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Abstract

The quantification of silicon isotopic fractionation by biotic and abiotic processes contributes to the understanding of the Si continental cycle. In soils, light Si isotopes are selectively taken up by plants, and concentrate in secondary clay-sized minerals. Si can readily be retrieved from soil solution through the specific adsorption of monosilicic acid (H₄SiO₄) by iron oxides. Here, we report on the Si-isotopic fractionation during H₄SiO₄ adsorption on synthesized ferrihydrite and goethite in batch experiment series designed as function of time (0–504 h) and initial concentration (ic) of Si in solution (0.21–1.80 mM), at 20 °C, constant pH (5.5) and ionic strength (1 mM). At various contact times, the δ²⁹Si vs. NBS28 compositions were determined in selected solutions (ic = 0.64 and 1.06 mM Si) by MC–ICP–MS in dry plasma mode with external Mg doping with an average precision of ±0.08 (±2σSEM). Per oxide mass, ferrihydrite (74–86% of initial Si loading) adsorbed more Si than goethite (37–69%) after 504 h of contact over the range of initial Si concentration 0.42–1.80 mM. Measured against its initial composition (δ²⁹Si = +0.01 ± 0.04 (±2σSD)), the remaining solution was systematically enriched in ²⁹Si, reaching maximum δ²⁹Si values of +0.70 ± 0.07 for ferrihydrite and +0.50 ± 0.08 for goethite for ic 1.06 mM. The progressive ²⁹Si enrichment of the solution fitted better a Rayleigh distillation path than a steady state model. The fractionation factor ²⁹ε (±1σSD) was estimated at −0.54 ± 0.03 for ferrihydrite and −0.81 ± 0.12 for goethite. Our data imply that the sorption of H₄SiO₄ onto synthetic iron oxides produced a distinct Si-isotopic fractionation for the two types of oxide but in the same order than that generated by Si uptake by biota and diatoms. They further suggest that the concentration of light Si isotopes in the clay fraction of soils is partly due to H₂SiO₄ sorption onto secondary clay-sized iron oxides.

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1. INTRODUCTION

Silicon is the second mass abundant element of Earth’s crust and a major solute in river discharge into oceans (Tréguer et al., 1995; Gaillardet et al., 1999). Breakdown of primary silicates, translocation of Si in solution, formation of secondary silicates, and Si uptake by plants are involved in the continental cycle of Si. Dissolved Si is present as monosilicic acid (H₄SiO₄) in natural solutions (Lindsay, 1979), where it is commonly a major solute (McKeague and Cline, 1963). Apart from clay formation and uptake by biota, monosilicic acid can be withdrawn from soil solution through its sorption onto aluminum and iron oxides (Beckwith and Reeve, 1963; Jones and Handreck, 1963; McKeague and Cline, 1963). Iron oxides are ubiquitous in sediments, weathered rocks and soils (Schwertmann and Taylor, 1989), where they partly control the concentration of aqueous silicic acid (McKeague and Cline, 1963; Gehlen and Van Raaphorst, 2002). Their surface OH groups specifically interact with silicic acid by exchanging ligands to form a bi-dendate innersphere complex involving mono-
meric SiO$_2$(OH)$_2$$^{2-}$ as main adsorbed species at low loading of aqueous Si (below ~0.9 mM) (Hingston et al., 1967; Sigg and Stumm, 1981; Hansen et al., 1994b; Swedlund and Webster, 1999; Davis et al., 2002; Pokrovski et al., 2003; Hiemstra et al., 2007). At higher loading of aqueous Si (>0.9 mM), however, Si tetramer can form on goethite surface (Hiemstra et al., 2007), whereas H$_2$SiO$_4^-$ polymerization occurs on ferrihydrite surface at H$_2$SiO$_4^-$ concentrations below those required for polymerization in solution (Swedlund and Webster, 1999). The adsorption of Si by Fe oxides strongly varies with pH, and commonly reaches a maximum around pH 9 (Jones and Handreck, 1963; Sigg and Stumm, 1981; Hansen et al., 1994a; Swedlund and Webster, 1999; Hiemstra et al., 2007).

Following weathering, iron oxides accumulate in soils (Bikerland, 1974), where they appear as both crystalline and short-range ordered (sro) minerals mainly in their clay-sized fraction (Schwertmann and Taylor, 1989). They can exert a crucial impact on the retention of Si and the control of aqueous H$_2$SiO$_4^-$ polymerization (McKeague and Cline, 1963). Quantifying the fractionation of Si stable isotopes by biotic and abiotic processes readily contributes to the understanding of the continental cycle of Si. As inferred from δ$^{30}$Si and/or δ$^{33}$Si values, river waters are depleted in light Si isotopes (De La Rocha et al., 2000; Ding et al., 2004; Alleman et al., 2005; Georg et al., 2006, 2007) compared with crustal rocks (Douthitt, 1982), following fractionating processes of silicate weathering and formation of clay-sized minerals (Ziegler et al., 2005a,b; Opfergelt et al., accepted for publication), silcrete formation (Basile-Doelsch et al., 2005), and Si uptake by biota (De La Rocha et al., 1997; Ding et al., 2005, 2008; Opfergelt et al., 2006a,b).

