Biogeosciences, 8, 597–620, 2011 www.biogeosciences.net/8/597/2011/ doi:10.5194/bg-8-597-2011 © Author(s) 2011. CC Attribution 3.0 License.



Global spatial distribution of natural riverine silica inputs to the coastal zone

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Received: 1 October 2008 – Published in Biogeosciences Discuss.: 27 January 2009 Revised: 8 January 2011 – Accepted: 19 January 2011 – Published: 7 March 2011

Abstract. Silica, SiO₂, in dissolved (DSi) and particulate (PSi) form, is both a major product of continental weathering as well as an essential nutrient in terrestrial and aquatic systems. Here we present estimates of the spatial distribution of riverine silica fluxes under natural conditions, i.e. without human influence, to \sim 140 segments of the global coastal zone. Focussing on the construction of the DSi budget, natural DSi concentration is multiplied with discharge of rivers for each segment for documented basins and segments. Segments with no documentation available are estimated using clustered information based mainly on considerations of local lithology, climate, and lake retention. We approximate fluxes of particulate silica in various forms (PSi) from fluxes of suspended matter, calculated from existing models. Results have been established for silica fluxes, concentrations and yields for drainage basins of the different continents, oceans basins as well as coastal segment basins. For the continental surfaces actually draining into the oceans (exorheic regions, representing 114.7 million (M) km²), 371 M t y⁻¹ of DSi and 8835 M t y⁻¹ of PSi are transported, corresponding to a mean concentration of $9.5 \,\mathrm{mg} \,\mathrm{l}^{-1}$ and $226 \,\mathrm{mg} \,\mathrm{l}^{-1}$, and to a mean yield of $3.3 \, t \, km^{-2} \, y^{-1}$ and $77 \, t \, km^{-2} \, y^{-1}$, respectively. DSi yields exceeding $6.6 \, t \, km^{-2} \, y^{-1}$, i.e. $> 2 \times$ the global average, represent 17.4% of the global continental icefree exorheic area but correspond to 56.0% of DSi fluxes. Pacific catchments hold most of the hyper-active areas ($>5\times$ global average), suggesting a close connection between tec-



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tonic activity and DSi fluxes resulting from silicate weathering. The macro-filters of regional and marginal seas intercept 33% and 46% of the total dissolved and particulate silica fluxes. The mass of DSi received from rivers per unit square area of various oceans ranges over more than one order of magnitude. When expressed per unit volume and when individual regional seas are considered this figure ranges over two to three orders of magnitude, an illustration of the heterogeneity of the land to sea connection.

1 Introduction

Dissolved silica (SiO₂ or DSi, expressed as mg SiO₂ l⁻¹ throughout this paper unless noted) is one of the major nutrients of continental aquatic systems (Schelske and Stoermer, 1971) and of marine systems (Tréguer et al., 1995), essential to the development of diatoms, radiolarians and sponges (Schlesinger, 1997). Silica is also a major component of the total dissolved solids found in continental waters resulting from the chemical weathering of soils and surficial rocks. As such, it provides information on weathering processes and rates (Wollast and Mackenzie, 1983; Gaillardet et al., 1999) and has been analysed since the 19th century by European geologists and chemists. Perhaps the first comprehensive overview of silica in the environment (rock, soils, springs, rivers and the ocean) is provided by the German geologist Roth (1879, 1893). In the early 1900's, both geochemists, as the US Geological Survey pioneers (Clarke, 1924) and freshwater ecologists since the 1960's (Schelske, 1985; Wetzel, 2001) enhanced the knowledge of silica in the environment.

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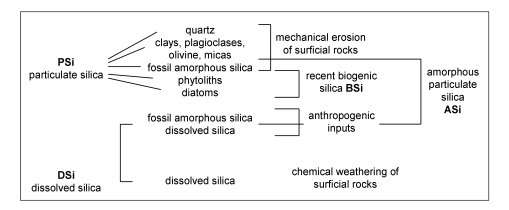


Fig. 1. Sources and forms of riverine silica.

In the last 25 years there has been a growing evidence of another major silica cycle in the terrestrial vegetation, litter and soils through the uptake and storage of silica in phytoliths (Bartoli, 1983; Meunier et al., 1999; Conley, 2002). Part of the phytoliths is not recycled and may be eroded, as the soil organic matter, and carried by rivers. Freshwater diatoms, living and as detritus, are also considered as biogenic particulate silica (BSi). It can be identified and quantified through visual analysis, a long and tedious procedure. A chemical proxy for BSi has been proposed, using a soda extraction (Paasche, 1973 modified by Nelson et al., 1989; Isaji, 2003; Saccone et al., 2006). The silica extracted by this method is referred to as amorphous silica (ASi).

In open ocean waters, ASi and BSi are nearly identical as there is little non-biogenic amorphous silica. Exceptions may temporarily occur in case significant amounts of long-range atmospheric dust, containing fly ash, are deposited on surface waters, as for East Asia downwind regions (e.g., Hartmann et al., 2008). For riverine material, ASi and BSi are not always identical as detrital amorphous silica can also originate from the mechanical erosion of sedimentary rocks and soils (e.g. chalk, cherts). Finally, there is an anthropogenic component of the silica cycle: either from diatomite earth mined from past lacustrine deposits, or artificially produced zeolith (Breese, 1994; van Dokkum et al., 2004). They are used in industries (pulp and paper production, water and waste water treatment, soil stabilization), pharmaceuticals and household products (mainly zeolith in detergents) and likely to reach the aquatic system either as dissolved or as particulate silica. A further impact of Humans on the silica cycle is the additional retention of BSi by reservoirs, mainly as diatoms detritus (Humborg et al., 2000; Straskrabova et al., 2003). This process is identical to the silica retention in natural lakes (Wetzel, 2001).

The dissolved silica fluxes are generally inferior to the particulate silica fluxes generated by the mechanical erosion of silica-containing minerals from soils and rocks (cf. Garrels and Mackenzie, 1971; Garrels et al., 1973; Martin and

Meybeck, 1979). River inorganic silicon-containing particulates of non-biogenic origin are composed mostly by quartz, feldspar or clay minerals (illite, chlorite, kaolinite, etc.) (e.g., Subramanian, 1980; Gölz, 1982; Gaiero et al., 2004). Minerals most resistant to chemical weathering are characterised by increased proportions in suspended matter, if compared to unweathered source rocks of a river catchment. Sometimes fine fragments of eroded rocks may contain amorphous detrital silica. All silica-containing components, including phytoliths and BSi, are part of the particulate silica (PSi), analysed after a complete digestion of river particulates.

There are therefore multiple ways to consider the riverine silica, from the analytical chemist, the ecologist, the geochemist, and the physical geographer (Fig. 1).

Since the first estimates of dissolved silica concentrations in rivers by Roth (1879) and Clarke (1924), there have been few studies on this topic, compared to the ones on nitrogen and phosphorus (Livingstone, 1963; Meybeck, 1979, 1987, 1988; Probst, 1992; Tréguer et al., 1995 and the special issue of Global Biogeochemical Cycles, edited by Seitzinger et al., 2005). Previous DSi budgets are not or little spatialized (e.g., Jennerjahn et al., 2006) while other nutrients and carbon budgets have been gradually spatialized over the last 15 years, first at the $2^{\circ} \times 2^{\circ}$ resolution, then at $0.5^{\circ} \times 0.5^{\circ}$ (Ludwig et al., 1996; Amiotte-Suchet et al., 2003; Green et al., 2004; Seitzinger et al., 2005). The first model-based spatially explicit DSi budget (Beusen et al., 2009) is based on the data presented here. Many of these recent spatialized nutrient budgets are based on a general register of river inputs to oceans, GEMS-GLORI (Meybeck and Ragu, 1995) that lists about 500 individual rivers with various levels of documentation.

In the case of spatially explicit model budgets for dissolved and particulate matter, in general monitoring data of the \sim 60–120 largest rivers draining to coastal waters are applied for model calibration and small catchments information is neglected. However, some studies provide evidence that regions characterised by small catchments draining to coastal

zones in humid and warm climates (with a specific lithologic characteristic) represent above average specific fluxes (Milliman and Syvitski, 1992; Hartmann et al., 2009, 2010a). In particular, the Ring of Fire, surrounding the Pacific Ocean, is characterised by small and steep catchments due to plate tectonics.

The only existing detailed global river data base on dissolved and particulate matter (Meybeck and Ragu, 1995) is also partly used here for the emphasized dissolved silica budget in order to

- i. reach a global spatial distribution of natural i.e. without (or low) human influence river dissolved silica concentrations and fluxes at the regional level (~ one million square kilometres);
- ii. compare these with the particulate silica fluxes;
- iii. identify areas most contributing to the global budget;
- iv. discuss the net silica inputs to oceans, focusing on DSi.

We therefore consider the following steps:

- collection and selection of individual DSi and PSi data for major and some minor rivers at the global scale, with a clear focus on constructing the DSi budget based on the earliest data available in order to avoid human overprint;
- extrapolation of documented data at regional resolution using a previously introduced coastal catchments delineation (COSCATs) (Meybeck et al., 2006), and calculation of silica river fluxes for each coastal catchment;
- iii. spatial analysis of silica fluxes from continents, for open oceans and regional seas, according to the methodology defined by Meybeck et al. (2007) and Meybeck and Dürr (2009).

The data compiled and the results presented here, have been used in several applications: Bernard et al. (2011) used the spatialized data as input for an OGCM. Beusen et al. (2009) used the data for a statistical approach on control factors of global DSi fluxes. Laruelle et al. (2009) used the data for an assessment of the global biogeochemical silicon cycle. Also regional studies may benefit as they need spatialized data for comparison with other, similar regions (e.g., Hartmann et al., 2009; Jansen et al., 2010). We also identify here major regional seas as significant filters of land-ocean DSi fluxes. The budget of DSi reaching the open ocean is significantly different from the amount that is delivered at river mouths to the coastal zones.

2 Data and methodology

2.1 Database for DSi and PSi in river waters and particulate matter

The dissolved silica found in continental waters originates from the chemical weathering of a great number of minerals, from the more easily weathered (e.g., volcanic glass, olivine) to the least (e.g. clays, quartz) (cf. Wollast and Mackenzie, 1983; Meybeck, 1987; Stallard, 1995). The weatherability of minerals generally has a stronger control on DSi mobilization from rocks than the silica content of the various, distinguished rock types or lithological classes (cf. Hartmann et al., 2010a, 2011). In general, the levels of silica concentrations transported from the terrestrial to the fluvial system depend on the relative proportions of minerals in soils and rocks and their weatherability, on temperature, water residence time in soils and rocks, drainage intensity, gradient of slope, land use/cover changes and acids involved in weathering processes, as e.g., carbonic, sulphuric or humic acids (Meybeck, 1979; Wollast and Mackenzie, 1983; Drever, 1994, 1997; Gaillardet et al., 1999; West et al., 2005; Jennerjahn et al., 2006; Conley et al., 2008; Hartmann et al., 2010a; Struyf et al., 2010; Cornelis et al., 2011).

The yield of dissolved silica exported by rivers per unit area ($t \operatorname{SiO}_2 \operatorname{km}^{-2} \operatorname{y}^{-1}$) is first dependant on the river runoff (mm y^{-1}), then on the control factors of concentration (Wollast and Mackenzie, 1983; Meybeck, 1994; Hartmann et al., 2010a and b; Cornelis et al., 2011).

