction of the global marine primary production occurring in the coastal zone and b is estimate of the global marine siliceous production	4, 15
F _{S2S1}	bSiO ₂ dissolution	31	$F_{S2S1} + F_{S3S1} + F_{O3S1} + F_{P1S1} - F_{S1S2}$	
F _{S1O1}	bSiO ₂ export from box3	9.6	$F_{S1S2} + F_{P2S2} - F_{S2S4} - F_{S2O2}$	
F _{S2S1}	bSiO ₂ dissolution	31	$F_{S4-burial} + F_{S4S3}$	
F _{S2S4}	Sedimentation	7.7		
F _{S2O2}	bSiO ₂ export from box3	4	assuming the same export rate as organic carbon, 10% of F _{S1S2}	15, 16, 17
F _{S3S1}	DSi efflux	5	within the range of 6 \pm 3 Tmol Si yr ⁻¹ given by 27 Tm ^{2(a)} \times 0.22 \pm 0.11 mol Si yr ⁻¹ ^(b) m ⁻² , where a is surface area of the distal coastal zone and b is estimate of benthic efflux rate for the continental shelf	11, 12
F _{S3-rw}	Reverse weathering	1		
F _{S4S3}	bSiO ₂ dissolution	6	$F_{S3-rw} + F_{S3S1}$	

Table 3. (continued)

	Flux	Flux (Tmol yr ⁻¹)	Calculation or Explanation	Reference ^b
F _{S4-burial}	bSiO ₂ Burial	1.7	$S_4 \times 0.016 \text{ cm yr}^{-1} \text{ (a)}/20 \text{ cm} \text{ (b)}$ (so that $F_{P4\text{-burial}} + F_{S4\text{-burial}} = 3.1$), where a is sediment accumulation rate in the proximal coastal zone and b is depth of the active sediment layer in the proximal coastal zone	11, 13
		<i>Open Ocean</i>		
F _{O1O2}	DSi uptake	200	$240 \text{ Tmol Si yr}^{-1} - F_{S1S2}$	4
F _{O2O1}	bSiO ₂ dissolution	104.9	within the range of 100–120 Tmol Si yr ⁻¹ (50 ^(a) to 60 ^(b) % of F_{O1O2}), where a is fraction of bSiO ₂ production preserved in the water column (lower estimate) and b is fraction of bSiO ₂ production preserved in the water column (upper estimate)	4, 10
F _{O2O4}	Sedimentation	99.1	$F_{O1O2} + F_{am} - F_{O2O4}$	
F _{O3S1}	Coastal upwelling	10	$[O_3/V(4b)] \times W_{4-3}$	
F _{O3O1}	Oceanic upwelling	85	$F_{O4O3} + F_{O5O3} - F_{O3S1}$	
F _{O4O3}	bSiO ₂ dissolution	52.5	$(F_{O6O5} + F_{O4O3} = 38\% \text{ of } F_{O1O2} \text{ (1)})$	4
F _{O4O6}	Sedimentation	46.6	$F_{O2O4} - F_{O4O3}$	
F _{O5O3}	Deep upwelling	42.5	$F_{O6O5} + F_{O7O5}$	
F _{O6O5}	bSiO ₂ dissolution	22.6	$(F_{O6O5} + F_{O4O3} = 38\% \text{ of } F_{O1O2} \text{ (1)})$	4
F _{O6O8}	Sedimentation	24	12% of F_{O1O2}	4
F _{O7O5}	DSi efflux	19.9	within the range of $23 \pm 15 \text{ Tmol Si yr}^{-1}$	4
F _{O8O7}	Sediment bSiO ₂ dissolution	19.3	$F_{O7O5} - F_{hyd}$	
F _{O8-burial}	bSiO ₂ burial	4.7	within the range of $4.1\text{--}4.3 \text{ Tmol Si yr}^{-1} \text{ (a)}$ and $6 \text{ Tmol Si yr}^{-1} \text{ (b)}$, where a is estimate of bSiO ₂ preservation in marine sediments and b is assuming 3% preservation of opal in marine sediments	4, 13
F _{hyd}	Hydrothermal input	0.6		4

^aV and VS refer to the volumes of water and sediment reservoirs, respectively, Si/C is the molar Si to carbon ratio of primary production, M(Si) and M(C) stand for the atomic mass of Si and C, respectively.

^bReferences are as follows: Alexandre et al. [1997]; Conley [2002b]; Appelo and Postma [1993]; Tréguer et al. [1995]; Arhonditsis and Brett [2005]; Berner and Berner [1996]; Garnier et al. [2002]; Conley et al. [1989]; Dürr et al. [2009]; Nelson et al. [1995]; Rabouille et al. [2001]; Berelson et al. [1987]; De Master [2002]; Heinze et al. [1999]; Biscaye et al. [1994]; Wollast [1991]; Michalopoulos and Aller [2004].

corresponding reservoirs. In some cases, however, the reservoir mass is calculated from flux estimates, assuming steady state conditions. For example, the mass of DSi in the water column of the distal coastal ocean (72 Tmol Si) is obtained from the export flux of DSi and the water flux to the open ocean (Table 2). This estimate, combined with the reservoir volume (3600 Tm³, Figure 1), then yields an average DSi concentration of the distal ocean of 20 μM, which can be compared to the wide range of observed, depth-integrated DSi concentrations in shelf waters from <5 μM [Alvarez-Salgado et al., 1997; Gibson et al., 1997; Lacroix et al., 2007] to ~15 μM [Heiskanen and Keck, 1996] to >30 μM, [Serebrennikova and Fanning, 2004; Zhang, 2002].

[12] The ultimate source of DSi for the global Si cycle is chemical weathering of silicate rocks of the continental and oceanic crust [Gerard and Ranger, 2002]. The total mass of silicate rock exposed at the Earth's surface largely exceeds that of the reactive DSi plus bSiO₂ reservoirs. Thus, on the time scales investigated (years to centuries), the reservoir size of silicate rocks remains essentially unchanged. A large fraction of DSi released by weathering is converted by plants into phytoliths and temporarily stored in soils [Sacccone et al., 2008]; (Figure 2). A rough estimate of the reservoir mass of bSiO₂ in soils is obtained based on average phytolith concentrations for different types of soils

[Conley, 2002b], the average soil bSiO₂ concentration, and the global volume of soils. The latter is derived from the FAO world soil map [Food and Agriculture Organization/ U.N. Educational, Scientific and Cultural Organization (FAO/UNESCO), 1986], assuming a mean soil depth of 1 m [Pouba, 1968; Batjes, 1997].

[13] The sediment reservoirs correspond to the topmost layers where decomposition of biogenic constituents drives the return of dissolved nutrient species to the water column (including DSi). The corresponding volumes (VS, Table 2) are estimated by assigning a mean thickness of 10 cm to the active layer of aquatic sediments on the continents and in the proximal coastal zone [Heinze et al., 1999], and 20 cm for distal coastal zone and deep sea sediments [De Master, 2002]. An average porosity of 0.7 and an average dry density of 2.5 g cm⁻³ are assumed for all sediments [Maher et al., 2004]. The calculated sediment volumes of boxes 1, 2, 3 and 4 are 1.73, 0.18, 5.5 and 75 Tm³, respectively.

2.3. Reactive Si Fluxes

[14] The fluxes of reactive Si are obtained from the literature, or constrained by assuming that the Si cycle is initially at steady state (Table 3). The assumption of an initial steady state is a common practice in the modeling and budgeting of global elemental cycles [e.g., Mackenzie et al., 1993; Tréguer et al., 1995; Yool and Tyrrell, 2003]. It is most likely that, given the oceanic residence time of reactive

Si is 15000–17000 years [Tréguer *et al.*, 1995], the marine Si cycle was not at steady state prior to 1950, due to glacial-interglacial changes. Nonetheless, considering the time scales investigated in the simulations (≤ 150 years), these much longer-term changes have little effect on the system's response to the imposed perturbations.

[15] The fluxes include the sources and sinks of reactive Si for the Earth's surface environment (Figure 2). The sources are chemical weathering on land (F_w) and flux of DSi to the oceans resulting from basalt-seawater interactions (F_{hyd}). The sinks are burial of bSiO₂ in sediments ($F_{C5-burial}$, $F_{P4-burial}$, $F_{S4-burial}$, $F_{O8-burial}$), and removal of DSi due to reverse weathering reactions in shelf sediments (F_{S3rw}) [Mackenzie and Garrels, 1966; Michalopoulos and Aller, 1995, 2004]. Note that, because we assume an initially steady state Si cycle, the sinks and sources of reactive Si balance each other exactly.

[16] All other fluxes either transform or transport reactive Si within the Earth's surface environment and are thus internal fluxes. Si fluxes are represented by the symbol “F” followed by a subscript that identifies the initial (source) and final (sink) reservoir. The reservoir symbols are listed in Tables 1 and 2, for water and reactive Si reservoirs, respectively. Fluxes describing the uptake of DSi by organisms to produce bSiO₂ (F_{C1C2} , F_{C3C4} , F_{P1P2} , F_{S1S2} , F_{O1O2}) scale to the primary production rates in the various compartments of the earth surface environment (Table 3). Most bSiO₂ is efficiently recycled via dissolution in the water column (F_{C4C3} , F_{P2P1} , F_{S2S1} , F_{O2O1} , F_{O4O3} , F_{O6O5}), soils (F_{C2C1}) and sediments (F_{C5C3} , F_{P4P3} , F_{S4S3} , F_{O8O7}). The accumulation of DSi in the pore waters of sediments and progressive loss in the reactivity of biogenic silica surfaces (aging) ultimately allows a small fraction of bSiO₂ production to be buried and preserved in sediments [Van Cappellen *et al.*, 2002].