The retention of silica in river systems, e.g. in lakes and flood plains, is linked to their trophic state (for DSi), to lake abundance, expressed by the limnic index (% of lake cover in a given area, i.e. % of area covered by lakes with respect to the total catchment basin area – this value can attain up to 20% at the $\sim 1~\rm M~km^2$ scale regarded here) (Conley et al., 2000), and by deposition of ASi in alluvial plains (Admiraal et al., 1990). The commonly reported retention for DSi extends from 4 to more than $100~\rm g~SiO_2$, with a median value of $20\pm10~\rm g~SiO_2$, per year per m² of lake area (Campy and Meybeck, 1995). Apart from the formerly glaciated shields in Canada, Northern Europe and Taymir Peninsula (Siberia), where it commonly ranges between 5 and 20%, the limnic index rarely exceeds few percent in other parts of the world (Meybeck, 1995).

In order to construct the DSi database, dissolved silica concentrations are first selected from the Meybeck and Ragu (1995) catalogue of river inputs to oceans ("GEMS-GLORI", www.gemswater.org). They selected 1 to 3 average analyses for the same river, depending on authors and period of records, in which we generally took the earliest silica average. We also used another data set on pristine rivers of the world (PRISRI, see the Supplement), already used by Meybeck (2003), to determine the control factors of river chemistry in natural conditions. In this second data set numerous important references on river chemistry prior 1950/1960

are given and reported here in our Supplement, i.e. before the main development of large reservoirs on world rivers (Vörösmarty et al., 1997), as (i) works on US rivers from a dozen of US Geological Survey reports prior to 1940, most of them used and cited by Clarke (1924), (ii) former Soviet rivers analysed between 1940 and 1960 by Alekin and Brazhnikova (1960, 1962 and 1968) and (iii) June Kobayashi's work on Asian rivers in the same period for Japan, Thailand, Indonesia, Sri Lanka (Kobayashi, 1959, 1960 and 1966). We also selected more recent analyses in regions with limited human impacts as in Alaska and Canada (Brunskill, 1986), Amazon and Orinoco basins (Lewis and Saunders, 1989), Patagonia (Depetris and Paolini, 1991), in West and South Africa (de Villiers, 1962; Olivry and Noah, 1978; Orange, 1992). Most of the presented analyses thus have a date before 1980/1990. Some carefully selected personnel communications were added where we could not track the original data source. The detailed list of references is given in Supplement 1A to F for each river.

Measuring dissolved silica is a relatively simple and standard colorimetric method, used since tens of years (e.g., Robinson and Thompson, 1948; Strickland et al., 1968; Koroleff, 1983). Contrary to other substances (as ammonium) potential error sources are small, and we believe that using relatively old data does not present a particular bias as to measuring methods. Furthermore, the data used and the according sources were included in a substantial data quality control and standardization efforts, based on principles used in previous studies (Meybeck and Ragu, 1995; Meybeck, 2003). For example, sufficient inter- and intra-annual data must be present to assure representative annual averages. Meybeck and Ragu (1995) have explained the detailed data quality requirements for the Gems-Glori database, which has been designed using a similar approach. The pristine river database used here follows the same data quality requirements (Meybeck, 2003), with the additional requirement that as little human influence as possible can be detected, and hence often the oldest available data have been recognized as more appropriate to avoid as far as possible human influences due to the construction of dams or eutrophication. As the documentation of all procedures for the data sampling, measurement, etc. is beyond the appropriate space and scope of this paper, we refer to the references provided.

Particulate silica (PSi) in rivers is not analysed within most water quality surveys as it is not listed as an indicator of contamination: it must be collected for each river from individual papers dealing with suspended particulate matter (SPM) chemistry. The total particulate silica obtained after complete digestion of filters or of dried sediments is the most common element found in any SPM analysis although it is not much analysed in academic literature, instead heavy metals and rare earth elements (REE) are the most commonly analysed. In case of large proportions of carbonate rocks in the river catchment, most of the carbonate minerals are dissolved during chemical weathering and only the insoluble material

– e.g., clays – is carried with SPM together with some fine carbonate detritus (e.g., Gölz, 1982). In the example case of France, the average SiO₂ concentration (DSi) in limestone and dolomite draining streams is 3.6 and 4.0 mg l $^{-1}$, respectively, and can go up to $12.0\,\mathrm{mg}\,\mathrm{l}^{-1}$ for chalk basins due to the presence of amorphous silica in this rock type; in gypsum marl DSi can also be high, between 8.0 and 9.6 mg l $^{-1}$ (Meybeck, 1986, 2003). As a result, even carbonated river basins do export particulate silica. The coarse fraction of river particulates that transits as bed load consists of quartz with few percent of heavy minerals.

As a result, in medium to large rivers silica is largely dominating all other elements in particulates (Martin and Meybeck, 1979; Canfield, 1997). Several world averages have been proposed for silica since 1979, they are very similar despite the growing data base: Gordeev and Lisitzin (1978) 255 000 ppm, Martin and Meybeck (1979) 285 000 ppm, Glazovsky (1990) 254 000 ppm. Viers et al. (2009) compiled a new global database on the chemical composition of suspended sediment fluxes, they found a quite similar value for Si content (254 000 ppm, with a variation of 117 000 ppm). A preliminary survey of hundreds of river analyses for major and trace elements (GLOMET, see the Supplement), based on data from 62 major basins and tributaries from 5 continents and from latitudes ranging from 40° S to 65° N (containing to a certain amount the same data used in the existing literature), gives a PSi median content of 260 000 mg Si per kg (or ppm) of river particulates. The major control factor of PSi export by unit river basin area (yield) is therefore the sediment supply that commonly ranges from 10 to $10\,000\,\mathrm{t\,km^{-2}\,y^{-1}}$ in medium and large rivers (Milliman and Syvitski, 1992). Among more than 50 chemical elements analysed in river particulates, Si is the least variable, either under different climate conditions or due to rock lithology, for rivers exceeding 10 000 km². Silicium, or silica, is also not largely affected by human impacts in the GLOMET database. The only exception is noted for rivers with very low SPM concentrations due to reservoir settling of detrital material and to the presence of biogenic silica due to diatom growth. The related fluxes at the global scale are quite limited. Therefore, we have considered here as a first approximation a constant PSi content of 260 000 mg Si kg⁻¹, i.e. 55.64 weight % SiO₂, for each coastal catchment, i.e. at the one million km² resolution at which this budget is realised. In this first estimate the river bed load, which consists mostly of quartz sands and gravel, has not been considered. It is generally higher in steeper headwaters and where mountainous rivers drain quickly to oceans, but this kind of data is seldom reported and is thought not to account significantly in the global budget, often considered as 10% of the river total sediment transport (cf., Zanke, 1982; Milliman and Syvitski, 1992).

References for the data base on silicon analysis in river SPM and/or freshly deposited fine-grained sediments can be found in the Supplement.

2.2 Calculating silica budgets for coastal catchments

For spatial analysis of material fluxes the exorheic part of the continents can be segmented, as done by Meybeck et al. (2006), resulting in a mega-puzzle of \sim 140 pieces, the coastal catchments or COSCATs. These entities have been specifically designed to allow the reporting of global river fluxes of water, of dissolved and particulate materials as nutrients and carbon species, and of contaminants in a standardized way. Previous coastal segmentations are not applicable for river budget reporting, while COSCATs delineations are based on multiple criteria combining basin boundaries, coastal morphology, ocean dynamics, regional seas limits etc. (see a complete discussion in Meybeck et al., 2006). The COSCATs are related to the exorheic part of the ~6200 individual river basins that have been delineated at the $0.5^{\circ} \times 0.5^{\circ}$ resolution and can be fully described through many databases that have been developed over the last 10 years, in particular within the IGBP-BAHC programme (Vörösmarty and Meybeck, 2004) as the global river network (Vörösmarty et al., 2000a, b), the global scale relief typology (Meybeck et al., 2001), or lithology (Dürr et al., 2005). Of particular importance is the water runoff field at the same resolution, resulting from a global-scale model combined to data from the Global Runoff Data Center at WHO/UNESCO (Fekete et al., 2002).

As such, each segment has a maximum homogeneity in its coastal area. Fixed segment boundaries allow easy description and mapping at coarse resolution and all land to ocean fluxes can be reported in the same format, e.g. for application in Earth System models. The median length of coastline of the segments (at the $0.5^{\circ} \times 0.5^{\circ}$ resolution) is a little over 2000 km, and the mean reach of the segments inland is 360 km (median 150 km). Some of the COSCATs are quite large in area, from 2 to 7 millions (M) km², since the largest river basins cannot be split as for the Amazon or Congo. The median size of the COSCATs is 0.83 M km². Each COSCAT is associated to a river runoff (mm y^{-1} or km³ y^{-1}) that can range from near zero (arheism) to $>2400 \,\mathrm{mm}\,\mathrm{y}^{-1}$ for very humid tropics (Meybeck et al., 2006). COSCATs have a 4 digit coding: the first two digits for individual continents, then for individual COSCATs numbered clockwise. COSCAT names are originating from oceanic and coastal morphological features (see original maps and details in Meybeck et al. (2006) and the auxiliary material pertaining to it). Note that COSCAT boundaries have also been specifically designed to allow for Regional Seas budgets (Meybeck and Dürr, 2009).

For each COSCAT the DSi fluxes result from the following steps:

 Product of documented DSi concentrations by the related natural river discharges (usually also available in Meybeck and Ragu, 1995; otherwise they are calculated from the Fekete et al. (2002) data set), and calculation of non-documented area and runoff.

- ii. If the river basin characteristics in the undocumented area are close to the documented ones then these known average concentrations are extrapolated to the remaining part of the COSCAT.
- iii. If the undocumented area is much different, a specific river or tributary can be chosen as representative of this area. It may even originate from a nearby COSCAT or river where general basin characteristics are similar. This informational clustering is based on lithology, climate, relief, water drainage, and abundance of lakes, which are the main control factors of silica fluxes (Wollast and Mackenzie, 1983; Meybeck, 1995, 2003; Drever, 1997; Hartmann et al., 2010a). This estimated DSi is then combined to the remaining runoff which can also be different from the one of the documented area (estimated DSi levels are given in Supplement 1 for each COSCAT). For instance, for very large COSCATs where climate and runoff can be heterogeneous, as for the Ob, Yenisseï and Lena, the undocumented coastal basins have not been based on the known DSi in these major rivers, but on much smaller north Siberian rivers that are documented as Khatanga and Olenek, coupled with a much lower runoff.
- iv. Few COSCATs remain completely undocumented: here the DSi concentration is chosen from rivers or a group of rivers characterised by similar weathering conditions, including similarity in catchment properties, and attributed to the whole runoff of the undocumented COSCAT.

The resulting discharge-weighted silica concentrations DSi* (mg SiO₂ l⁻¹) and yields Y_{DSi} (t SiO₂ km⁻² y⁻¹) are then calculated for each COSCAT (Eqs. 1 and 2):

$$DSi_{j}^{*} = (\Sigma \text{ fluxes}_{cij})/(COSCAT_{j} \text{ discharge})$$
 (1)

$$Y_{\text{DSi}j} = (\Sigma \text{ fluxes}_{\text{cij}})/(\text{COSCAT}_j \text{ area})$$
 (2)

With "ci" being the running index for catchments within a COSCAT_i.