[17] The groundwater discharge flux of DSi to the coastal zone (F_{C1P1}) is derived from the corresponding water flux (W_{1a-2} in Figure 1) and the average DSi concentration in groundwater. The riverine supply of DSi to the proximal zone (F_{C3P1}) is derived by averaging the estimated river DSi delivery fluxes computed for 150 coastal segments in the GEMS-GLORI [Meybeck and Ragu, 1995] and GEMS-PRISRI [Meybeck *et al.*, 2003] databases, under pristine conditions, that is, prior to human perturbation [Dürr *et al.*, 2009]. The estimated river DSi flux (6.2 Tmol yr^{-1}) thus implicitly corrects for the drop in DSi concentration in the downstream reaches of rivers that has accompanied the extensive building of dams since the 1950s [Humborg *et al.*, 2006]. Note that, while the main source of reactive Si for the oceans is in the form of DSi, the contribution of riverine bSiO₂ delivery (F_{C4P2}) is far from negligible [Conley *et al.*, 2000]. Reactive Si is also supplied to the oceans via the atmosphere with eolian dust (F_{C2O2}), although this flux is most likely very small [Tréguer *et al.*, 1995] and its origin (biogenic versus mineral) remains poorly known [Cole *et al.*, 2009; Dürr *et al.*, 2009].

[18] Transport fluxes of DSi into the ocean (F_{P1S1} , F_{S1O1} , F_{O3S1} , F_{O3O1} , F_{O5O3}), as well as export fluxes of bSiO₂ from the proximal zone to the distal zone (F_{P2S2}) and from the distal zone to the open ocean (F_{S2O2}), are assumed to be directly coupled to the water cycle. That is, the flux of DSi

or bSiO₂ exiting the reservoir is related to the mass of DSi or bSiO₂ in that reservoir according to:

$$\frac{F_{ij}}{S_i} = \frac{Q_{ij}}{V_i} \quad (1)$$

where F_{ij} and Q_{ij} are the fluxes of reactive Si and water from reservoir i to reservoir j , respectively, S_i is the mass of DSi or bSiO₂ in reservoir i , and V_i is the volume of the reservoir.

[19] The remaining transport fluxes correspond to sedimentation (F_{O2O4} , F_{O4O6}) and deposition of bSiO₂ (F_{C4C5} , F_{P2P4} , F_{S2S4} , F_{O6O8}), and the efflux of DSi from sediments (F_{C5C3} , F_{P3P1} , F_{S3S1} , F_{O7O6}). In the marine realm, these fluxes are relatively well constrained by observations. Sedimentation rates and DSi fluxes from sediments can be determined directly with sediment traps and benthic chambers, respectively [Koning *et al.*, 1997; Rao and Jahnke, 2004]. Furthermore, numerous estimates of benthic DSi efflux have been calculated from measured pore water profiles [Rabouille *et al.*, 1993; Dixit and Van Cappellen, 2003].

[20] A widely used approach in biogeochemical box modeling is to relate the reservoir sizes and fluxes via linear expressions,

$$F_{ij} = k_{ij}S_i \quad (2)$$

where k_{ij} is a first-order rate constant [Lasaga, 1981; Chameides and Perdue, 1997; Mackenzie *et al.*, 1998]. Values of k_{ij} range from values of 1 yr^{-1} , for example for biological DSi uptake and bSiO₂ dissolution in aquatic environments, to values of 10^{-3} yr^{-1} or less for ground-water transport of DSi or burial of bSiO₂ in the deep-sea sediments.

2.4. Model Simulations

[21] The mass balance equations for the various reactive Si reservoirs, based on the linear flux equations (2), are solved in MATLAB using Euler's method. The steady state silica cycle represented in Figure 2 is taken as the initial condition. After verifying that the state variables exhibit no drift under time-invariant conditions, a time-dependent change in forcing is imposed, as detailed below. The system behavior is monitored for a period of 150 years, using an integration time step of 0.01 years. The starting time of the imposed perturbations is nominally set to 1950.

[22] To simulate the response of the Si cycle to a global temperature increase, three different time courses for mean surface air temperature are considered (low, medium and high; Figure 3), based on projections of the Intergovernmental Panel on Climate Change [Houghton *et al.*, 1995]. The three scenarios diverge after the year 2000. For simplicity, linear functions are used to describe the rising air temperature. Temperatures of surface waters (box 1a, box 2, box 3 and box 4a) are assumed to follow air temperature, while the magnitude of the temperature rise of the intermediate oceanic waters is four times lower [Levitov *et al.*, 2000]. The initial temperature of the intermediate water is set to 5°C [Yool and Tyrrell, 2003].

[23] The processes that are directly affected by temperature in the simulations are biological DSi uptake, bSiO₂

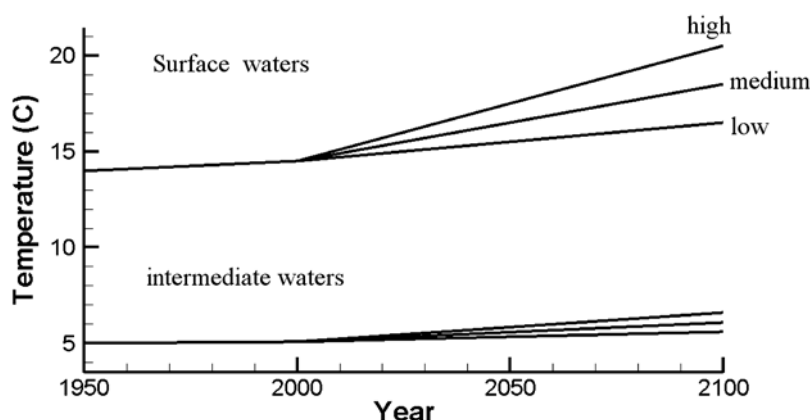


Figure 3. Temperature scenarios for surface and intermediate waters used as forcing functions in the simulations.

dissolution and chemical weathering. The model assumes that, at the spatial and temporal scales resolved, an increase (decrease) in siliceous phytoplankton production results in an increase (decrease) in DSi fixation. In particular, we assume that a temperature-dependent change in primary production by diatoms causes a proportional change in DSi uptake. The temperature dependence of DSi uptake in continental and marine environments is described by the Eppley function (uptake rate $\propto e^{(0.07 \cdot T)}$, where T is temperature in $^{\circ}\text{C}$) [Eppley, 1972]. This exponential function is widely used to describe the response of planktonic communities to temperature variations, except under extreme temperature conditions [Pasquer *et al.*, 2005]. This formulation implies the assumption that changes in DSi uptake linearly follow diatom growth. In the absence of relationships specifically describing the temperature dependence of DSi uptake by higher plants on land, we opt for a simple Q_{10} function, whereby the uptake rate doubles with every 10°C temperature increase [Winkler *et al.*, 1996]. The Arrhenius equation is used to correct the rates of bSiO₂ dissolution and silicate weathering [Lasaga, 1998]. Reported activation energies for the dissolution of framework silicates and bSiO₂ fall mostly in the range 25–90 kJ mol⁻¹ [Blum and Stillings, 1995; Van Cappellen *et al.*, 2002]. Here, a single Arrhenius activation energy of 60 kJ mol⁻¹ is imposed, to account for

the effect of temperature changes on both silicate rock weathering and bSiO₂ dissolution.

[24] Another major effect of human activity on the cycling of reactive Si at the Earth's surface is the construction of dams, known to trap large quantities of bSiO₂ [Conley *et al.*, 2000; Humborg *et al.*, 2006]. To test the sensitivity to damming, a correction coefficient is added to the flux equation describing bSiO₂ accumulation in sediments on the continents (F_{C4C5}). This coefficient is assigned a value of 1 at the start of the simulation and afterward varies proportionally with changes in the number of dams. Gleick [2003] has projected future damming pressure over the next 25 years by relating global water use to the number of new dams [Rosenberg *et al.*, 2000]. Based on this work, and the assumption that global water use is proportional to the world's population, we estimate that the number of dams should increase by 20% with an increase of the world's population by 1.9 billion people. Three damming scenarios are then considered, based on three projections for the change in the world population until the year 2100 (United Nations, World Population Prospects: The 2004 Revision Population Database, 2004). The low, medium, and high damming scenarios are shown in Figure 4.

[25] Additional effects of anthropogenic modifications of the earth surface environment are likely to affect Si cycling

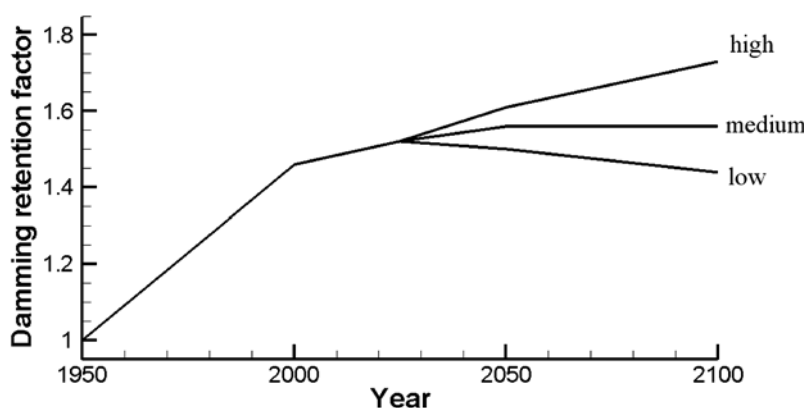


Figure 4. River damming scenarios used in the simulations. The three curves represent the relative change in damming pressure, relative to 1950, for low, medium and high damming scenarios.

along the land-to-ocean continuum. On the time scales considered here (≤ 150 years), shifts in precipitation patterns and vegetation, changes in land use and erosion will affect the cycling of Si on land and the delivery of reactive Si to the oceans [Conley *et al.*, 2008]. On even longer time scales, changes in thermohaline circulation accompanying a warming of the surface ocean will modify the exchanges of DSi between the surface and deeper parts of the oceans, thereby affecting marine biosiliceous productivity [see, e.g., Yool and Tyrrell, 2005]. A complete assessment of the response of the biogeochemical Si cycle to human-induced global change will thus require further work.

3. Results and Discussion

3.1. Global Silica Cycle

[26] Most previous global-scale mass balance studies of the Si cycle have focused on the oceans [Tréguer *et al.*, 1995; De Master, 2002; Ragueneau *et al.*, 2002; Yool and Tyrrell, 2003, 2005; De La Rocha and Bickle, 2005]. A novelty of the Si cycle presented here is that it includes an explicit representation of DSi and bSiO₂ cycling on the continents. Nevertheless, due to the relative scarcity of data, the estimates of the continental reservoir masses and fluxes have large uncertainties associated with them. For instance, the calculation of bSiO₂ stock in soils assumes an average concentration of 5 mg phytoliths per g of soil. While the latter value is consistent with the bSiO₂ determinations in soils that have been made so far [Alexandre *et al.*, 1997; Conley, 2002b; Clarke, 2003; Sferratore *et al.*, 2006], it remains to be seen how representative the relatively limited set of existing measurements is for the global soil reservoir.

[27] According to our estimates, phytoliths in soils and living terrestrial biomass constitute the largest fraction (65%) of the continental reactive Si reservoir. The amount of Si fixed by terrestrial and aquatic organisms on the continents on a yearly basis is estimated to be 89 Tmol yr⁻¹. This value lies within the range of 60–209 Tmol yr⁻¹ given in the literature [Conley, 2002b] and is of the same order of magnitude as the rate of Si fixation in the oceans (244 Tmol yr⁻¹). Thus, Si biomineralization on the continents is an important component of biological Si cycling on Earth [Conley, 2002a]. As in the marine realm, siliceous productivity on the continents relies on the efficient regeneration of nutrient DSi through bSiO₂ dissolution. In our budget, 80% of the continental bSiO₂ produced is recycled, while the remainder accumulates in lacustrine sediments and in soils [Kendrick and Graham, 2004], or is exported to the oceans. Based on the estimates in Figure 2, the residence time of reactive Si on the continents is estimated to be 775 years.

[28] Most reactive Si is delivered to the oceans by rivers under the form of DSi (F_{C1P1} ; 6.2 Tmol yr⁻¹). Nonetheless, the alternative supply routes of riverine bSiO₂ delivery (F_{C4P2} ; 1.1 ± 0.2 Tmol yr⁻¹), submarine groundwater discharge (F_{C1P1} ; 0.4 ± 0.4 Tmol yr⁻¹) and atmospheric transport (F_{C2O2} ; 0.5 ± 0.5 Tmol yr⁻¹), together contribute about 25% of the transfer of reactive Si from the continents to the oceans. When hydrothermal inputs are also included (F_{hyd} ; 0.6 ± 0.4 Tmol yr⁻¹), we estimate the total reactive Si delivery to the ocean to be 8.8 ± 1.5 Tmol yr⁻¹. In

comparison, Tréguer *et al.* [1995] estimated the total reactive Si supply to the oceans to be 6.7 ± 1.5 Tmol yr⁻¹. These authors, however, did not account for the reactive Si input from groundwater discharge and riverine bSiO₂. It should further be recognized that all regional sources of Si may not have been identified yet. For example, the venting of crustal fluids in the North Pacific has only recently been suggested to contribute as much as 1.5 ± 0.5 Tmol yr⁻¹ to the global oceanic Si budget [Johnson *et al.*, 2006].

[29] Assuming an initial steady state, the sum of the inputs to the ocean is balanced by that of the outputs, and thus burial and reverse weathering should together yield a total removal flux of 8.8 Tmol yr⁻¹. This value falls within the range for total reactive Si removal from the ocean of 8.4 to 9.4 Tmol yr⁻¹, obtained when combining the estimated sediment burial fluxes of biogenic Si of DeMaster [2002] ($7.4\text{--}8.4$ Tmol yr⁻¹) with that for reverse weathering (1.0 Tmol yr⁻¹).

[30] The explicit consideration of the proximal coastal zone enables us to account for the important filter function of estuaries, lagoons and embayments in nutrient cycles [Rabouille *et al.*, 2001; Wollast, 1993; Seitzinger and Giblin, 1996; Mackenzie *et al.*, 2000; Arndt *et al.*, 2009]. Significant Si processing decreases the DSi/bSiO₂ ratio from around 3 in rivers to 1.2 in the proximal zone. A net transformation of DSi into bSiO₂ is commonly observed in nearshore environments and causes a large fraction (43%, according to our model) of reactive Si to be delivered from the proximal to the distal coastal zone in the form of bSiO₂.

[31] The fluxes in Figure 2 emphasize the role of the continental margins in removing reactive Si by sediment burial. Although the proximal and distal zones are only responsible for about 18% of the total biological fixation of DSi in the oceans, they may account for 40% of the total marine bSiO₂ burial, in line with the assessment of DeMaster [2002]. The sum of the bSiO₂ burial fluxes in the proximal and distal coastal zones in our budget (3.1 Tmol yr⁻¹) corresponds to the maximum of the range estimated by DeMaster [2002] for bSiO₂ burial along the continental margins (2.4–3.1 Tmol yr⁻¹). It should be noted, however, that DeMaster's oceanic silica budget omits riverine supply of bSiO₂ and groundwater DSi discharge. Our relatively high estimate for bSiO₂ accumulation in coastal sediments is consistent with the inclusion of these additional reactive Si inputs to the coastal zone, as well as the upward revision of the riverine DSi supply (6.2 versus 5.6 Tmol yr⁻¹). The preferential burial of bSiO₂ in nearshore and shelf sediments is not only due to the relatively high sedimentation rates, but also to enhanced preservation resulting from interactions between deposited bSiO₂ and constituents solubilized from lithogenic minerals and the formation of new aluminosilicate phases [Van Cappellen *et al.*, 2002; Dixit and Van Cappellen, 2003; Michalopoulos and Aller, 1995, 2004]. In the proximal zone and especially in deltaic settings, there is a tight coupling between biogenic Si burial and reverse weathering, and analytical procedures for the measurement of biogenic silica account for reverse weathering products [Michalopoulos *et al.*, 2000; Michalopoulos and Aller, 1995, 2004; Presti and Michalopoulos, 2008]. Thus, the

Table 4. Reactive Silica Contents and Residence Times in the Various Compartments of the Earth Surface Environment

	DSi + bSiO ₂ (Tmol Si)	Export + Burial (Tmol Si yr ⁻¹)	Residence time (years)	Uptake/ (Export + Burial)
<i>Continents (Box 1)</i>				
Terrestrial	11310	14.6	775	0.9
Rivers and Lakes	24.1	10.1	2.1	5.5
River and Lakes + Sediment	1441.1	10.1	143	-
<i>Proximal Coastal Zone (Box 2)</i>				
Water Column	5.6	7.7	0.7	0.6
Water Column + Sediment	115.6	7.7	15	-
<i>Distal Coastal Zone (Box 3)</i>				
Water Column	104	16.3	4.9	2.5
Water Column + Sediment	2295	16.3	141	-
<i>Open Ocean (Box 4)</i>				
Water Column	96875	14.6	6635	13.7
Water Column + Sediment	147517	14.6	10104	-
<i>World Ocean (Box 2 + 3 + 4)</i>				
Water Column	96984.6	8.8	11021	27.8
Water Column + Sediment	149927.8	8.8	17037	-

Si burial flux used here for the proximal zone may include reverse weathering products.

[32] The main inflow of DSi to the distal zone is caused by coastal upwelling, which is estimated to be on the order of 10 Tmol yr⁻¹ (Figure 2). The intermediate water masses of the open ocean (100–1000 m water depth) are assumed to be the source region for coastal upwelling. This assumption is consistent with mesopelagic DSi concentrations (25–30 $\mu\text{mol kg}^{-1}$) [Dittmar and Birkicht, 2001; Brzezinski *et al.*, 1997], and Si/N ratios close to one [Hill and Wheeler, 2002; Brzezinski *et al.*, 1997] reported for coastal upwelling waters. Deeper source regions (i.e., >1000 m) would yield higher DSi concentrations and Si/N ratios between 2 and 3 [Sverdrup *et al.*, 1942]. Tréguer *et al.* [1995] estimated the whole-ocean residence time of reactive Si to be on the order of 15000 years. The latter value likely represents a lower limit, however, as these authors excluded the active surface layer of marine sediments in their calculation. According to the reservoir masses considered here, reactive Si in the water column and surface sediments of the proximal coastal zone, distal coastal zone, plus the open ocean amounts to 149927 Tmol. The removal rate of reactive Si through burial and reverse weathering of 8.8 Tmol yr⁻¹ then implies a whole-ocean residence time of 17037 years. If the proximal coastal zone is excluded, the oceanic residence time of reactive Si is 20245 years. Estimated residence times for various marine reservoirs and their combinations are summarized in Table 4.

[33] Interestingly, the residence times of reactive Si in the open ocean (10104 years) and the distal coastal zone (141 years) alone are significantly lower than the whole-ocean residence time (17037 years). This reflects the large exchange fluxes of reactive Si between the continental margins and the open ocean. For the distal zone in particular, these exchanges dominate the inputs and outputs of reactive Si and, hence, explain the relatively short residence time of reactive Si on the continental shelves.

[34] Water column residence times of reactive Si are somewhat shorter than the corresponding water residence times (Table 4), because sinking of bSiO₂ by sedimentation decouples the Si cycle from the water cycle. Nonetheless, they are significantly lower than the values obtained when

including sediments. The relative differences are particularly large for the proximal and distal coastal zone, because of the importance of benthic exchange fluxes of reactive Si. This is especially the case in the proximal coastal zone, where benthic regeneration of silica accounts for nearly one-third of the reactive Si influx and is, therefore, a major source of reactive Si sustaining siliceous productivity in the overlying water column [Ragueneau *et al.*, 2005; Laruelle *et al.*, 2009].

3.2. Sensitivity Analysis

[35] To identify the most sensitive processes controlling Si cycling across the continent-ocean transition, we compute the relative changes in water column DSi and bSiO₂ concentrations of the coastal proximal and distal zones, induced by varying the rate constants k_{ij} in the flux equations (equation (2)). In each simulation the value of one rate constant is increased by 50%, while all other model parameters remain unchanged. The sensitivity of the model to continental rock weathering (F_w) is similarly tested by increasing the value of F_w by 50%. Sensitivities are expressed as relative changes in DSi and bSiO₂ concentrations after 150 years of simulation time, relative to the initial, steady state values.

[36] The rate constants included in the sensitivity analysis correspond to the reactive Si fluxes that are not directly coupled to the water cycle via equation (1). These fluxes include uptake and biomineralization of DSi, dissolution, sedimentation, and burial of bSiO₂, plus DSi efflux across the sediment-water interface. In addition, on the time scale considered (150 years), Si cycling in the proximal zone is not affected by processes occurring in the downstream distal coastal zone and open ocean reservoirs. Hence, the sensitivity analysis for the proximal zone is further limited to rate constants k_{ij} corresponding to processes occurring in the upstream, continental reservoir or within the proximal zone itself. In contrast, Si cycling in the distal coastal zone may also be affected by processes in the downstream open ocean reservoir, because of the return of oceanic waters onto the shelves via coastal upwelling.