The fluxes and yields of particulate silica for each COSCAT have been calculated on the basis of SPM loads estimated from the Ludwig and Probst (1998) model which takes into account several control factors as relief, lithology, runoff, vegetation cover and is used to characterise the particulate inputs for each COSCAT. This spatialized model $(0.5^{\circ} \times 0.5^{\circ})$ does not include the sediment trapping in reservoirs which is believed to become more and more important (Vörösmarty et al., 2003; Syvitski et al., 2005), it is therefore compatible with our target of estimating the silica budget as close as possible to pre-anthropogenic conditions. SPM yields and loads are then combined to the median PSi content $(260\,000\,\mathrm{mg}\,\mathrm{Si}\,\mathrm{kg}^{-1})$ to generate the corresponding PSi yields and loads re-converted into SiO₂ (Eq. 3) in order to

Table 1. Average dissolved and particulate silica fluxes, concentrations (runoff weighted averages) and yields (area weighted averages) for the exorheic parts of the continents. Catchment surface area, discharge and runoff are given as well as % of volcanic rocks from Dürr et al. (2005). The fraction of PSi that possibly dissolves and becomes biologically available might be <1% (see discussion).

	Catchment area 10 ⁶ km ²	Discharge km ³ y ⁻¹	$\begin{array}{c} \text{Runoff} \\ \text{mm y}^{-1} \end{array}$	Volcanic rocks (Vb+Va+1/2 Cl) ^b %	DSi mean concentration $mg l^{-1} SiO_2$	DSi flux $M t y^{-1}$ SiO_2	DSi yield $t \mathrm{km}^{-2} \mathrm{y}^{-1}$ SiO_2	PSi flux $M t y^{-1}$ SiO_2	PSi yield $t \mathrm{km}^{-2} \mathrm{y}^{-1}$ SiO_2
Africa	26.2	4130	157	4.2	12.6	52.1	2.0	528	20.1
Europe ^a	8.4	2410	285	6.9	5.6	13.5	1.6	367	43.5
North America	21.9	5880	268	11.3	8.0	47.2	2.2	1713	78.2
South America	17.4	11 850	681	10.0	9.0	106.5	6.1	1633	93.9
Asia	34.0	12900	379	12.8	10.0	129.4	3.8	3951	116.2
Australasia	6.7	1920	284	8.4	11.8	22.6	3.3	643	95.3
Total Exorheic	114.7	39080	341	9.3	9.5	371	3.3	8835	77.0

^a Including a small non-glaciated fraction of Greenland.

facilitate the comparison between Y_{DSi} and Y_{PSi} , both expressed as SiO₂:

$$Y_{\text{PSi}\,i} = Y_{\text{SPM}\,i} \times 0.26 \times 2.14$$
 (3)

With

 $Y_{\text{SPM}j}$ = sediment yield in COSCAT_j in t SPM km⁻² y⁻¹ 0.26 = constant factor for all COSCATs: 26% of Si in SPM or 260 000 ppm Si

2.14 = conversion factor from Si to SiO₂.

Example:

$$Y_{\text{SPM}x} = 100 \text{ t km}^{-2} \text{ y}^{-1} \rightarrow Y_{\text{PSi}x} = 55.6 \text{ t SiO}_2 \text{ km}^{-2} \text{ y}^{-1}$$

3 Results

The combination of COSCATs and their silica budgets allows for the calculation of various types of budgets: per continent, per ocean drainage basin, per latitudinal zones, and for different regional seas. It is common among geochemists and geographers to calculate riverine fluxes per continents (Berner and Berner, 1996; Gerasimov et al., 1964): this type of reporting is used in Table 1. In this table the exorheic catchment area of continents – i.e. the area facing the oceans – includes areas that are not presently characterized by active river runoff – i.e. arheic regions. This operational cut-off is set here at 3 mm y⁻¹ (Vörösmarty et al., 2000a, b) which corresponds to an average of one flood occurrence every 10 years in desert regions. Presently glaciated areas as Antarctica and major parts of Greenland are not considered in our budget.

Endorheic regions – i.e. facing internal regions – as the Caspian Sea basin (Volga, Terek, Sulak, Kura Rivers), the Rift Valley (Omo River), Lake Chad (Chari and Logone Rivers) and Okawango (Okawango River) basins in Africa, the Lake Eyre Basin (Cooper River) in Australia, the Altiplano (Deseaguadero River) and the Great Basin (Sevier and Humboldt Rivers) in the Americas, the Caspian and Aral Sea

basins and other regions of Central Asia, are not considered in our budget although they contribute to the weathering of continents at the global scale. The few major rivers with available data (Meybeck, unpublished report to WHO, Div. Envir. Health, Geneva, "Water quality of endorheic basins", 1996: Chari $19.2 \text{ mg SiO}_2 \, l^{-1}$ for $24.2 \, \text{km}^3 \, \text{y}^{-1}$ discharge; Logone $19.3 \, \text{mg SiO}_2 \, l^{-1}$ for $15.8 \, \text{km}^3 \, \text{y}^{-1}$, Okawango $16.0 \,\mathrm{mg} \,\mathrm{SiO}_2 \,\mathrm{l}^{-1}$ for $10.0 \,\mathrm{km}^3 \,\mathrm{y}^{-1}$; Omo $18.0 \,\mathrm{mg} \,\mathrm{SiO}_2 \,\mathrm{l}^{-1}$ for $14.0 \text{ km}^3 \text{ y}^{-1}$; Sevier $38.4 \text{ mg SiO}_2 \text{ l}^{-1}$ for $0.02 \text{ km}^3 \text{ y}^{-1}$; Humboldt $32.0 \,\mathrm{mg} \,\mathrm{SiO}_2 \,\mathrm{l}^{-1}$ for $0.42 \,\mathrm{km}^3 \,\mathrm{y}^{-1}$ discharge) have a discharge-weighted DSi average of 18.6 mg SiO₂ l⁻¹ (1.2 Mt total annual DSi flux for 64.5 km³ of annual discharge, i.e. 7.1% of the total annual endorheic discharge for 6.1% of the total endorheic catchment area). Most of these regions are characterised by semi-arid to arid conditions (63% of the total endorheic area are arheic) with elevated DSi, compared to the global average of the exorheic regions.

The exorheic parts of individual continents are characterized as follows:

Although North America has an average runoff of 268 mm y^{-1} , much higher than Africa (157 mm y^{-1}) , their silica yields, in $t \text{SiO}_2 \text{ km}^{-2} \text{ y}^{-1}$, are of the same order: $2.2 \text{ tkm}^{-2} \text{ y}^{-1}$ for North America, $2.0 \text{ tkm}^{-2} \text{ y}^{-1}$ for Africa. This is in part due to the silica retention by lakes observed in most of the Canadian rivers where the limnic index of lake occurrence is commonly between 5 to 20% (Meybeck, 1995; Lehner and Döll, 2004): most Canadian rivers have a DSi level between 1.1 and $3.6 \text{ mg SiO}_2 \text{ l}^{-1}$ (see Supplement 1C, COSCATs #0815 to 0818) while Alaska and British Columbia rivers are characterised by much higher silica levels (COSCATs #0809, 0810, 0812, 0813) and much lower limnic index (Table 6).

European rivers have the lowest silica levels on average: $5.6 \text{ mg SiO}_2 \text{ l}^{-1} \text{ vs. } 9.5 \text{ mg SiO}_2 \text{ l}^{-1} \text{ for the global exorheic}$ average. This is probably due to several factors: cold to

b Vb - Basic volcanic rocks, Va - Acid volcanic rocks, Cl - Complex lithology (consisting 1/2 of volcanic rocks, see Dürr et al., 2005).

Table 2. Average dissolved and particulate silica fluxes, concentrations (runoff weighted averages) and yields (area weighted averages) from catchments to coastal zones related to ocean basins. Catchment surface area, discharge and runoff are given as well as % of volcanic rocks from Dürr et al. (2005).

	Catchment area 10 ⁶ km ²	Discharge km ³ y ⁻¹	Runoff mm y ⁻¹	Volcanic rocks (Vb+Va+1/2 Cl)* %	DSi mean concentration $mg l^{-1} SiO_2$	DSi flux Mty ⁻¹ SiO ₂	DSi yield t km ⁻² y ⁻¹ SiO ₂	$\begin{array}{c} \text{PSi flux} \\ \text{Mty}^{-1} \\ \text{SiO}_2 \end{array}$	PSi yield tkm ⁻² y ⁻¹ SiO ₂
Arctic	16.7	3310	199	8.3	6.1	20	1.2	122	7.3
Atlantic	57.5	19 950	347	5.4	8.4	169	2.9	2748	47.8
Indian	20.8	5200	250	7.9	12.0	62	3.0	2138	102.8
Pacific	19.8	10 620	1010	22.9	23.2	120	6.1	3826	193.5
Total Exorheic	114.7	39 080	341	9.3	9.5	371	3.3	8835	77.0

^{*} Vb – Basic volcanic rocks, Va – Acid volcanic rocks, Cl – Complex lithology (consisting 1/2 of volcanic rocks, see Dürr et al., 2005).

temperate climate, retention of silica in Scandinavian lakes (Conley et al., 2000), scarcity of (young) volcanic rocks and abundance of carbonate rocks (Dürr et al., 2005). Removal of DSi by plankton communities by elevated nutrient (N and P) levels may impact observed low concentrations, too. Despite our attempt to choose reported DSi concentrations with low anthropogenic impact in general, for Europe it remains difficult to cope with this objective, because of the anthropogenic influence since the 19th century. E.g., Roth (1879) reports nitrate concentration for the Rhine (1848), Seine (1848) and Thames (no date) of 3.8, 14.6 and 3.9 mg l⁻¹, respectively.

Africa silica concentrations are the highest (12.6 mg I^{-1} on average), despite the scarcity of volcanic rocks. This can be explained by the prevalence of warm climate conditions that favour chemical weathering of crystalline rocks abundant on this continent, for the exorheic regions. In addition, an evaporation effect is likely to influence concentrations (cf. White and Blum, 1995).

South America silica yields are near twice the world average (6.1 vs. $3.3 \, \text{t} \, \text{km}^{-2} \, \text{y}^{-1}$), with silica concentrations very close to the world average (9.0 vs. $9.5 \, \text{mg} \, \text{l}^{-1}$). This is related to the very high runoff on this continent, $681 \, \text{mm} \, \text{y}^{-1}$ vs. $340 \, \text{mm} \, \text{y}^{-1}$ for the global average: the silica yield of river systems is predominantly controlled by their runoff (Meybeck, 1994). Most of the observed DSi fluxes originate in the humid regions of the Andes (cf. data from different regions of South America in Mortatti and Probst, 2003; Depetris et al., 2005; Jennerjahn et al., 2006), characterised by high proportions of volcanics (Dürr et al., 2005).

DSi fluxes from Asia are not easy to evaluate, because of the great heterogeneity of the continent. The very wet southern flanks of the Himalayan region are characterised by high fluxes due to intensive influence of the monsoon (Kobayashi, 1959), whereas much drier central parts like Tibet are characterised by low fluxes (Galy and France-Lanord, 1999). Large endorheic areas are located in the centre of Asia and cold climate in the northern parts leads to low DSi fluxes.

The silica budgets for ocean drainage basins (Table 2) generate the following comments:

Due to its extended drainage area and water inputs (50% of world exorheic), the Atlantic ocean receives 45% of the silica inputs, with an average concentration (8.4 mg l $^{-1}$) very close to the world average (9.5 mg l $^{-1}$), despite a lower than average outcrop proportion of volcanic rocks (5.4% vs. 9.3% on average). The presence of the three world's major rivers in terms of discharge, Amazon (world's first), Congo (second) and Orinoco (third), which are characterised by high runoff values and higher dissolved silica contents (45.5 M t, 12.4 M t and 7.2 M t SiO $_2$ flux per year, carrying alone 18% of the global exorheic DSi flux), compensates the limited inputs from the dry regions of the Atlantic Ocean basin.