[37] The results of the sensitivity analysis are summarized in Table 5. DSi and bSiO₂ concentrations of proximal

Table 5. Sensitivity Analysis^a

Modified Parameter	Type of Process	Percent DSi in Box 2	Percent bSiO ₂ in Box 2	Percent DSi in Box 3	Percent bSiO ₂ in Box 3
F _w	Weathering	7	6	2	2
k _{C1C2}	Uptake	−13	−11	−3	−4
k _{C2C1}	Dissolution	14	12	3	4
k _{C3C4}	Uptake	−15	−9	−3	−4
k _{C4C3}	Dissolution	13	8	3	3
k _{C4C5}	Sedimentation	−10	−13	−3	−4
k _{C5-burial}	Burial	−1	−1	0	0
k _{C5C3}	Dissolution	5	5	1	1
k _{P1P2}	Uptake	−21	15	−2	−1
k _{P2P1}	Dissolution	5	−4	0	0
k _{P2P4}	Sedimentation	0	−15	−2	−2
k _{P3P1}	Efflux	0	0	0	0
k _{P4-burial}	Burial	−2	−2	−1	−1
k _{P4P3}	Dissolution	3	3	1	1
k _{S1S2}	Uptake	−	−	−25	11
k _{S2S1}	Dissolution	−	−	21	−10
k _{S2S4}	Sedimentation	−	−	−9	−17
k _{S3-rw}	Reverse weathering	−	−	−2	−2
k _{S3S1}	Efflux	−	−	2	2
k _{S4-burial}	Burial	−	−	−1	−1
k _{S4S3}	Dissolution	−	−	10	9
k _{O1O2}	Uptake	−	−	0	0
k _{O2O1}	Dissolution	−	−	0	0
k _{O4O3}	Dissolution	−	−	1	1
k _{O6O5}	Dissolution	−	−	0	0
k _{O6O8}	Sedimentation	−	−	0	0
k _{O7O5}	Efflux	−	−	0	0
k _{O8-burial}	Burial	−	−	0	0
k _{O8O7}	Dissolution	−	−	0	0

^aPercent change in DSi and bSiO₂ concentrations in proximal (box 2) and distal coastal zone (box 3) 150 years after increasing the corresponding flux or rate constant by 50%.

coastal waters are most sensitive to chemical weathering of continental rocks (F_w), terrestrial (F_{C1C2}), riverine (F_{C3C4}) and proximal coastal siliceous production (F_{P1P2}), and bSiO₂ dissolution on the continents (F_{C2C1} , F_{C4C3} , F_{C5C3}). Overall, enhanced production and sedimentation lead to lower DSi and bSiO₂ concentrations, while increased dissolution results in larger stocks of reactive Si in the water column. The main difference in the sensitivity of DSi and bSiO₂ concentrations in the proximal zone is related to the deposition of bSiO₂ in nearshore sediments (F_{P2P4}). While increasing k_{P2P4} causes a significant drop (−15%) of the bSiO₂ concentration in proximal waters, the DSi concentration is hardly affected. The latter reflects the fact that internal recycling of DSi via dissolution of bSiO₂ within the proximal zone (F_{P2P1} , F_{P4P3}) is much less important than continental DSi input (Figure 2). Thus, with the exception of nearshore siliceous production, reactive Si dynamics in the proximal zone are primarily controlled by processes taking place on the continents.

[38] The sensitivity analysis reveals a different picture for the distal coastal zone (Table 5). DSi and bSiO₂ concentrations in the distal coastal waters are most sensitive to internal processes. These include DSi uptake and bSiO₂ dissolution in the water column (F_{S1S2} , F_{S2S1}), as well as deposition (F_{S2S4}) and dissolution of bSiO₂ in the sediments (F_{S4S3}). Among the sensitive upstream processes, those on the continents, especially terrestrial bSiO₂ production and dissolution (F_{C1C2} , F_{C2C1}) and weathering (F_w), are more important than those in the adjacent proximal coastal zone,

although absolute changes larger than 4% are not observed. Reactive Si cycling in the distal coastal zone is also sensitive to downstream processes, foremost open ocean water column dissolution and sinking of bSiO₂ (F_{O4O3} , F_{O4O6}), as these processes control the accumulation of DSi in the source waters of coastal upwelling.

[39] On the time scale investigated, permanent removal of reactive Si through burial plays a minor role in Si cycling at the Earth's surface. This contrasts with the dominant role of sedimentary burial in the global biogeochemical Si cycle on geological time scales [De Master, 2002; Van Cappellen, 2003]. Nonetheless, even on the time scale of decades and centuries, benthic-pelagic coupling is crucial to Si cycling in distal coastal waters [Conley, 1997; Ragueneau *et al.*, 2002, 2005], as indicated by the high sensitivities to the deposition (F_{S2S4}) and subsequent benthic dissolution of bSiO₂ (F_{S4S3}).

[40] Changes in Si cycling at the land-ocean transition may also be caused by changes in the water cycle. Nevertheless, reliable scenarios for the future evolution of the water cycle are difficult to constrain. In addition to bSiO₂ retention, river damming causes a decrease of net river flow to the oceans. In the proximal coastal zone, a reduction by 20% of the river discharge (W_{1a2}) to the oceans results, after 150 years, in a relatively small decrease (4%) in the concentration of DSi, but no change in that of bSiO₂. For the distal coastal zone, the corresponding changes are 3 and 4% reductions in the concentrations of DSi and bSiO₂. The latter concentrations are further reduced (by 5 and 4%, respectively) when the decrease in river discharge by 20% is

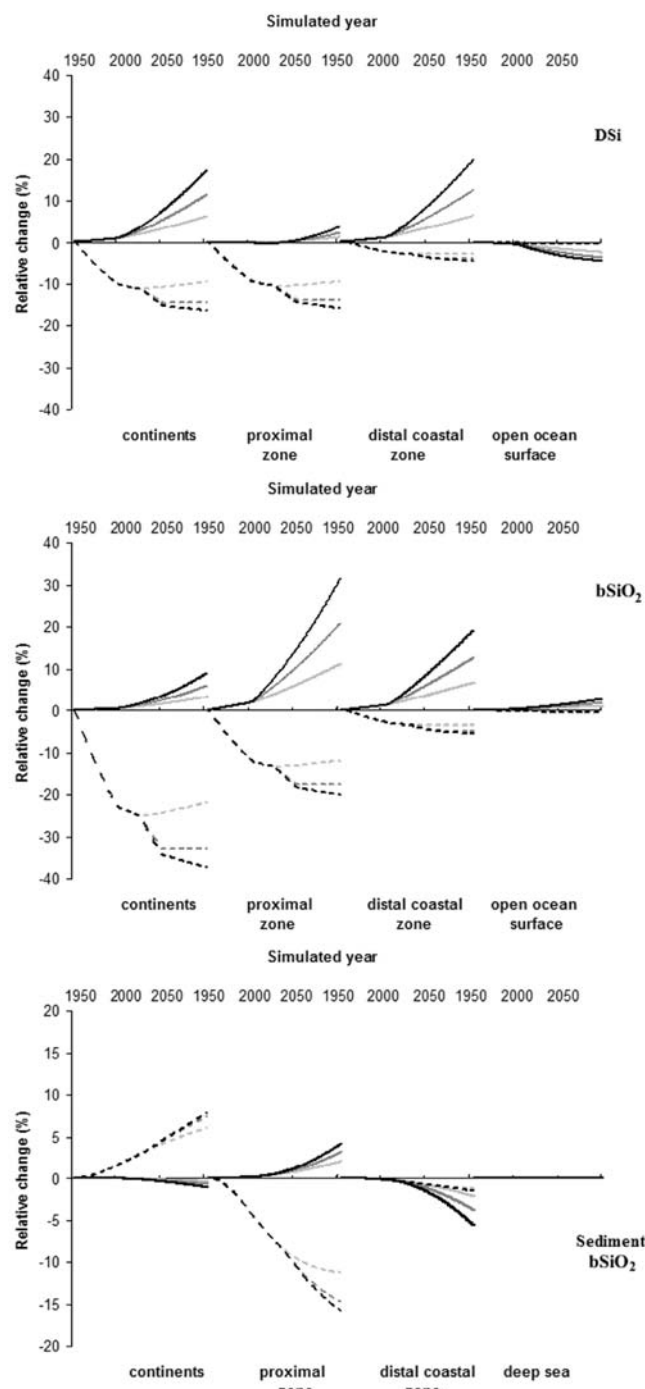


Figure 5. Relative changes in (top) DSi, (middle) water column bSiO₂, and (bottom) sediment bSiO₂ reservoir sizes versus time for the three different temperature (continuous lines) and damming (dashed lines) scenarios. In black, the high temperature or damming scenario, in gray the medium temperature or damming scenario, and in light gray the low temperature or damming scenario.

accompanied by a 20% reduction in the coastal upwelling water flux (W_{43}). On longer time scales, changes in the water cycle (e.g., upwelling) may have significantly larger effects on global Si cycling [e.g., Yool and Tyrrell, 2005].

3.3. Applications

3.3.1. Temperature Rise

[41] The three temperature scenarios yield the same general trends, but with different magnitudes (Figure 5). Except for open surface ocean DSi, increasing temperatures result in higher water column DSi and bSiO₂ concentrations. bSiO₂ concentrations in sediments of the continents and deep sea are hardly affected, while they show opposite trends in the proximal (increase) and distal coastal ocean (decrease).

[42] Rising concentrations of DSi and bSiO₂ in the continental aquatic environment (box 1a, Figure 2) reflect enhanced DSi fluxes from weathering and bSiO₂ dissolution in soils. In combination with the relatively small volume of the aquatic environment, this causes significant, and parallel, increases in the DSi and bSiO₂ concentrations. Because of more rapid bSiO₂ dissolution kinetics, the additional reactive Si mobilized does not accumulate in sediments and soils, but is exported to the ocean.

[43] Increased continental supply of reactive Si enhances siliceous productivity in proximal and distal coastal ecosystems. The largest relative change is observed in the bSiO₂ concentration of nearshore waters. That is, increased temperatures further decrease the water column DSi/bSiO₂ ratio of the proximal zone. The higher-deposition flux of bSiO₂ offsets the faster bSiO₂ dissolution kinetics, resulting in a net increase in the bSiO₂ concentration of proximal coastal sediments. Sediment bSiO₂ concentrations in the distal coastal zone show an opposite response, reflecting the very different recycling efficiencies of reactive Si in the two coastal systems (Figure 2). The latter is also reflected in the relative increases in water column DSi and bSiO₂ concentrations. In the proximal zone, rising temperatures cause a larger relative increase in bSiO₂, compared to DSi, while the reverse is observed for the distal zone. Possible indicators of a global warming effect on the Si cycle thus include opposite changes of water column DSi/bSiO₂ ratios and sediment bSiO₂ accumulation rates in proximal versus distal coastal environments.

[44] Because of its much larger volume and of the buffering effect of the coastal zone, the open ocean exhibits much smaller modifications in Si cycling. The most pronounced changes are in the surface ocean, as it directly experiences changes in air temperature. Both DSi uptake and bSiO₂ dissolution rates are enhanced by rising temperature, resulting in faster Si turnover in the upper ocean. For the model structure and parameter values used here, the net effect is a slight increase in the bSiO₂ standing stock, at the expense of the DSi pool.

3.3.2. River Damming

[45] River damming leads to opposite trends of DSi and bSiO₂ concentrations compared to those of temperature rise, with the exception of bSiO₂ in distal coastal sediments (Figure 5, broken lines). Predicted concentration changes also imply that damming should increase water column DSi/bSiO₂ ratios in continental aquatic systems and the proximal coastal zone. A number of studies have indeed reported measurable increases in DSi/bSiO₂ ratios of riverine and nearshore waters [Conley, 1997, 2002b; Friedl and Wüest, 2002].

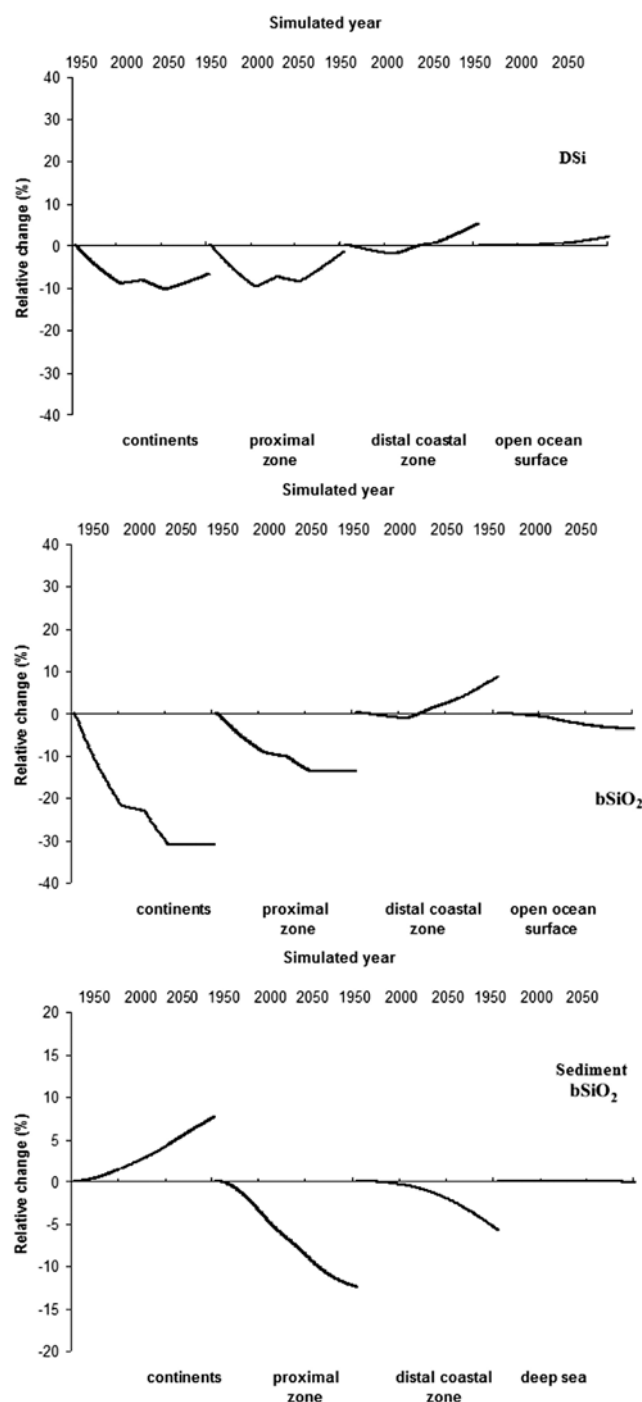


Figure 6. Relative changes in (top) DSi, (middle) water column bSiO₂, and (bottom) sediment bSiO₂ reservoir sizes versus time for the combined scenario with high damming and medium temperature forcing.

[46] As expected, the sedimentary bSiO₂ pool on the continents increases as biosiliceous debris accumulates behind the growing number of dams. The decreased continental supply of reactive Si causes a relative drop in sediment bSiO₂ in nearshore marine sediments and an increase on the continents. It should be borne in mind,

however, that the reservoir size of proximal coastal zone sediment bSiO₂ (110 Tmol Si) is much smaller than that of continental sediments (1417 Tmol Si). At the scale investigated in our simulations, damming causes more bSiO₂ to be trapped in continental sediments than lost from proximal sediments.

[47] A general feature of the response of Si cycling to changes in damming is the “dilution” of the relative effects from rivers to the open ocean (Figure 5). Typically, the largest changes in reactive Si concentrations are observed in aquatic systems on land and in the proximal coastal zone, while the open ocean system experiences little changes. For example, the open ocean surface DSi concentration is predicted to change by less than 1% after 150 years, even for the highest damming scenario.

[48] According to the damming scenarios, the largest increase in the number of dams should have taken place between the 1950s and the present. After 2000, the different damming scenarios hypothesize the same decreasing damming pressure until 2025, after which the three scenarios diverge (Figure 4). The time-dependent features imposed to the damming scenarios are recorded nearly instantaneously by the water column DSi and bSiO₂ concentrations in the continental aquatic system and the proximal coastal zone, because of the correspondingly very short residence times of reactive Si (2.1 and 0.7 years, Table 4). This is not the case for the sediment bSiO₂ concentrations in the same reservoirs. Because of much longer residence, and hence response times, the initial trends (i.e., for the period 1950–2000) are projected into the future with still little differentiation in sediment bSiO₂ concentrations among the scenarios by the year 2050.

3.3.3. Combined River Damming and Temperature Rise

[49] A set of nine permutations of the three temperature and three damming scenarios were run. The results are illustrated in Figure 6 for the intermediate temperature rise plus maximum damming scenario. This particular scenario was selected because it exhibits the main features observed in all the various simulations. The comparison between Figures 5 and 6 suggests that river damming is driving the changes in Si cycling in the combined scenario, particularly during the initial period (1950–2000). On the continents, the results of the combined simulation closely follow those of the damming-only simulation, except for the slight rise in water column DSi concentration simulated beyond 2050. The slowing down of dam construction after 2000 and continued rise in temperature cause much more pronounced reversals in the water column DSi concentrations of the coastal ocean. By the end of the simulation, the DSi concentration in the proximal zone has returned to within 2% of its starting value, while in the distal zone the DSi concentration increases above the initial value.

[50] The water column bSiO₂ concentration in the proximal coastal zone mainly records the changing damming pressure. This is no longer the case for the distal coastal zone, since reactive Si cycling in this reservoir is largely driven by internal recycling, via temperature-dependent DSi uptake and bSiO₂ dissolution. Thus, as far as the water column DSi and bSiO₂ concentrations are concerned, when

moving from the continents to the open ocean, the relative influence of changes in river damming decreases, while that of global warming increases.

[51] Changes in sediment bSiO_2 concentrations in the combined scenario are essentially the same as in the damming-only scenario, for all reservoirs. The lack of temperature-induced changes reflects the much longer residence times of reactive Si in the sediment reservoirs, relative to the water column. That is, over the 150 years of simulation time, the sediments only record the initial rapid growth in the number of dams after the 1950s. The observed loss of sediment bSiO_2 in coastal environments over the simulation period is therefore mainly due to reactive Si retention by dams.

[52] The results presented in Figures 5 and 6 illustrate the complex response of Si cycling to human influences. In particular, the temporal trends in DSi and bSiO_2 concentrations in the combined damming plus temperature rise scenario are not simply the sum of the individual responses to the two perturbations. Nonetheless, the results also indicate that by combining temporal trends in DSi and bSiO_2 concentrations in multiple reservoirs it may become possible to extricate the relative effects of the different anthropogenic forcings acting on the biogeochemical Si cycle.

4. Conclusions

[53] Silicon is a key nutrient whose biogeochemical cycling is closely coupled to those of carbon, nitrogen, phosphorus, iron and trace compounds. Large amounts of dissolved Si (DSi) are biologically fixed annually as biogenic silica (bSiO_2), both on land (89 Tmol yr^{-1}) and in the oceans (240 Tmol yr^{-1}). The estimated residence time of reactive Si on the continents (775 years), however, is about 20 times smaller than for the oceans (17037 years), reflecting the much larger marine reservoir of reactive Si.

[54] While reactive Si is mainly exported from the continents as riverine DSi (6.2 Tmol yr^{-1}), a non-negligible fraction is delivered to the oceans as bSiO_2 in river suspended matter and in eolian dust deposits (1.6 Tmol yr^{-1}), and as DSi in submarine groundwater discharge (0.4 Tmol yr^{-1}). Because of the net transformation of DSi into bSiO_2 in nearshore waters, nearly half (43%) of land-derived reactive Si reaching the distal coastal zone is in the form of bSiO_2 . Nevertheless, the major input of reactive Si to the continental shelves is via coastal upwelling.

[55] The coastal ocean represents a dynamic interface between the continents and the open ocean. Although coastal and shelf environments account for only 18% of all biological Si fixation in the oceans, an estimated 40% of all marine bSiO_2 burial takes place in nearshore and shelf sediments (3.2 Tmol yr^{-1}). Nearshore ecosystems also attenuate the downstream propagation of land-based perturbations of the Si cycle that results from damming of rivers or land use changes.