The relatively high dissolved silica concentrations in the Pacific ocean drainage basin rivers $(23.2 \, \text{mg SiO}_2 \, l^{-1})$ on average) are probably linked to its higher volcanic outcrops (22.9%) and are thus related to the tectonic activity of the Ring of Fire, surrounding the Pacific. Silica yields in this basin are also the highest, due to a runoff three times the world's average $(1010 \, \text{vs.} \, 340 \, \text{mm y}^{-1})$, but its drainage area is relatively small. As a result the absolute DSi flux to the Pacific Ocean is less than the one from the Atlantic drainage basin.

Arctic rivers are characterised by relatively low silica levels, $6.1\,\mathrm{mg\,SiO_2\,l^{-1}}$ on average, for an average volcanic rock outcrop. Multiple factors can be invoked: possible lower weathering rates due to colder climate, the extent of permafrost, and the areal proportion of lakes in Canada and Scandinavia, which is an order of magnitude higher than in the other ocean drainage basins (Meybeck, 1995; Lehner and Döll, 2004), resulting from the occurrence of past ice caps in the Arctic drainage basin. The low DSi yield $(1.2\,\mathrm{t\,km^{-2}\,y^{-1}}$ vs. the world's average of 3.3) is linked also to a lower runoff (199 mm y⁻¹). The absolute flux of DSi received by this ocean is only 5.4% of the world's river fluxes $(371\,\mathrm{M\,t\,SiO_2\,y^{-1}})$.

It is important to note that, when the DSi inputs are compared to ocean volumes and/or to ocean area, these relative inputs are ranked in a very different manner (Table 3). Ocean volumes and area are those given by Kossinna (1921) and reported by Viglieri (1966) in Fairbridge (1966). Ocean

Table 3. Dissolved and particulate silica fluxes received by coastal zones related to ocean basins, rated to coastline length, ocean areas and ocean volumes. Retention of silica in estuaries, on shelves and in regional seas is not considered here. Coastline length, ocean area and ocean volume according to Meybeck et al. (2006, 2007).

	Arctic	Atlantic	Indian	Pacific	Global Ocean
t DSi silica received per km y ⁻¹ of coastline	353	1133	920	935	923
t per $km^2 y^{-1}$ of ocean area	1.64	1.79	0.84	0.66	1.03
$t \text{ per km}^3 \text{ y}^{-1} \text{ of ocean volume}$	1.47	0.50	0.22	0.17	0.28
t PSi received per km y ⁻¹ of coastline	2144	18 541	31 613	29 706	21 992
t per km ² y ⁻¹ of ocean area	9.97	29.3	28.9	21.1	24.4
t per km ³ y ⁻¹ of ocean volume	8.92	8.18	7.51	5.36	6.55

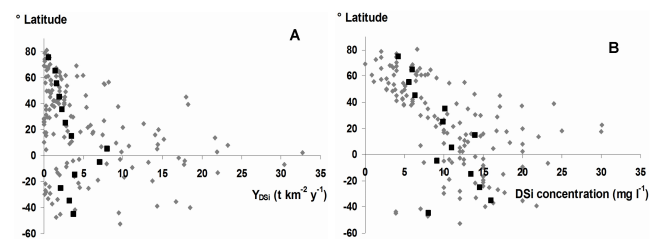


Fig. 2. Latitudinal distribution of dissolved silica inputs to coastal zones (at barycentre of the 140 COSCATs). (**A**): Y_{DSi} in t SiO₂ km⁻² y⁻¹, (**B**): DSi concentration in mg SiO₂ l⁻¹. Grey dots: individual COSCATs, black dots: averages per 10° latitudinal bands.

coastline has been determined here for each COSCAT at the $0.5^{\circ} \times 0.5^{\circ}$ resolution (Meybeck et al., 2006) and may be somewhat lower than the actual coastline delineated at a finer resolution: it varies between 57 000 km for the Arctic Ocean and 149 000 km for the Atlantic Ocean (Meybeck et al., 2007). The Arctic Ocean is actually receiving the maximum DSi inputs with regards to its size and the relative silica inputs to the Pacific Ocean are more than one order of magnitude less.

The global estimate for river SPM fluxes to coastal zones is 16 billion tons per year (Ludwig and Probst, 1998), corresponding to a mean concentration of $226\,\mathrm{mg}\,\mathrm{l}^{-1}$ or about 9 billion tons of particulate Si, expressed as SiO_2 , i.e. the DSi/PSi ratio at the global scale is 4.2% (371/8835 = 0.042). As the PSi concentration in SPM is nearly constant (26% \pm 4% of Si), the distribution of PSi concentration per liter of water ranges over three orders of magnitude in rivers since it is similar to the one of SPM which commonly varies between less than $10\,\mathrm{mg}\,\mathrm{l}^{-1}$ to more than $10\,000\,\mathrm{mg}\,\mathrm{l}^{-1}$ in world rivers. We refer here to the large body of literature describing the characteristics of SPM yields and concentra-

tions in world rivers (e.g., Milliman and Syvitski, 1992; Ludwig and Probst, 1998; Meybeck et al., 2003; Syvitski et al., 2005 and references therein).

The latitudinal distribution of the measured and/or estimated DSi from the ~140 coastal segments is reported on Fig. 2, using the latitude of the COSCAT barycentre (or gravity centre). Maximum concentrations are found in the subtropical regions while the boreal regions have much lower concentrations: minimum concentrations of the sub-tropics (around $10 \text{ mg SiO}_2 l^{-1}$) are higher than maximum concentrations north of 50° N and south of 45° S. Due to the runoff minimum for the 20-30° N and 20-30° S belts, the DSi export rate is relatively low at these latitudes despite medium DSi concentrations. The maximum DSi is actually linked to the wet sub-tropics where both high DSi concentrations and runoff are encountered. With few exceptions, regions with high DSi yields correspond to high runoff values as well as high DSi concentrations, but this is a very general picture for all latitudes. For example, most COSCATs with DSi yields $\ge 2 \times$ the world exorheic average, i.e. $\ge 6.6 \,\mathrm{t\,km^{-2}\,y^{-1}}$, also have runoff values $\ge 2 \times$ the world exorheic average, i.e.

Table 4. Percentages of documented area, discharge and corresponding dissolved SiO₂ fluxes for continents, regional seas and ocean drainage basins.

	% Area	% Discharge	% DSi flux
Africa	46.3	46.9	43.5
Europe*	53.0	47.0	44.3
North America	60.2	51.6	49.0
South America	69.4	78.3	70.1
Asia	63.1	54.6	44.7
Australasia	25.8	25.2	25.0
Regional Seas	53.6	48.5	42.2
Arctic	77.3	81.6	76.8
Atlantic	59.4	68.5	66.8
Indian	36.5	41.0	35.2
Pacific	53.0	41.4	33.1
Total Exorheic	56.7	58.6	51.1

^{*} Including a small non-glaciated fraction of Greenland.

 $>680 \,\mathrm{mm}\,\mathrm{y}^{-1}$ (see discussion). The relative minimum in silica concentrations around the equator might be due to the often deeply weathered environments in the Brazilian and African shields located in the central tropics (cf. Boeglin and Probst, 1998). For example, the two largest tropical rivers, i.e. the Amazon and Congo, both show relatively low concentrations at their river mouth. They are characterised by large proportions of deeply weathered regolith in areas contributing large proportions of runoff and, in the case of the Congo river, also some proportions of carbonates are present (Probst et al., 1994; Stallard, 1995; Mortatti and Probst, 2003; Dürr et al., 2005). Low concentrations can further be attributed to tectonic stability (>100 My) and to the high precipitation in these wet tropical regions (cf. Stallard, 1995; Mortatti and Probst, 2003). The northern Hemisphere is characterised by a strong latitudinal correlation of DSi concentrations and yields towards the equator and is probably indicating the influence of temperature and vegetation, pointed out by missing values in the lower concentration range in the subtropical and tropical zone.

4 Discussion

4.1 Global coverages and approaches to river silica budgets

For each coastal catchment we have determined the sum of documented silica fluxes vs. undocumented ones and their related areas and runoff. These area and discharge coverage rates are tabulated for individual continents and ocean drainage basins on Table 4, and in the Supplement for individual COSCATs. The resulting percentages of dissolved silica flux coverage range from 25% for Australasia – i.e. Aus-

tralia, New Guinea and New Zealand - to 81% for the Arctic river runoff. The mean coverage rates are 56.7% for the continental area, 58.6% for the continental runoff, and 51% for the silica flux, i.e. half of the silica flux is based on actual water analyses combined with measured runoff, half has been extrapolated. This extrapolation has been realized by the product of estimated regional or local DSi concentrations and the river runoff over undocumented areas, now known from global models with a satisfactory accuracy (Fekete et al., 2002). If we assume that DSi analytical uncertainties and flux calculation uncertainties for documented rivers are negligible, the uncertainty induced by the frequency of regular river measuring campaigns results in errors around 10% for major ions as expressed by the electrical conductivity (Moatar and Meybeck, 2007), and is likely to be identical for DSi. Based on extrapolation studies for dissolved silica fluxes (Hartmann et al., 2010a; Jansen et al., 2010; Moosdorf et al., 2010), it can be assumed that the performance of the budget approach is weaker. The maximum uncertainty will thus be somewhat higher: sometimes not many analyses are used for a certain value or no measurements over several seasons exist, or only arithmetic averages are available instead of discharge-weighted averages. While it is difficult to put actual numbers on the overall uncertainties, considering available data and that the budget approach was designed to overcome gaps of data for certain local and regional areas, some aspects are better known, such as the uncertainty induced by extrapolating annual averages of concentrations (cf. Moatar and Meybeck, 2007). As a rough approximation, we estimate that the additional uncertainty on extrapolated yields, using our method, yields in a total error probably not much greater than 30% for the regional scale. Locally, and in less well documented areas, the uncertainty is of course higher. However, as the budget approach closes gaps in knowledge of DSi-concentrations from certain areas, a reliable and calculable number can not be provided for these areas.

The silica content of river SPM also depends on the grain size: coarser bed sediments are more siliceous with contents around 300 000 ppm while SPM is mostly in the silt and clay fraction and has a lesser quartz content and a higher content in Al, and therefore a lower content of Si. Viers et al. (2009) found low carbonate contents, also with very little variation. The stream bed sediments in European rivers are around 286 000 while river alluvial deposits are at 302 000 ppm (median values, De Vos et al., 1996; Salminen et al., 2005; De Vos et al., 2006). The hypothesis of constant PSi content in river particulates induces thus an uncertainty of $\pm 15\%$ on average and probably lower than the uncertainty of particulate matter fluxes. The total PSi flux of rivers depends on their suspended particulate budget. The inputs of PSi to oceans thus greatly depend on the estimates of SPM fluxes by rivers. Ludwig and Probst's (1998) estimate might underestimate some inputs, especially from the smallest basins (basin area <10 000 km²), particularly in South

Table 5. Existing global budgets and values of ranges for dissolved silica (DSi).

Reference	Global average concentration (mg l ⁻¹ SiO ₂)	Approach		
Clarke (1924)	8.3	Extrapolated from few, big temperate rivers		
Livingstone (1963)	13.1	As Clarke (1924)		
Meybeck (1979)	10.4	Biomes typology, 60 rivers, Amazon included		
Probst (1992)	8.9	Multiregression (Meybeck's data)		
Meybeck (1992, unpublished)	9.2	Data later published as Meybeck and Ragu (1995); plus 9 morphotectonic types (lithological control)		
Tréguer et al. (1995)	9.1	Meybeck and Ragu (1995) data		
Meybeck (2003)	8.75	On the totality of the analyses from Meybeck and Ragu (1995) (~260 rivers and tributaries, exo- and endorheic)		
Beusen et al. (2009)	9.7	Same data as here, multiple linear regression model		

East Asia coasts (Milliman and Syvitski, 1992). Recent models give varying numbers (Vörösmarty et al., 2003; Syvitski et al., 2005 and references therein), due to the balance between enhanced inputs owing to land use changes on the one hand, and to enhanced retention by reservoirs on the other (see discussion in Vörösmarty and Meybeck, 2004).

As for any global river budget, it is very much dependant on the analyses of the largest rivers, i.e. Amazon, Congo, Orinoco, Ganges and Brahmaputra, Mississippi, Yangtze, and Great Arctic rivers which are surveyed by water quality authorities and/or by scientists. An error on the Amazon average concentration (15% of the world river discharge) may affect the whole budget. The use of a complex model linking all control factors to silica yields as developed by Probst (1992) or more recently - using the data presented here - by Beusen et al. (2009) for DSi, would not have improved much our mode of extrapolation, since previous multi-regression models are globally not more accurate than 30%. Compare for results of global land-ocean flux estimations of carbon and nutrients and individual discussions (Global Biogeochemical Cycles special issue on Global Nutrient Fluxes from Watersheds (Global-NEWS), edited by Seitzinger et al., 2005; individual discussions are provided on sediment fluxes by Beusen et al. (2005) on nitrogen by Dumont et al., 2005, and on phosphorus by Harrison et al., 2005).

When comparing global average silica concentrations that have been proposed since Clarke's first estimate in 1924, these figures have not changed much: they range between 8.3 and $13.1 \,\mathrm{mg} \,\mathrm{l}^{-1} \,\mathrm{SiO}_2$ (Table 5). Clarke's figure $(8.3 \,\mathrm{mg}\,\mathrm{l}^{-1}\,\mathrm{SiO}_2)$ was an arithmetic average based on the silica content of dry residue from few major rivers, mostly located in the North temperate regions of Europe and North America. Livingstone made first an arithmetic average of DSi for each continent, then these means were weighted by the average runoff of the continents (six continents including Australia) in order to calculate the "global weighted DSi". He obtained a higher value (13.1 mg l^{-1} SiO₂). Since continents are generally heterogeneous, this approach is not very reliable (e.g., Asia average of $11.7 \text{ mg l}^{-1} \text{ SiO}_2$, mostly based on tropical Asia since Livingstone's data set does not contain any Russian rivers). The first average made by Meybeck in 1979 (DSi = $10.4 \text{ mg l}^{-1} \text{ SiO}_2$) was based on 60 rivers distributed in most continents with less weight from the big Arctic rivers, and developed for the first time a simple typology linking silica yields with temperature and runoff to extrapolate the documented data set to unknown regions. The first multi-regression approach was made by Probst in 1992 on the same set of rivers, mostly large ones. It resulted in a $8.9 \,\mathrm{mg}\,\mathrm{l}^{-1}\,\mathrm{SiO}_2$ average, very close to the present average. A previous unpublished attempt for global silica budgets has been made by one of our co-authors (M. Meybeck, unpublished data, 1992) on the basis of a new and larger data set (n = 250), later published as the GEMS-GLORI register (Meybeck and Ragu, 1995). It was based on another typology, based on tectonics and global geomorphology: 9 morpho-tectonic types were defined, such as old and recent sedimentary basins, Hercynian, Caledonian and Alpine mountain ranges, active volcanism etc. The corresponding silica average by this approach was $9.2 \,\mathrm{mg} \,\mathrm{l}^{-1} \,\mathrm{SiO}_2$. A similar level (9.1 mg l^{-1} SiO₂) has been reached by Tréguer et al. (1995) on the basis of the GEMS-GLORI data that was communicated to these authors. The exorheic plus endorheic rivers have also been considered recently (Meybeck, 2003, see also the Supplement), also using exorheic rivers selected by Meybeck and Ragu (1995) plus major endorheic rivers, the corresponding world weighted average is $8.7 \text{ mg } 1^{-1} \text{ SiO}_2.$

Recently, and partly also based on the data compiled here, the major control factors on local, regional and global DSi fluxes have been analysed using empirical techniques. Control factors are usually extracted by using small to medium sized catchments (<10000 km²). Some of the controlling factors, including runoff or precipitation, temperature, relief and lithology, have been identified for specific regions of the globe (e.g., Bluth and Kump, 1994; Hartmann et al., 2010a, for Japan; Jansen et al., 2010 for the USA), as well as globally (Beusen et al., 2009). Runoff/precipitation and lithology were identified in most regional to global scale studies as strong predictors for DSi fluxes. However, it remains difficult to disentangle the strength of further effects at the regional to global scale such as vegetation, temperature or relief from the strong influence of lithology and runoff. For example, while correlations of DSi concentrations (not fluxes) with temperature have been observed for certain data (Garnier et al., 2006), the influence of temperature was not always identified in large-scale statistical models (cf., Beusen et al., 2009; Hartmann et al., 2010a). There is thus still a discussion on how and to what extent this assumed temperature effect can be identified, its strength on DSi mobilization can be estimated robustly, and how it could be employed in regional and global scale models (cf. discussions in Godderis et al., 2009; Hartmann and Moosdorf, 2010; Hartmann et al., 2010a).

Comparing the recent literature, it becomes clear that the findings about controlling factors from smaller catchments are not all representative for large basins or the total global DSi budget. The findings of the recent studies confirm in general our results for the global scale. It should be mentioned that there is still an absence of entirely process-based models for the global scale due to difficulties to globalize or

regionalize important parameters (cf. Sferratore et al., 2005). We believe that the budget approach presented here, considering local and regional characteristics, is an appropriate tool for deriving a global spatial representation of land-ocean flux estimates. This is supported by the outcomes of the work of Beusen et al. (2009) on DSi, if compared with the results presented here.

It must be reminded that riverine budgets should either associate concentrations and discharges, or yields and area. Extrapolations are then carried out based on the hypothesis of constant concentration or of constant yields. The first hypothesis therefore greatly depends on the global water budget, which is now more and more addressed (Oki et al., 2001; Fekete et al., 2002) and has therefore been chosen here. Although the second one is easier to use, as drainage areas are well defined (e.g. in Meybeck's budget, 1979), it greatly depends on the sample of documented rivers, which should have representative average runoff, sediment yields and lithology (for a more detailed discussion see Meybeck, 1988).

4.2 Relations between dissolved and particulate silica exports and yields

Silica yields $Y_{\rm DSi}$ and $Y_{\rm PSi}$, expressed in t SiO₂ km⁻² y⁻¹, are calculated at the river mouths upstream of the land-ocean interface. Actually, there can be differences between whole river yields at mouth and those measured in headwaters, if retention processes of river-borne material are significant, as for particulates (Walling, 1983; Walling and Webb, 1996), organic carbon (Stallard, 1998), and nutrients, including dissolved silica, which can be processed and stored within the aquatic system (Sferratore et al., 2005).

Long-term average dissolved silica mobilized into river systems ultimately origins from silicate mineral weathering, while particulate Si mostly concerns quartz and clay mineral particles 10 to 50 times less soluble than silicates. The ratio DSi (in $\rm mg\,l^{-1}$) to PSi (in ppm) is thus essentially regulated by grain size and not the weatherability of silicates, as they are rarely dominant on quartz and clay minerals in rivers (if this is the case the resulting rocks are arkoses). The ratio of DSi flux to PSi flux is very low as soon as the SPM fluxes increase (ex. Madeira) and much higher when SPM fluxes are low.

The ratio between average $Y_{\rm DSi}$ and $Y_{\rm PSi}$ in COSCATs (Fig. 3) varies over more than three orders of magnitude, from 0.005 to 10. DSi export nearly matches PSi exports in few regions as in the Zaire-Congo river basin (COSCAT #0014) and in some Siberian COSCATs, due to the very limited river transport of particulates in downstream areas of these regions.

PSi export largely dominates in most regions and the $Y_{\rm PSi}/Y_{\rm DSi}$ can exceed 1000 for the Pohai Gulf (COSCAT #1325), which is dominated by the HuangHe (Yellow River) sediment inputs. Such an anomaly in the dissolved vs.

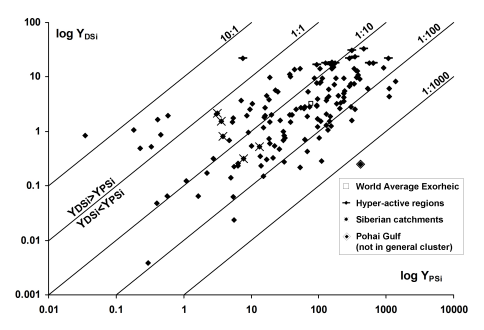


Fig. 3. General relation between dissolved (Y_{DSi}) and particulate (Y_{PSi}) silical yields in the 140 coastal catchments and in the global budget. Y_{DSi} and Y_{PSi} in t SiO₂ km⁻² y⁻¹, log scales.

particulate river material relationship has also been observed for the organic carbon (Meybeck, 1993). This observation is attributed to high physical erosion rates from the loess regions within the catchment. The observed weak positive relation between DSi and PSi yields ($r^2 = 0.41$ for the ~ 140 loglog COSCAT couples) is partly due to their common control factor, runoff.

The general positive relationship between dissolved and particulate Si fluxes (Fig. 3) is in accordance with observed correlations between chemical weathering rates and physical erosion for certain regions or geological settings (Gaillardet et al., 1999; Riebe et al., 2001; Millot et al., 2002; Lyons et al., 2005; West et al., 2005). However, in specific regions of the world opposite trends have been found as for the USA (see a complete discussion in Meybeck, 1994).

The world weighted average DSi/PSi ratio in COSCATs of 4.2% (median 4.0%) is slightly biased towards the particulate material. Regions in which the dissolved silica export is dominant (DSi > PSi) are very rare and correspond essentially to Hudson Bay drainage, i.e. to lake outlets where PSi is completely trapped (also $Y_{\rm PSi}$ is not measured in those areas, but estimated from the Ludwig and Probst (1998) model with many uncertainties in those lake-covered regions). In these environments PSi at lake outlets may be largely biogenic (diatoms).

4.3 Hot spots of river silica inputs to coastal zones

Export rates of silica by rivers vary over more than 3 orders of magnitude for particulates and 2 to 3 orders for the dissolved component (Fig. 3). After yields normalisation to

the world average, the relative rates, scaled to the world's weighted average, $Y_{\rm DSi}/\overline{Y_{\rm DSi}}$, can be compared for any type of river material. Meybeck et al. (2006) have proposed a general classification of relative rates in 7 classes, centred around the world average $(\overline{Y_{\rm DSi}})$:

- Hot regions have $10 < Y_i / \overline{Y_i}$
- Hyper-active regions are characterised by $5 < Y_i/\overline{Y_i} < 10$ (see Fig. 3)
- Eury-active regions by $2 < Y_i / \overline{Y_i} < 5$
- Meso-active regions by $0.5 < Y_i / \overline{Y_i} < 2$
- Hypo-active regions by $0.2 < Y_i/\overline{Y_i} < 0.5$
- Oligo-active regions by $0.1 < Y_i / \overline{Y_i} < 0.2$
- Steno-active regions by $0.01 < Y_i / \overline{Y_i} < 0.1$
- Inactive regions have $Y_i/\overline{Y_i} < 0.01$

For dissolved silica, at the spatial resolution of COSCATs, i.e. circa one million km², there are no true hot spots, but several hyper-active regions are found, as in humid tropical regions and/or wet volcanic regions (Fig. 4). Of particular interest is the Pacific Ocean drainage basin where most hyperactive COSCATs are found: they coincide with the dominance of volcanic outcrops related to the Pacific Ring of Fire from Kamchatka to Java, in Central America, and in the wet coast of South Chile.