[56] The biogeochemical Si cycle is currently undergoing significant modifications due to human activities. The proposed model can help delineate the expected changes, through sensitivity analyses and scenario simulations. A major difficulty is that multiple anthropogenic perturbations

are simultaneously acting on the Si cycle. The results presented here, and observed changes in rivers and near-shore waters, indicate that riverine export of reactive Si, and the riverine bSiO_2/DSi ratio, are likely to continue to drop in the near future, as a result of reactive Si retention by dams. However, enhanced bSiO_2 dissolution due to global warming may ultimately allow coastal siliceous productivity to recover from the downward trend caused by river damming.

[57] Our work represents a first step toward modeling the global biogeochemical Si cycle along the entire land to ocean continuum. Further progress will rely especially on the increased understanding and characterization of the spatial heterogeneity of continental Si cycling, caused by differences in lithology, vegetation, land use and hydrology. This information is needed to account for the regional variability in reactive Si delivery to the coastal zone by rivers and submarine groundwater discharge.

[58] **Acknowledgments.** This project was initiated during the 2004 Summer School of the EU-funded Research Training Network SiWEBS (contract HPRN-CT-2002-000218). Further work on the project was made possible by financial support from the EU (SiWEBS), Utrecht University (High Potential project G-NUX to C.P. Slomp), and the Netherlands Organisation for Scientific Research (NWO Pioneer grant to P. Van Cappellen). We thank the Editor and the anonymous reviewers for their useful and constructive comments.

References

- Alexandre, A., J.-D. Meunier, F. Colin, and J. M. Koud (1997), Plant impact on the biogeochemical cycle of silicon and related weathering processes, *Geochim. Cosmochim. Acta*, **61**, 677–682, doi:10.1016/S0016-7037(97)00001-X.
- Alvarez-Salgado, X. A., C. G. Castro, F. F. Pérez, and F. Fraga (1997), Nutrient mineralization patterns in shelf waters of the western Iberian upwelling, *Cont. Shelf Res.*, **17**, 1247–1270, doi:10.1016/S0278-4343(97)00014-9.
- Anikouchine, W. A., and R. W. Sternberg (1981), *The World Ocean: An Introduction to Oceanography*, Prentice-Hall, Upper Saddle River, N. J.
- Appelo, C. A. J., and D. Postma (1993), *Geochemistry, Groundwater and Pollution*, Balkema, Amsterdam.
- Arhonditsis, G. E., and M. T. Brett (2005), Eutrophication model for Lake Washington (USA). Part II: Model calibration and system dynamics analysis, *Ecol. Modell.*, **187**, 179–200, doi:10.1016/j.ecolmodel.2005.01.039.
- Arndt, S., P. Regnier, and J.-P. Vanderborght (2009), Seasonally-resolved nutrient filtering capacities and export fluxes in a macrotidal estuary, *J. Mar. Syst.*, **78**, 42–58.
- Bartoli, F. (1983), The biogeochemical cycle of silicon in two temperate forest ecosystems, *Ecol. Bull.*, **35**, 469–476.
- Batjes, N. H. (1997), A world data set of derived soil properties by FAO UNESCO soil unit for global modeling, *Soil Use Manage.*, **13**, 9–16, doi:10.1111/j.1475-2743.1997.tb00550.x.
- Berelson, W. M., D. E. Hammond, and K. S. Johnson (1987), Benthic fluxes and the cycling of biogenic silica and carbon in the two southern California Borderland Basins, *Geochim. Cosmochim. Acta*, **51**, 1345–1363, doi:10.1016/0016-7037(87)90320-6.
- Berner, E. A., and R. A. Berner (1996), *Global Environment: Water, Air and Geochemical Cycles*, Prentice-Hall, Upper Saddle River, N. J.
- Biscaye, P. E., C. N. Flagg, and P. G. Falkowski (1994), The shelf edge exchange processes experiment, SEEP-II: An introduction to hypotheses, results and conclusions, *Deep Sea Res., Part II*, **41**, 231–252, doi:10.1016/0967-0645(94)90022-1.
- Blum, A. E., and L. L. Stillings (1995), Feldspar dissolution kinetics, in *Chemical Weathering Rates of Silicate Minerals*, edited by A. E. White and S. L. Brantley, pp. 291–351, Mineral. Soc. of Am., Washington, D. C.
- Bonn, W. J. (1995), Biogenic opal and barium: Indicators for late Quaternary changes in productivity at the Antarctic continental margin, Atlantic Sector, *Rep. Polar Res.*, **180**, 186 pp., Alfred Wegener Inst. for Polar and Mar. Res., Bremerhaven.
- Brink, K. H., F. F. G. Abrantes, P. A. Bernal, R. C. Dugdale, M. Estrada, L. Hutchings, R. A. Jahnke, P. J. Muller, and R. L. Smith