In contrast, most of the northern COSCATs are hypoto steno-active (e.g. West Hudson Bay). The Amazon and

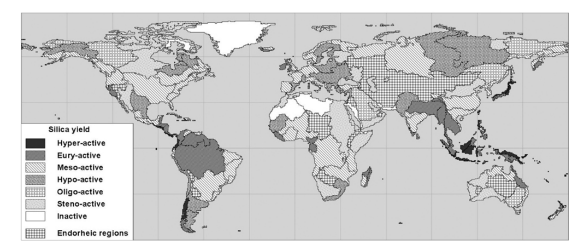


Fig. 4. Relative natural dissolved silica yield for coastal catchments (COSCATs) (115 M km², $n \approx 140$, global mean yield 3.3 t DSi km⁻² yr⁻¹). Inactive areas 0–0.01× global mean yield, steno-active 0.01–0.1x, oligo-active 0.1–0.2x, hypo-active 0.2–0.5x, meso-active 0.5–2x, eury-active 2–5x, hyper-active 5–10x. No Hot spots (>10× global mean yield) are observed at this scale. Silica yield classes boundaries see also Fig. 5. Antarctica and glaciated parts of Greenland not considered. Endorheic regions are set apart.

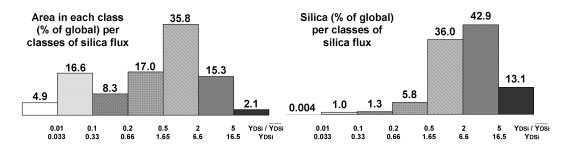


Fig. 5. Relative distribution of dissolved silica yields in exorheic basins (115 M km², $n \approx 140$). Classes are constructed after the mean yield of 3.3 t DSi km⁻² yr⁻¹ (see text). Antarctica and glaciated parts of Greenland not considered.

Orinoco regions and the Asian monsoon regions, from the Ganges to the Mekong, are eury-active, i.e. between 2 and 5 times the world average DSi yields. Maximum dissolved silica yields are found in subtropical and tropical, as well as in wet volcanic regions (Jennerjahn et al., 2006; Hartmann et al., 2010a). For example, the yield of the Japanese river Tokachi, included in the database, is $37 \, t \, SiO_2 \, km^{-2} \, y^{-1}$ $(DSi = 29.9 \text{ mg SiO}_2 \text{ l}^{-1}, \text{ runoff } q = 1240 \text{ mm y}^{-1}, \text{ catch}$ ment area 8800 km²) (Kobayashi, 1960). As silica concentrations are even higher in volcanic and hydrothermal regions as in Kamchatka (e.g., 41.9 mg l⁻¹ SiO₂ for a spring close to Koryaksky volcano; Hartmann, 2006), and fresh unweathered ashes are capable to release large amounts of DSi, high $Y_{\rm DSi}$, exceeding 50 t km⁻² y⁻¹, are possible at the local scale (Hartmann et al., 2010a). Other hot spots of silica fluxes combine extreme runoff and active tectonics as for the Purari river (New Guinea) which exports 35.2 t SiO₂ km⁻² y⁻¹ $(DSi = 13.8 \text{ mg SiO}_2 \text{ l}^{-1} \text{ and } q = 2750 \text{ mm y}^{-1}, \text{ data from } 1.2 \text{ mg SiO}_2 \text{ l}^{-1}$ Petr, 1983, see also Supplement 1F).

A similar mapping as for DSi can be made for PSi. If the particulate Si content in river SPM could be considered as

constant (±26 wt%) for the regional scale, the pattern of PSi yield is identical to the one observed for SPM fluxes from models, which show that most land-ocean fluxes originate from the erosion of the Himalayas, South East Asia island arcs and archipelagos, from Taiwan to New Zealand (Milliman and Syvitski, 1992; Ludwig and Probst, 1998). Note that we have not focused on spatial BSi distribution here, because the bioavailable part of PSi (=BSi) is actually still under discussion, without a definitive emerging number and we found that the few available studies providing local data do not provide a large enough data base for an appropriate global spatialization (cf. Conley, 1997).

When the global silica budget is disaggregated for the 7 relative flux classes, from inactive to hyper-active regions (Fig. 5), the following observations can be made:

- 17.4% of the exorheic area contribute to 55% of the dissolved silica fluxes (eury- to hyper-active regions).
- 28.9% of the exorheic area contribute to only 2.3% of fluxes (inactive, steno- and oligo-active regions).

This skewed distribution of riverine fluxes is very general and has been observed for water, suspended matter and nutrients as total nitrogen (Meybeck et al., 2006).

4.4 Towards spatial estimates of net river inputs to oceans

At the beginning of global bio- and geochemical cycles studies, significantly influenced by the work of Mackenzie and Garrels (Mackenzie and Garrels, 1966; Garrels et al., 1973), the amount of data on world rivers was scarce, GIS did not exist nor global models. Therefore it was difficult to spatialize river inputs at global scale. The global budget of silica (Tréguer et al., 1995) was also not spatialized. The first major improvement was the use of GIS and multi-regressions to build up spatialized models of river inputs to oceans at the basin scale (Probst, 1992; Ludwig et al., 1996; Amiotte-Suchet et al., 2003; Beusen et al., 2009). These river material budgets were then used in regional ocean biogeochemical models as for organic carbon (Aumont et al., 2001). However, they did not take into account the natural filters between river basins and the open ocean as estuaries, shelves and regional seas (Meybeck and Dürr, 2009).

The processing and retention of dissolved silica in estuaries and its potential impact on coastal food webs has been demonstrated since several decades (Nixon, 1987; Billen and Garnier, 1997; Rabalais and Turner, 2001). The retention of riverine silica and other nutrients on the shelf is also well documented (DeMaster et al., 1983; Rabouille et al., 2001; Dagg et al., 2004). However, none of these processes have been addressed at the global scale.

The main transformations occurring to silica in near-shore and estuarine environments are (i) its uptake by silicifying organisms such as diatoms under its dissolved form to produce biogenic silica (Roubeix et al., 2007 and references therein), (ii) the dissolution of the particulate pool within the water column or the upper layer of the sediment which regenerates dissolved silica (Yamada and D'Elia, 1984) and (iii) the permanent storage of the particulate silica through sediment accumulation which constitutes the sink term of the system and leads to long-term Si retention (Nixon, 1987). The importance of a proper representation of the particulate biogenic pool of silica was pointed out by several authors (Conley, 2002 and references therein). The fraction of PSi that is considered possibly dissolvable and biologically available is currently not well known. Most of the PSi is in mineral form that will dissolve only very slowly and mainly be buried and re-mineralized in marine sediments. A small fraction of PSi is constituted of biological material (i.e., BSi), mostly phytoliths and diatom frustules. This BSi is considered orders of magnitude more easily soluble than the remainder of PSi, i.e. it can become bioavailable, and can thus contribute to algal growth (van Cappellen, 2003). The BSi fraction in PSi is difficult to estimate as it depends very much on the river basin considered and will be linked to control factors such as forest distribution, land use, water turbidity, or any factor likely to influence diatom productivity. Conley (1997) estimated the global BSi load to estuaries at $63\pm7.2\,M\,t\,SiO_2\,y^{-1}\,$ (1.05 $\pm\,0.2\,Tmol\,Si\,y^{-1}$), i.e. 17% of the total DSi flux. This represents just 0.75% of the total PSi export from the rivers, and, considering the lithogenic fraction of the PSi as biologically inert, the available fraction might thus be <1% (see Table 1), due to the overwhelming mineral composition of PSi. This fraction can be seen as the maximum amount of PSi potentially recycled and biologically available at shorter timescales and is 5 to 6 times smaller than the pool of DSi.

However, direct measurements of BSi fluxes in budgets are still scarce and, for instance, BSi production in coastal environments is often deduced from primary production or DSi consumption (Beucher, 2003). Estimates of the net uptake of silica in estuaries deduced from estuarine mixing curves, i.e. DSi vs. salinity relationship from rivers to sea waters, are still the most common type of studies available in the literature and both excess and depletion patterns of DSi in estuarine environments have been observed, they vary seasonally and between stations. Net budgets to oceans, combining silica patterns and river discharges are very limited and/or biased (for example, summer profiles are much more common than profiles at river flood stage) (DeMaster, 1981; Aston, 1983; Balls, 1994; Rendell et al., 1997; Kimmerer, 2005; Chou and Wollast, 2006). However, primary production is a typically seasonal process driven by light and temperature conditions and varies greatly through the year as do river inputs. Hence, extrapolating an annual flux from these estimates is a difficult task which also does not account for the potential transient storage within the estuary itself (Webster et al., 2000). Moreover, the net uptake is a transfer from the dissolved to the particulate pool but does not indicate the real retention of silica within the system. For that matter, the work of De-Master (2002) to quantify the accumulation of particulate silica within the sediment remains one of the only references at global scale but is essentially based on carbon measurements and molecular ratios. Recent studies also indicate that this retention can be influenced by "reverse weathering", a process suggested by Mackenzie and Garrels (1966), and describing formation of new aluminosilicate minerals from BSi in marine sediment by chemical combination of elemental material from oceans and marine sediments (Mackenzie and Kump, 1995; Michalopoulos and Aller, 1995; Michalopoulos et al., 2000). The quantity that is removed by reverse weathering reactions is assumed to be a small fraction of the total river input, but is currently not precisely known (Holland, 2005). Results for the Amazon delta suggest that \sim 90% of the biogenic silica originally present in deposits is converted to clay or otherwise altered and the deltaic storage of riverine Si is estimated to be ~22% of the Amazon River input (Michalopoulos and Aller, 2004). Further retention may result from the activity of benthic organisms that enhance benthic-pelagic coupling (Ragueneau et al., 2005) but large

Table 6. Dissolved silica fluxes, concentration and yield for selected regional sea catchments: North Atlantic Regional Seas and Japan Sea.
Catchment surface area, discharge and runoff are given from Meybeck et al. (2007) as well as % of volcanic rocks from Dürr et al. (2005)
and lake density from Lehner and Döll (2004).

	Catchment area 10 ⁶ km ²	Discharge km ³ y ⁻¹	Runoff mm y ⁻¹	Lake density %	Volcanic rocks (Vb+Va+1/2 Cl) %	SiO ₂ flux Mt y ⁻¹	SiO_2 mean concentration $mg l^{-1}$	SiO ₂ yield t km ⁻² y ⁻¹
Baltic Sea	1.6	390	240	6.4	2.8	1.5	3.9	0.9
Black Sea	2.4	410	170	0.78	8.7	2.1	5.1	0.9
(incl. Azov)								
Mediterranean Sea	8.3	800	97	1.3	6.0	3.3	4.1	0.4
Hudson Bay	4.0	850	210	10.5	2.4	1.7	1.9	0.4
St. Lawrence Gulf	1.6	630	402	19.4	2.4	3.3	5.2	2.1
N. American Med. Seas	6.5	1875	290	0.95	6.9	22.7	12.1	3.5
(Gulf of Mexico, Caribbean)								
Japan Sea	0.4	240	580	_	39.2	4.1	17.1	9.9

scale extrapolations remain to be calculated. One may also refer to the concept of proximal and distal limited filters for the coastal ocean (Rabouille et al., 2001).