- (1995), Group report: How do coastal upwelling systems operate as integrated physical, chemical, and biological systems and influence the geological record?, in *Upwelling in the Ocean: Modern Processes and Ancient Records*, edited by C. P. Summerhayes et al., pp. 103–124, Wiley, New York.
- Broecker, W. S., and T.-H. Peng (1982), *Tracers in the Sea*, Eldigio Press, Palisades, New York.
- Brzezinski, M. A., D. R. Phillips, F. P. Chavez, G. E. Friederich, and R. C. Dugdale (1997), Silica production in the Monterey, California, upwelling system, *Limnol. Oceanogr.*, **42**, 1694–1705.
- Chameides, W. L., and E. M. Perdue (1997), *Biogeochemical Cycles: A Computer-Interactive Study of Earth System Science and Global Change*, Oxford Univ. Press, New York.
- Chauvaud, L., F. Jean, O. Ragueneau, and G. Thouzeau (2000), Long-term variation of the Bay of Brest ecosystem: Benthic–pelagic coupling revisited, *Mar. Ecol. Prog. Ser.*, **200**, 35–48, doi:10.3354/meps200035.
- Clarke, J. (2003), The occurrence and significance of biogenic opal in the regolith, *Earth Sci. Rev.*, **60**, 175–194, doi:10.1016/S0012-8252(02)00092-2.
- Cole, J. M., S. L. Goldstein, P. B. deMenocal, S. R. Hemming, and F. E. Grousset (2009), Contrasting compositions of Saharan dust in the eastern Atlantic Ocean during the last deglaciation and African Humid Period, *Earth Planet. Sci. Lett.*, **278**, 257–266, doi:10.1016/j.epsl.2008.12.011.
- Conkright, M., S. Levitus, and T. P. Boyer (1994), *NOAA Atlas NESDIS 1. World Ocean Atlas 1994*, vol. 1, *Nutrients*, U.S. Gov. Print. Off., Washington, D. C.
- Conley, D. J. (1988), Biogenic silica as an estimate of siliceous microfossil abundance in Great-Lakes sediments, *Biogeochemistry*, **6**(3), 161–179, doi:10.1007/BF02182994.
- Conley, D. J. (1997), Riverine contribution of biogenic silica to the oceanic silica budget, *Limnol. Oceanogr.*, **42**, 774–777.
- Conley, D. J. (2002a), The biogeochemical silica cycle: Elemental to global scales, *Oceanis*, **28**, 353–368.
- Conley, D. J. (2002b), Terrestrial ecosystems and the global biogeochemical silica cycle, *Global Biogeochem. Cycles*, **16**(XX), 1121, doi:10.1029/2002GB001894.
- Conley, D. J., and T. C. Malone (1992), Annual cycle of dissolved silicate in Chesapeake Bay—implications for the production and fate of phytoplankton biomass, *Mar. Ecol. Prog. Ser.*, **81**, 121–128, doi:10.3354/meps081121.
- Conley, D. J., and C. L. Schelske (2001), Biogenic silica, in *Tracking Environmental Changes Using Lake Sediments: Biological Methods and Indicators*, edited by J. P. Smol, H. J. B. Birks, and W. M. Last, pp. 281–293, Kluwer Acad., Dordrecht.
- Conley, D. J., S. S. Kilham, and E. Theriot (1989), Differences in silica content between marine and freshwater diatoms, *Limnol. Oceanogr.*, **34**, 205–213.
- Conley, D. J., C. L. Schelske, and E. F. Stoermer (1993), Modification of the biogeochemical cycle of silica with eutrophication, *Mar. Ecol. Prog. Ser.*, **101**, 179–192, doi:10.3354/meps101179.
- Conley, D. J., P. Stålnacke, H. Pitkänen, and A. Wilander (2000), The transport and retention of dissolved silicate from rivers in Sweden and Finland, *Limnol. Oceanogr.*, **45**, 1850–1853.
- Conley, D. J., G. E. Likens, D. Buso, L. Saccone, S. W. Bailey, and C. Johnson (2008), Deforestation causes increased dissolved silicate losses in the Hubbard Brook Experimental Forest, *Global Change Biol.*, **14**, 2548–2554.
- Cossins, A. R., and K. Bowler (1987), *Temperature Biology of Animals*, Chapman and Hall, New York.
- Datnoff, L. E., G. H. Snyder, and G. H. Komdörfer (2001), *Silicon in Agriculture*, edited by L. E. Datnoff, G. H. Snyder, and G. H. Komdörfer, Elsevier Sci., Amsterdam.
- Del Amo, Y., and M. A. Brzezinski (1999), The chemical form of dissolved Si taken up by marine diatoms, *J. Phycol.*, **35**(6), 1162–1170, doi:10.1046/j.1529-8817.1999.3561162.x.
- De La Rocha, C. L., and M. J. Bickle (2005), Sensitivity of silicon isotopes to whole-ocean changes in the silica cycle, *Mar. Geol.*, **217**, 267–282, doi:10.1016/j.margeo.2004.11.016.
- DeMaster, D. J. (2002), The accumulation and cycling of biogenic silica in the Southern Ocean: Revisiting the marine silica budget, *Deep Sea Res., Part II*, **49**, 3155–3167, doi:10.1016/S0967-0645(02)00076-0.
- Dittmar, T., and M. Birkicht (2001), Regeneration of nutrients in the northern Benguela upwelling and the Angola-Benguela Front areas, *S. Afr. J. Sci.*, **97**(5–6), 239–246.
- Dixit, S., and P. Van Cappellen (2003), Predicting benthic fluxes of silicic acid from deep-sea sediments, *J. Geophys. Res.*, **108**(C10), 3334, doi:10.1029/2002JC001309.
- Dürr, H. H., M. Meybeck, J. Hartmann, G. G. Laurelle, and V. Roubex (2009), Global spatial distribution of natural riverine silica inputs to the coastal zone, *Biogeosci. Disc.*, **6**, 1–57.
- EGGE, J. K., and D. L. Aksnes (1992), Silicate as a regulating nutrient in phytoplankton competition, *Mar. Ecol. Prog. Ser.*, **83**, 281–289, doi:10.3354/meps083281.
- Eppley, R. W. (1972), Temperature and phytoplankton growth in the sea, *Fish. Bull.*, **70**, 1063–1085.
- Epstein, E. (1999), Silicon, *Annu. Rev. Plant Physiol. Plant Mol. Biol.*, **50**, 641–664, doi:10.1146/annurev.arplant.50.1.641.
- Fekete, B. M., C. J. Vörösmarty, and W. Grabs (2002), High-resolution fields of global runoff combining observed river discharge and simulated water balances, *Global Biogeochem. Cycles*, **16**(3), 1042, doi:10.1029/1999GB001254.
- Food and Agriculture Organization/U.N. Educational, Scientific and Cultural Organization (FAO/UNESCO) (1986), Gridded FAO/UNESCO soil units: UNEP/GRID, FAO soil map of the world in digital form, digital raster data on 2-minute geographic (lat × lon) 5400 × 10800 grid, Carouge, Switzerland.
- Friedl, G., and A. Wüest (2002), Disrupting biogeochemical cycles: Consequences of damming, *Aquat. Sci.*, **64**, 55–65, doi:10.1007/s00027-002-8054-0.
- Friedl, G., C. Teodoru, and B. Wehrli (2004), Is the Iron Gate I reservoir on the Danube River a sink for dissolved silica?, *Biogeochemistry*, **68**, 21–32, doi:10.1023/B:BIOG.0000025738.67183.c0.
- Garnier, J., A. D'Aygués, J. Billen, D. J. Conley, and A. Sferratore (2002), Silica dynamics in the hydrographic network of the Seine River, *Oceanis*, **28**, 487–508.
- Garrels, R. M., and F. T. Mackenzie (1971), *Evolution of Sedimentary Rocks*, edited by W. W. Norton, New York.
- Gerard, F., and J. Ranger (2002), Silicate weathering mechanisms in a forest soil, *Oceanis*, **28**, 384–415.
- Gibson, C. E., B. M. Stewart, and R. J. Gowen (1997), A synoptic study of nutrients in the north-west Irish Sea, *Estuar. Coastal Shelf Sci.*, **45**, 27–38, doi:10.1006/ecss.1996.0164.
- Gleick, P. H. (2003), Global freshwater resources: Soft-path solutions for the 21st Century, *Science*, **302**, 1524–1528, doi:10.1126/science.1089967.
- Heinze, C., E. Maier-Reimer, A. M. E. Winguth, and D. Archer (1999), A global oceanic sediment model for long-term climate studies, *Global Biogeochem. Cycles*, **13**(1), 221–250, doi:10.1029/98GB02812.
- Heiskanen, A. S., and A. Keck (1996), Distribution and sinking rates of phytoplankton, detritus, and particulate biogenic silica in the Laptev Sea and Lena River (Arctic Siberia), *Mar. Chem.*, **53**, 229–245, doi:10.1016/0304-4203(95)00091-7.
- Hill, J. K., and P. A. Wheeler (2002), Organic carbon and nitrogen in the northern California current system: Comparison of offshore, river plume and coastally upwelled water, *Prog. Oceanogr.*, **53**, 369–387, doi:10.1016/S0079-6611(02)00037-X.
- Houghton, J. T., L. G. Meira Filho, B. A. Callender, N. Harris, A. Kattenberg, and K. Maskell (Eds) (1995), *Climate Change 1995: Contribution of Working Group I to the Second Assessment Report of the Intergovernmental Panel on Climate Change*, 572 pp., Cambridge Univ. Press, U. K.
- Humborg, C., D. J. Conley, L. Rahm, F. Wulff, A. Cociasu, and V. Ittekkot (2000), Silicon retention in river basins: Far-reaching effects on biogeochemistry and aquatic food webs in coastal marine environments, *Ambio*, **29**(1), 45–51.
- Humborg, C., M. Pastuszak, J. Aigars, H. Siegmund, C. M. Morth, and V. Ittekkot (2006), Decreased silica land-sea fluxes through damming in the Baltic Sea catchment: Significance of particle trapping and hydrological alterations, *Biogeochemistry*, **77**(2), 265–281, doi:10.1007/s10533-005-1533-3.
- Jahnke, R. A., S. R. Emerson, and J. W. Murray (1982), A model of oxygen reduction, denitrification, and organic matter mineralization in marine sediments, *Limnol. Oceanogr.*, **27**, 610–630.
- Johnson, H. P., S. L. Hautala, T. A. Björklund, and M. R. Zarnetske (2006), Quantifying the North Pacific silica plume, *Geochem. Geophys. Geosyst.*, **7**, Q05011, doi:10.1029/2005GC001065.
- Kendrick, K. J., and R. C. Graham (2004), Pedogenic silica accumulation in chronosequence soils, southern California, *Soil Sci. Soc. Am. J.*, **68**, 1295–1303.
- Koning, E., G.-J. Brummer, W. Van Raaphorst, J. Van Bennekom, W. Helder, and J. Van Iperen (1997), Settling, dissolution and burial of biogenic silica in the sediments off Somalia (northwestern Indian Ocean), *Deep Sea Res., Part II*, **44**, 1341–1360, doi:10.1016/S0967-0645(97)00018-0.
- Lacroix, G., K. Ruddick, Y. Park, N. Gypens, and C. Lancelot (2007), Validation of the 3D biogeochemical model MIROCO with field nutrient and phytoplankton data and MERIS-derived surface chloro-

- phyll a images, *J. Mar. Syst.*, 64(1–4), 66–88, doi:10.1016/j.marsys.2006.01.010.
- Laruelle, G. G., P. Regnier, O. Ragueneau, M. Kempa, B. Moriceau, S. Ni Longphuiert, A. Leynaert, G. Thouzeau, and L. Chauvaud (2009), Benthic-pelagic coupling and the seasonal silica cycle in the bay of Brest (France): New insights from a coupled physical-biological model, *Mar. Ecol. Progr. Ser.*, 385, 15–32, doi:10.3354/meps07884.
- Lasaga, A. C. (1981), Rate laws of chemical reactions, in *Kinetics of Geochemical Processes*, vol. 8, edited by A. C. Lasaga and R. J. Kirkpatrick, pp. 1–169, Mineral. Soc. of Am., Washington, D. C.
- Lasaga, A. C. (1998), *Kinetic Theory in the Earth Sciences*, Princeton Univ. Press, Princeton, N. J.
- Ledford-Hoffman, P. A., D. J. DeMaster, and C. A. Nittrouer (1986), Biogenic silica accumulation in the Ross Sea and the importance of Antarctic continental-shelf deposits in the marine silica budget, *Geochim. Cosmochim. Acta*, 50, 2099–2110, doi:10.1016/0016-7037(86)90263-2.
- Levitus, S., J. I. Antonov, T. P. Boyer, and C. Stephens (2000), Warming of the world ocean, *Science*, 287, 2225–2229, doi:10.1126/science.287.5461.2225.
- Leynaert, A., P. Tréguer, and C. Lancelot (2001), Silicic acid limitation of Equatorial Pacific diatom populations: Evidence from ^{32}Si kinetic experiments, *Deep Sea Res.*, 48, 639–660, doi:10.1016/S0967-0637(00)00044-3.
- Macdonald, A. M. (1998), The global ocean circulation: A hydrographic estimate and regional analysis, *Prog. Oceanogr.*, 41, 281–382, doi:10.1016/S0079-6611(98)00020-2.
- Mackenzie, F. T., and R. A. Garrels (1966), Chemical mass balance between rivers and oceans, *Am. J. Sci.*, 264, 507–525.
- Mackenzie, F. T., L. M. Ver, C. Sabine, M. Lane, and A. Lerman (1993), C, N, P, S global biogeochemical cycles and modeling of global change, in *Interactions of C, N, P and S Biogeochemical Cycles and Global Change*, edited by R. Wollast, F. T. Mackenzie, and L. Chou, pp. 1–62, Springer, New York.
- Mackenzie, F. T., A. Lerman, and L. M. Ver (1998), Role of the continental margin in the global carbon balance during the past three centuries, *Geology*, 26, 423–426, doi:10.1130/0091-7613[1998]026<0423:ROTCMI>2.3.CO;2.
- Mackenzie, F. T., L. M. Ver, and A. Lerman (2000), Coastal-zone biogeochemical dynamics under global warming, *Int. Geol. Rev.*, 42, 193–206, doi:10.1080/00206810009465077.
- Maher, K., D. J. DePaolo, and J. C.-F. Lin (2004), Rates of silicate dissolution in deep-sea sediment: In situ measurement using $^{234}\text{U}/^{238}\text{U}$ of pore fluids, *Geochim. Cosmochim. Acta*, 68(22), 4629–4648, doi:10.1016/j.gca.2004.04.024.
- Meunier, J.-D., F. Colin, and C. Alarcon (1999), Biogenic silica storage in soils, *Geology*, 27(9), 835–838, doi:10.1130/0091-7613[1999]027<0835:BSSIS>2.3.CO;2.
- Meybeck, M., and A. Ragu (1995), *River Discharges to Oceans: An Assessment of Suspended Solids, Major Ions and Nutrients*, report, U.N. Environ. Programme, Nairobi.
- Meybeck, M., L. Laroche, H. H. Dürr, and J. P. M. Syvitski (2003), Global variability of daily total suspended solids and their fluxes in rivers, *Global Planet. Change*, 39(1–2), 65–93, doi:10.1016/S0921-8181(03)00018-3.
- Michalopoulos, P., and R. C. Aller (1995), Rapid clay mineral formation in Amazon delta sediments: Reverse weathering and oceanic elemental fluxes, *Science*, 270, 614–617, doi:10.1126/science.270.5236.614.
- Michalopoulos, P., and R. C. Aller (2004), Early diagenesis of biogenic silica in the Amazon delta: Alteration, authigenic clay formation, and storage, *Geochim. Cosmochim. Acta*, 68, 1061–1085, doi:10.1016/j.gca.2003.07.018.
- Michalopoulos, P., R. C. Aller, and R. J. Reeder (2000), Conversion of diatoms to clays during early diagenesis in tropical, continental shelf muds, *Geology*, 28, 1095–1098, doi:10.1130/0091-7613[2000]28<1095:CODTCD>2.0.CO;2.
- Nelson, D. M., P. Tréguer, M. A. Brzezinski, A. Leynaert, and B. Queguiner (1995), Production and dissolution of biogenic silica in the ocean: Revised global estimates, comparison with regional data and relationship to biogenic sedimentation, *Global Biogeochem. Cycles*, 9, 359–372, doi:10.1029/95GB01070.
- Paasche, E. (1980), Silicon, in *The Physiological Ecology of Phytoplankton*, edited by I. Morris, pp. 259–284, Blackwell Sci. Publ., Oxford, U. K.
- Pasquer, B., G. Laurelle, S. Bequevort, V. Schoemann, H. Goosse, and C. Lancelot (2005), Linking ocean biochemical cycles and ecosystem structure and function: Results of the complex SWAMCO model, *J. Sea Res.*, 53, 93–108, doi:10.1016/j.seares.2004.07.001.
- Piperno, D. R. (1988), *Phytolith Analysis—An Archeological and Geological Perspective*, 280 pp., Academic, London.
- Pouba, Z. (1968), *Geologische Kartierung* (in Czech), Naklad Ceskoslovenske Akad. Ved., Prague.
- Presti, M., and P. Michalopoulos (2008), Estimating the contribution of the authigenic mineral component to the long-term reactive silica accumulation on the western shelf of the Mississippi River Delta, *Cont. Shelf Res.*, 28, 823–838, doi:10.1016/j.csr.2007.12.015.
- Rabouille, C., P. Crassous, A. Kripounoff, J.-F. Gaillard, R. Jahnke, C. Pierre, and J.-C. Relexans (1993), A model of early diagenesis in the tropical North Atlantic: Processes and mass balances in the sediments of the EUMELI program, *Chem. Geol.*, 107, 463–466, doi:10.1016/0009-2541(93)90232-8.
- Rabouille, C., F. T. Mackenzie, and L. M. Ver (2001), Influence of the human perturbation on carbon, nitrogen, and oxygen biogeochemical cycles in the global coastal ocean, *Geochim. Cosmochim. Acta*, 65(21), 3615–3641, doi:10.1016/S0016-7037(01)00760-8.
- Ragueneau, O., N. Dittert, L. Corrin, P. Tréguer, and P. Pondaven (2002), Si:C decoupling in the world ocean: Is the Southern Ocean different?, *Deep Sea Res., Part II*, 49(16), 3127–3154, doi:10.1016/S0967-0645(02)00075-9.
- Ragueneau, O., L. Chauvaud, B. Moriceau, A. Leynaert, G. Thouzeau, A. Donval, F. Le Loc'h, and F. Jean (2005), Biodeposition by an invasive suspension feeder impacts the biogeochemical cycle of Si in a coastal ecosystem (Bay of Brest, France), *Biogeochemistry*, 75, 19–41, doi:10.1007/s10533-004-5677-3.
- Ragueneau, O., S. Schultes, K. Bidle, P. Claquin, and B. Moriceau (2006a), Si and C Interactions in the world ocean: Importance of ecological processes and implications for the role of diatoms in the biological pump, *Global Biogeochem. Cycles*, 20, GB4S02, doi:10.1029/2006GB002688.
- Ragueneau, O., D. J. Conley, S. Ni Longphuiert, C. P. Slomp, and A. Leynaert (2006b), A review of the Si biogeochemical cycle in coastal waters, I. Diatoms in coastal food webs and the coastal Si cycle, in *Land-Ocean Nutrient Fluxes: Silica Cycle*, SCOPE 66, edited by V. Ittekkot, C. Humborg, and J. Garnier, pp. 163–195, Island Press, Washington, D. C.
- Ragueneau, O., D. J. Conley, S. Ni Longphuiert, C. P. Slomp, and A. Leynaert (2006c), A review of the Si biogeochemical cycle in coastal waters, II. Anthropogenic perturbation of the Si cycle and responses of coastal ecosystems, in *Land-Ocean Nutrient Fluxes: Silica Cycle*, SCOPE 66, edited by V. Ittekkot, C. Humborg, and J. Garnier, pp. 197–213, Island Press, Washington, D. C.
- Rao, A. M. F., and R. A. Jahnke (2004), Quantifying porewater exchange across the sediment-water interface in the deep sea with in situ tracer studies, *Limnol. Oceanogr. Methods*, 2, 75–90.
- Rickert, D. (2000), Dissolution kinetics of biogenic silica in marine environments, in *Reports on Polar Research*, vol. 351, 211 pp., Alfred Wegener Inst. for Polar and Mar. Res., Bremerhaven.
- Rosenberg, D. M., P. McCully, and C. M. Pringle (2000), Global-scale environmental effects of hydrological alterations: Introduction, *Biogeochem.*, 50, 746–751, doi:10.1641/0006-3568[2000]050[0746:GSEEOH]2.0.CO;2.
- Saccone, L., D. J. Conley, G. E. Likens, S. W. Bailey, D. C. Buso, and C. E. Johnson (2008), Distribution of amorphous silica in soils of the Hubbard Brook Experimental Forest, *Soil Sci. Soc. Am. J.*, 72, 1637–1644, doi:10.2136/sssaj2007.0117.
- Schroeder, D. (1978), *Bodenkunde in Stichworten*, 154 pp., Verlag Ferdinand Hirt, Vienna.
- Seitzinger, S. P., and A. E. Giblin (1996), Estimating denitrification in North Atlantic continental shelf sediments, *Biogeochemistry*, 35, 235–259, doi:10.1007/BF02179829.
- Serebrennikova, Y. M., and K. A. Fanning (2004), Nutrients in the Southern Ocean GLOBEC region: Variations, water circulation, and cycling, *Deep Sea Res., Part II*, 51, 1981–2002, doi:10.1016/j.dsr2.2004.07.023.
- Sferratore, A., J. Garnier, G. Billen, D. Conley, and S. Pinault (2006), Silica diffuse and point sources in the Seine watershed, *Environ. Sci. Technol.*, 40, 6630–6635, doi:10.1021/es060710q.
- Simpson, T. L., and B. E. Volcani (1981), *Silicon and Siliceous Structures in Biological Systems*, Springer, New York.
- Slomp, C. P., and P. Van Cappellen (2004), Nutrient inputs to the coastal ocean through submarine groundwater discharge: Controls and potential impact, *J. Hydrol.*, 295, 64–86, doi:10.1016/j.jhydrol.2004.02.018.
- Sverdrup, H. V., M. W. Johnson, and R. H. Fleming (1942), *The Oceans*, Prentice-Hall, Englewood Cliffs, N. J.
- Tréguer, P., D. M. Nelson, A. J. Van Bennekom, D. J. DeMaster, A. Leynaert, and B. Queguiner (1995), The silica balance in the world ocean: A reestimate, *Science*, 268(5209), 375–379.
- Van Cappellen, P. (2003), Biomineralization and global biogeochemical cycles, in *Biomineralization, Rev. Mineral. Geochem.*, vol. 54, edited by P. Dove, J. DeYoreo, and S. Weiner, pp. 357–381, Mineral. Soc. Am., Washington, D. C.

- Van Cappellen, P., S. Dixit, and J. Van Beusekom (2002), Biogenic silica dissolution in the oceans: Reconciling experimental and field-based dissolution rates, *Global Biogeochem. Cycles*, 16(4), 1075, doi:10.1029/2001GB001431.
- Ver, L. M. (1998), Global kinetic models of the coupled C, N, P, and S biogeochemical cycles: Implications for global environmental change, Ph.D. diss., Univ. of Hawaii, Honolulu.
- Winkler, J. P., R. S. Cherry, and W. H. Schlesinger (1996), The Q_{10} relationship of microbial respiration in a temperate forest soil, *Soil Biol. Biochem.*, 28, 1067–1072, doi:10.1016/0038-0717(96)00076-4.
- Wollast, R. (1974), The silica problem, in *The Sea*, edited by E. D. Goldberg, pp. 359–392, Wiley.
- Wollast, R. (1991), The coastal organic carbon cycle: Fluxes, sources, and sinks, in *Ocean Margin Processes in Global Change*, edited by R. F. C. Mantoura, J. M. Martin, and R. Wollast, pp. 365–381, Wiley-Intersci., Malden, Mass.
- Wollast, R. (1993), Interactions of carbon and nitrogen cycles in the coastal zone, in *Interactions of C, N, P and S Biogeochemical Cycles and Global Change*, edited by R. Wollast, F. T. Mackenzie, and L. Chou, pp. 195–210, Springer, New York.
- Woodwell, G. M., P. H. Rich, and C. A. S. Hall (1973), Carbon in estuaries, in *Carbon and the Biosphere; Proceedings of the 24th Brookhaven Symposium in Biology*, edited by G. M. Woodwell and E. V. Pecan, pp. 221–240, Tech. Inf. Cent., U.S. At. Energy Comm., Washington, D. C.
- Yool, A., and T. Tyrrell (2003), The role of diatoms in regulating the ocean's silicon cycle, *Global Biogeochem. Cycles*, 17(4), 1103, doi:10.1029/2002GB002018.
- Yool, A., and T. Tyrrell (2005), Implications for the history of Cenozoic opal deposition from a quantitative model, *Palaeogeogr. Palaeoclimatol. Palaeoecol.*, 218, 239–255, doi:10.1016/j.palaeo.2004.12.017.
- Zhang, J. (2002), Biogeochemistry of Chinese estuarine and coastal waters: Nutrients, trace metals and biomarkers, *Reg. Environ. Change*, 3, 65–76, doi:10.1007/s10113-001-0039-3.
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