Overall, this relative lack of specific data, combined with the heterogeneity of the coastal zone and the question of its precise definition (Smith and Hollibaugh, 1993), makes the quantification of global budgets a very difficult task and stresses out the need for a coastal typology of coastal systems especially for small to medium-sized systems. The world's largest rivers, however, often generate large plumes which consist in an "external" estuary dominating the biogeochemical transformations occurring in the adjacent continental shelf. Therefore, a good understanding of these Riverdominated Ocean Margin (RiOMar, McKee, 2003) systems and their plumes would help significantly understanding and quantifying the fate of silica in the coastal zone (Johnson et al., 2006). Differences between estuarine systems where the major processing occurs near-shore and large systems such as the Amazon (DeMaster et al., 1983; Michalopoulos and Aller, 2004), the Zaire/Congo with its submarine canyon (Chou and Wollast, 2006), the Mississippi (Rabalais and Turner, 2001) or the Danube delta (Ragueneau et al., 2002), should be regarded with greatest attention (Liu et al., 2010).

It remains thus very difficult to quantify these processes and resulting fluxes at the global scale, considering the number of transformations involving silica that can potentially occur in estuaries, river plumes or shelf areas. To our knowledge, only one other study has summarised the existing knowledge of the global silica cycle for the various silica pools and transformation processes (Laruelle et al., 2009).

In addition to the silica transformation and retention processes in the coastal zone, one should also consider the retention in major regional seas of the world. They have been considered as mega filters within an Earth System analysis (Meybeck et al., 2007; Meybeck and Dürr, 2009). Five types of mega filters have been defined: three types of regional

seas, depending on their connection to the open ocean, plus extended platforms (e.g. Siberian seas) and coastal archipelagos (e.g. Canadian Archipelago and South Chile Coast). Their related drainage area and riverine fluxes have been tabulated using a specific re-aggregation of the COSCATs mega puzzle. On the basis of simple assumptions relating the retention of particulates to the morphology of these mega filters and the retention of nutrients to their theoretical river water residence time, which exceeds three orders of magnitude between the Saint Lawrence Gulf (fastest renewal) and the Red Sea (longest renewal), it has been estimated that (Meybeck et al., 2007):

- Regional seas intercept 39% of the exorheic continental landmass (glaciated areas as most of Greenland and Antarctica excluded), and 35% of the exorheic river runoff.
- ii. Regional seas store about 80% of the incoming related particulate fluxes; this proportion will be similar for the riverine particulate silica.
- iii. Regional seas intercept 33% of the related dissolved silica of which about 75% might not be exchanged with the open ocean.

Some examples of the river silica fluxes to regional seas as calculated by COSCATs aggregations are featured on Table 6. Their ranges in terms of average concentration (1.9 to $17.1 \, \text{mg} \, l^{-1}$) and yields (0.4 to $9.9 \, t \, \text{km}^{-2} \, y^{-1}$) are over one order of magnitude and illustrate the multiple control factors already presented.

For each individual ocean drainage basin the actual silica retention in regional seas should now be estimated; it may be quite different from these global figures (Meybeck et al., 2007): the Southern Atlantic Ocean has no mega filter except for the Patagonia platform, therefore retention is very limited.

The North Atlantic Ocean is very different from the South Atlantic, as it is connected to many regional seas, often interconnected in nested basins as the Mediterranean plus Black Sea, the Gulf of Mexico plus Caribbean – also termed North American Mediterranean –, the Baltic Sea, Hudson Bay and St. Lawrence Gulf (see Table 6 for individual budgets). Their potential silica retention is large (40% for PSi, 20% for DSi, applying the filter types proposed by Meybeck et al., 2007).

The Pacific Ocean is also very much affected by regional seas: (i) its drainage area is already very limited (19.8 M km²) drainage basin area/181.3 M km² Pacific Ocean area = 0.11), compared to the world average (114.7 M km² total exorheic basin area/362 M km 2 total ocean area = 0.32) or to the Arctic Ocean (16.7 M km² drainage basin area/12.3 M km² Arctic Ocean area = 1.4); (ii) a larger proportion of its western drainage area (83%) is actually linked to the West Pacific Mediterranean Regional Seas from Okhotsk to Banda and Sulu Sea (Meybeck et al., 2007); (iii) the Bering Sea, the British Columbia and S. Alaska coast archipelago type of mega filter may also retain an important share of the riverine dissolved silica from the Yukon River to the Fraser River. The net inputs to the open Pacific are probably only half of the riverine inputs arriving at the mouths of the rivers. This is taking into account the position of the regional seas basins only. If the estuarine and shelf filter is considered, this retention might even be higher.

In the Arctic Ocean the mega filters consist mostly of the Siberian platform – a limited mega filter, also termed Siberian Seas, much more connected to the open ocean than the enclosed and semi-enclosed West Pacific regional seas. The consideration of mega-filters is therefore increasing the discrepancy between the Arctic Ocean, relatively well fed by river silica per unit volume of open sea – i.e. without the regional seas and other mega filters –, and the open Pacific Ocean, very much deprived of river silica.

The Indian Ocean has a limited connection to regional seas (e.g. Andaman Sea, Persian Gulf, Red Sea), which are often characterized by minimal river inputs, as for the Red Sea. To our knowledge, the retention of silica inputs from the Ganges-Brahmaputra-Meghna rivers in the Bay of Bengal remains to be established. If it is recognized, the open waters of the Indian Ocean might receive less silica than those of the South Atlantic (Liu et al., 2010).

Open oceans are generally deprived of DSi at their surface due to bioactivity when compared to deeper waters (Levitus et al., 1993). Only in some regions, where COSCATs characterised by high proportions of volcanic rocks drain to the open ocean (e.g., Indonesia, Hawaii, Kamchatka), some increases in surface water DSi concentrations can be observed (if Antarctica and northern Arctic waters are disregarded). Upwelling onto continental margins and subsequent storage due to sedimentation can be a sink for oceanic silica (Tréguer et al., 1995; Laruelle et al., 2009), but these "reverse" fluxes of silica in the land-ocean continuum can currently only be estimated by ocean models and since they do not concern

continental silica fluxes to the oceans, they are not discussed further here.

Besides riverine inputs, atmospheric deposition of DSi and PSi is a further important source of "new Si" to the surface oceanic system. Atmospheric PSi dust deposition estimates range globally from 278-489 Mt yr⁻¹ of dust, and 79 to 128 Mt yr⁻¹ PSi from dust is deposited onto the oceans (Tegen and Kohfeld, 2006), with highest amounts reported for the North Atlantic and the western North Pacific. However, the contribution of atmospheric deposition to the oceanic DSi budget is difficult to estimate and can be attributed to large spatial and temporal differences in DSi in atmospheric deposition as well as uncertainties in quantification of DSi removal from PSi dissolution (cf. Zhang et al., 2005; Baker et al., 2006; Hartmann et al., 2008). For example, it was found that the atmospheric dissolved silica deposition rate (DSi-R), including relatively fast soluble DSi from solid matter in the bulk deposition sampler, on a Japanese watershed, close to the sea, was higher for periods with increased abundance of dust storm events. However, the ratio of DSi-R to solid matter deposition rate (SM-R) decreased with increasing SM-R. The overall observation range (monthly samples over 12 years) suggests a DSi-R/SM-R ratio of about 8 wt%. This high observed ratio is to some part attributed to influences of anthropogenic fly ash with notable amounts of amorphous silica for the East Asian region (cf. Misran et al., 2007; Hartmann et al., 2008), but this amount is globally not equally distributed in aerosols. In general, reported Si-solubility from deposited particulate matter into surface waters ranges from 0.02-1.1% for aerosols from Saharan dust events deposited over the Atlantic (Baker et al., 2006) to theoretical estimates of 5–10% (Harrison, 2000; Tréguer et al., 1995). However, these solubility rates are based on the assumption of different dust mineral compositions. For example, Saharan dust contains large amounts of quartz whereas Harrison (2000) uses kinetic considerations to derive potential solubility estimates and assumes that aerosol particles contain large quantities of feldspar. For global ocean models calculating at time scales ranging from years to decades, DSi input from dust is often neglected as it is assumed that deposited aeolian silica would be deposited as sand particles whose sinking speed would not allow any dissolution resulting in a significant input of silicic acid (Johnson et al., 2006). In consequence, a fair estimate of DSi input via the atmosphere into the oceanic system can not be provided, except for the value provided by Tréguer et al. (1995) for DSi inputs via rain water $(0.5 \,\mathrm{Tmol}\,\mathrm{y}^{-1}, \,\mathrm{i.e.}\,30\,\mathrm{Mt}\,\mathrm{SiO}_2\,\mathrm{y}^{-1}\,(\mathrm{DSi})),\,\mathrm{which}$ will be used here for our attempt of a global land-to-ocean budget (see below). However, estimated global DSi and PSi deposition on the ocean is much lower than the estimates for riverine DSi. If 30 Mt SiO₂ y⁻¹ are deposited over oceans, and we assume a similar deposition over land, $11 \,\mathrm{Mt} \,\mathrm{SiO}_2 \,\mathrm{y}^{-1}$ (9.5 $\,\mathrm{Mt} \,\mathrm{SiO}_2 \,\mathrm{y}^{-1}$ over the exorheic area) might be deposited via rain over continents (362 M km² of ocean area, $133 \,\mathrm{M\,km^2}$ of continental area with $115 \,\mathrm{M\,km^2}$ being exorheic, according to the numbers in Meybeck et al., 2006, 2007 and Vörösmarty et al., 2000a, b).

The amount of DSi retained in lakes and floodplains can be estimated as a rough first-order calculation as follows: from the work by Lehner and Döll (2004), lake surface can be calculated, plus a certain proportion of the share of a region covered by floodplains; swamps and wetlands. From the share in a given area that is covered by floodplains, swamps and wetlands, we assume 15% of this share to be permanently flooded – similar to the value proposed by Laruelle et al. (2008) using a global silica box model. This calculation (100% of classes 1, 3, and 15% of classes 4, 5, 9-12 in Lehner and Döll, 2004) results in a total surface area of 3.4 M km² of exorheic area occupied by lakes and permanently inundated floodplains, swamps and wetlands. If an average dissolved silica retention of $20 \pm 10 \,\mathrm{g}\,\mathrm{m}^{-2}\,\mathrm{y}^{-1}$ is considered (see above, Campy and Meybeck, 1995), this results in $69 \pm 35 \,\mathrm{Mt} \,\mathrm{SiO}_2 \,\mathrm{y}^{-1}$ retained in continental areas connected to the oceans. This number corresponds to 1.2 Tmol Si and is somewhat lower than the 2.8 Tmol calculated by Laruelle et al. (2009) as preserved in sediments, but their number includes the volume of rivers, in addition to lakes and floodplains. Yet the average DSi retention in lakes should be confirmed with a larger data set.

Human impacts on the silica fluxes have not been targeted here. They can affect both sources and sinks. As diatomites and/or artificially produced zeoliths are often used in washing powders and other household products (2-3 million metric tons of SiO₂, Breese, 1994; van Dokkum et al., 2004), an increase of silica due to urban impacts is possible. Such inputs have been estimated to 780 g SiO₂ (for DSi) per capita per year (1 g of Si per capita per day) by Sferratore et al. (2006) on Paris megacity (10 million people). The extrapolation to 1 or 2 billions of similar users would correspond to a load of 0.78 to 1.56 Mt DSiO₂ per year, compared to a river flux of 371 MtDSiO₂ y⁻¹ to coastal zones. Based mainly on data from Europe, van Dokkum et al. (2004) estimate the total additional flux due to anthropogenic emissions to be <=2% of riverine DSi fluxes. These fluxes might thus be of local importance (e.g., Seine), and are very minor at global scale, but such extrapolation remains poorly constrained.

Additional sinks correspond to the 100 000 and more reservoirs already registered (Vörösmarty et al., 2003). They correspond to at least 0.5 M km² area of registered impoundments, and have been globally estimated to a total area of 1.5 M km², if smaller and unregistered reservoirs are taken into account (St. Louis et al., 2000). Considering again, as for lakes, an average dissolved silica retention of $20 \pm 10 \, \mathrm{g} \, \mathrm{m}^{-2} \, \mathrm{y}^{-1}$ (Campy and Meybeck, 1995), the total DSi retention would be of the order of $30 \pm 15 \, \mathrm{Mt} \, \mathrm{SiO}_2$ retained each year in reservoirs. This number can be adjusted to $26.4 \pm 13.2 \, \mathrm{Mt} \, \mathrm{SiO}_2$ (i.e., less than 10% of the total annual DSi flux to coastal areas) if the distribution of 88% reservoir area to exorheic regions is taken into account (calculated

using the database - containing major reservoirs only - by Lehner and Döll, 2004). This number is lower than the 18-19% retention calculated by Beusen et al. (2009) based on sediment or inorganic phosphorus retention rates. It must be noted that the retention rate of silica per unit lake or reservoir area considered is about seven times higher than the average weathering rate of silica-bearing minerals. This retention has already been suggested as responsible for the decrease of silica in some large river basins as the Mississippi, where the N:Si ratio has widely fluctuated over the last 100 years (Rabalais and Turner, 2001). Other cases of silica-depleted coastal waters have also been reported for Swedish rivers and in the Danube River delta (Humborg et al., 1997, 2000, 2006), invoking also alternative human impact mechanisms such as reduced connectivity with usually DSi-rich aquifers due to hydraulic alterations, resulting in less exchanges with the riparian zone or lower natural water levels controlling silicon weathering rates (Teodoru et al., 2006). While there is now a considerable amount of literature (e.g., Vörösmarty et al., 2003; Syvitski et al., 2005) on the anthropogenic overprint of suspended matter fluxes, it is outside the scope of this paper to add to this discussion. For suspended particulate matter (SPM), and thus detrital PSi, global retention in reservoirs (yet partly balanced by increased deforestation leading to enhanced denudation) is now estimated between 20% (Syvitski et al., 2005) and 30% (Vörösmarty et al., 2003). Additional retention of DSi can occur in main stems of rivers and is closely related to anthropogenic influence on nitrogen (N) and phosphorus (P) levels. Significant amounts of BSi resulting from plankton blooms can be sedimented in flood plains (Admiraal et al., 1990). As early data from European rivers suggests (see above), DSi retention due to N and P fertilization may possibly occur since the middle of the 19th century (Roth, 1879). N and P fertilization might thus cause decreases in DSi levels in urbanized or agricultural catchments. When recent decreases of N and P in managed rivers are observed, it has been noted that DSi concentrations might eventually come back to nearly pristine levels, as data from the Rhine River suggest (Hartmann et al., 2010b). It must be noted that long-term records of DSi in rivers are much less common than those for N and P.

Retention in lakes, floodplains, and human-built reservoirs is very much dependant on water residence times in these bodies, and our simple calculation for the global scale should be verified by further studies. The assumed retention rate of $20\pm10\,\mathrm{g\,m^{-2}\,y^{-1}}$ by Campy and Meybeck (1995) is a crude estimate that has been derived for few lakes, and is probably lower for reservoirs and highly anthropised floodplains, due to lower residence times.

If our first-order back-of-the-envelope estimate, proposed here as a working hypothesis, is validated in forthcoming studies, the total amount of dissolved silica in headwaters, derived from weathering on the exorheic parts of the continents should be increased by as much as $59 \, \text{Mt SiO}_2$ per year (additional retention of $69 \pm 35 \, \text{Mt SiO}_2$ as BSi in lakes,

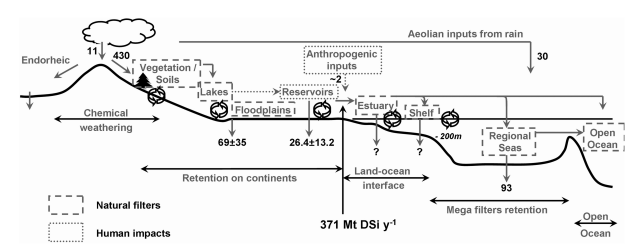


Fig. 6. Conceptual pathways of dissolved silica from land to oceans with cascading filters on land and nested filters at the land-ocean interface. Dotted: anthropogenic components. All numbers are in Mt DSi per year.

floodplains, swamps and wetlands for the natural situation, minus $9.5\,\mathrm{Mt}\,\mathrm{SiO}_2$ from atmospheric deposition), assuming that riverine DSi derived from re-dissolution of BSi ultimately also stems from weathering sources. While there is still considerable uncertainty on control factors, the budget seems well constrained.

A schematic view on the silica transfers from land to ocean with the cascading filters, both natural and anthropogenic, and new anthropogenic inputs that characterize the Anthropocene (Crutzen and Stoermer, 2000) situation of river systems described by Meybeck and Vörösmarty (2005), is presented in Fig. 6. It is combined with the nested filters of river inputs to oceans: estuaries, shelf, regional seas and other mega filters.

5 Conclusions and perspectives

The spatialized distribution of riverine dissolved silica fluxes has been realised for \sim 140 coastal catchments (COSCATs). The documented area and river runoff correspond to 56 and 58% of the globe, respectively, and the documented river DSi fluxes to 51%. The undocumented part is extrapolated on the basis of modelled runoff, and of regional similarities of dissolved silica based on lithology, climate and lake occurrence. Some of the COSCATs are still lacking analyses (e.g. Madagascar island and parts of Australasia), but the spatial coverage of the 250+ rivers used in this budget is not biased for any of the control factors. The resulting overall dissolved silica average (9.5 mg l $^{-1}$) is not much different from previous attempts of the last decade, but the spatialization of average concentrations and yields for each piece of the COSCAT puzzle is new.

At such resolution (ca. 1 M km²), the average DSi concentrations for COSCATs range over one order of magnitude and a factor of 50 for DSi yields. The highly skewed distribution

of yields results in a marked imbalance of riverine fluxes: at the global scale, 55% of dissolved silica fluxes originate from only 17.4% of the land area facing the oceans, while 28.9% of this area contribute to only 2.3% of fluxes. Similar skewed distributions have been found for many other river-borne materials. Higher fluxes originate from different COSCATs, encountered on many continents: they correspond to high to very high runoff rates, combined with medium and high relief, volcanic and/or highly active tectonic regions, the major control factors of silica sources. The occurrence of multiple lakes or of single large lakes (e.g. Saint Lawrence basin, COSCAT #825, with the Great Lakes, or Gulf of Finland basin, COSCAT #0406 with the Neva River showing one of the world's lowest silica yields due to the presence of Lake Ladoga close to its outlet, combined to a cold climate) is probably a major control factor of silica retention on continents.

Partitioning silica outputs between individual continents has little interest since they generally combine both high and low yields as demonstrated when considering individual COSCATs.

The riverine silica inputs to individual oceans, required by ocean biogeochemists, show great disparities when considering relative fluxes received per unit ocean area or volume: there is nearly one order of magnitude difference between the fluxes to the Arctic Ocean (silica over-fed) and the Pacific (silica-starving). This type of computation should now also take into account the nested filtering effects of estuaries, shelves and regional seas: the discrepancy between the silica fluxes received by the Pacific after the potential retention of DSi by regional seas may be even greater, compared to those received by the Arctic.

When comparing the riverine yields of dissolved silica (Y_{DSi}) and of particulate silica (Y_{PSi}) – at the COSCAT resolution, assuming a constant silica content around 55% SiO₂

in river suspended matter – a weak positive correlation is found, with the Pohai Gulf (COSCAT #1325) including the HuangHe basin set aside. The $Y_{\rm DSi}/Y_{\rm PSi}$ ratio in COSCATs ranges from more than unity to 10^{-4} (HuangHe). A majority of COSCATs has a ratio between 0.005 and 0.5, compared to the global flux ratio ($\overline{Y_{\rm DSi}}/\overline{Y_{\rm PSi}} = 0.042$). 9% of the COSCAT area (10.3 M km²) are constituted by regions where the particulate transfers are very high ($\overline{Y_{\rm DSi}}/\overline{Y_{\rm PSi}} < 0.01$), as for the Pohai Gulf.

Future development of ocean budgets should therefore aim at regional budgets in which river inputs are spatially distributed (Liu et al., 2010). These biogeochemical models combine all river nutrients (N, P, Si) and carbon species (Sferratore et al., 2005), including the retention impact of reservoirs that is not yet fully quantified at the global scale.

The improvement of the silica budget coverage, e.g. from the actual 50–60% coverage to 75%, will not be easily achieved. There are only few major rivers, such as the Salween and Irrawaddy, that remain undocumented in the database used here and the inclusion of hundreds of small and medium basins (0.01 to 0.1 M km²) would be needed: the first 10 rivers, ranked by basin area, correspond to about 26% of the global exorheic area, the first 100 to 62% and the first 1000 to 74% only (Milliman and Syvitski, 1992; Meybeck and Ragu, 1995).

Further research should first estimate each of the components of the global silica cycle (Fig. 6), then establish these fluxes for individual regions and/or ocean basins and we have seen that major disparities exist in silica concentrations or yields.

A most promising approach is represented by biogeochemical models that are spatially distributed at very high resolution linking sources (weathering) and sinks (uptake, sedimentation of biogenic silica), as already developed for the Seine River (Sferratore et al., 2005) and the Red River (Hong) in Vietnam, in connection with N and P models (Quynh et al., 2005). This type of model is often datalimited and should now be validated for different types of river basins. Also the analysis of riverine particulate biogenic silica, either detrital or phytoliths, or autochthonous, should be further developed, particularly in highly eroded river basins, and compared to the particulate organic carbon. These models will be able to simulate the silica retention in reservoirs and explore Global Change scenarios on the riverine components of the silica cycle.

Supplementary material related to this article is available online at: http://www.biogeosciences.net/8/597/2011/bg-8-597-2011-supplement.pdf.

Acknowledgements. This paper is dedicated to Fred Mackenzie and Robert Berner who paved the way of global riverine chemistry. The original idea for this work came from the Si-WEBS project, an EU Research Training Network. This project was made possible by

financial support from Utrecht University (High Potential project G-NUX) and the EU (Si-WEBS, contract number HPRN-CT-2002-000218). The work of Jens Hartmann is funded by the German Science Foundation (DFG-project HA 4472/6-1 and the Cluster of Excellence "CliSAP" (EXC177), Universität Hamburg). Charles Vörösmarty (Water Systems Analysis Group, University of New Hampshire, USA) is acknowledged for the communication of various data sets. We thank two anonymous reviewers who helped to clarify and improve the manuscript.

Edited by: J. Middelburg

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