In this paper, we report on an experimental study on the isotope fractionation of Si by H$_2$SiO$_4^-$ adsorption onto iron oxide. The experiments were designed to answer four questions of environmental significance: (1) Does Si isotope fractionation occur during Si adsorption onto Fe oxide at common pH for soil solutions? (2) How much does this fractionation compare with that generated by Si uptake by biota? Can this fractionation contribute to (3) the depletion of light Si isotopes in river waters, and (4) the enrichment of light Si isotopes in clay-sized soil fractions? For these purposes, we use goethite and ferrihydrite, crystalline and sro minerals, respectively. Our experimental study involves the prior synthesis and characterization of pure Fe oxides and the quantitative determination of H$_2$SiO$_4^-$ adsorption in controlled conditions of temperature, solid:liquid ratio, pH and ionic strength.

2. MATERIALS AND METHODS

At all steps, acid-washed high density polyethylene (HDPE) ware and analytical grade Pro Analysis chemicals were used to minimize Si contamination.

2.1. Synthesis and characterization of ferrihydrite and goethite

Ferrihydrite and goethite were synthesized according to the methods described in Cornell and Schwertmann (1996). For ferrihydrite, a freshly prepared 0.1 M Fe(NO$_3$)$_3$ solution (250 ml) was slowly neutralized by 1 M NaOH (70 ml), i.e. brought drop wise to pH 7.5. The red/brown precipitate was separated by centrifugation (3500 rpm, 10 min), dialyzed (dialysis-membrane SPECTRA/POR 4, MWCO = 12–14000, Φ = 29 mm) against deionized water until electrical conductivity was stable at 1–2 µS for 4 h. The dialyzed product was freeze-dried. For goethite, a freshly prepared 1 M Fe(NO$_3$)$_3$ (100 ml) solution was neutralized by 5 M NaOH (70 ml) under intense stirring and brought to pH > 12. The red/brown precipitate was dried in the mother liquid at 70 °C for 60 h, and turned to yellow/brown. The precipitate was washed with deionized water through washing-centrifugation cycles (13,100 rpm, 15 min) until pH and electrical conductivity were stable, respectively, at 5.5 and 1–2 µS. The precipitate was oven-dried at 50 °C for 48 h. Mineralogical and chemical characterizations of both products were done by (1) X-ray diffraction (XRD, Bruker D8 Advance diffractometer), (2) transmission electron microscopy (TEM, Philips 420 STEM), (3) elemental analysis (inductively coupled plasma/atomic emission spectrometry: ICP–AES, Jarrell Ash Iris Advantage) after Na$_2$O$_2$ fusion in vitrified graphite crucibles at 1000 °C, (4) dithionite–citrate–bicarbonate extraction (DCB, Mehra and Jackson, 1965), (5) dark oxalate extraction (Blakemore et al., 1981), and (6) surface area determination using ethylene glycol monoethylether (EGME, Carter et al., 1965).

The X-ray diffraction (XRD) patterns were consistent with those for 2-line ferrihydrite and goethite in Cornell and Schwertmann (1996), and in Jambor and Dutrizac (1998) (Fig. 1). Observations by TEM revealed the common microaggregated shape of ferrihydrite, and the typical euhe- deral acicular crystals (0.5–1.5 mm length) of goethite, constituted of parallel subunits (Fig. 2) (Schwertmann and Taylor, 1989; Cornell and Schwertmann, 1996). The content of poorly crystalline Fe was assessed through the determination of the ratio of oxalate extractable Fe (Fe$_{ox}$) to DCB extractable Fe (Fe$_{dc}$) (Cornell and Schwertmann, 1996) (Table 1). The Fe$_{ox}$/Fe$_{dc}$ ratio was 0.004 for goethite, certifying a well crystalline form, and 0.864 for ferrihydrite, attesting a large dominance of sro mineral particles. The average concentration of Si in the synthesized ferrihydrite and goethite was below 0.2 g kg$^{-1}$ (Table 1), revealing the very low level of Si contamination. A specific extraction of initial adsorbed (i.e. potentially contaminating) Si by KH$_2$PO$_4$ (Delfosse et al., 2005) showed that this Si background was negligible and represents less than 0.5% of the total Si budget per experiment (Table 1). The EGME specific surface was 338 and 147 m$^2$ g$^{-1}$ for ferrihydrite and goethite, respectively.

2.2. Adsorption experiments

H$_2$SiO$_4^-$ solutions were prepared by dissolving Na$_2$SiO$_3$·5H$_2$O in MilliQ water, and further leaching on an H$^+$ cation exchanger (Amberlite® IR-120) to fix Na$^+$ ions. The leaching was terminated when the threshold level of sodium was below 10$^{-2}$ mM Na, as determined by ICP–AES (Henriët et al., 2006). The acidic Si stock solution
(35.61 mM Si, pH below 6) was free of polymers (Beckwith and Reeve, 1963; Stumm and Morgan, 1996). Eight solutions of distinct initial concentrations (ic) of Si were considered: 0.21, 0.42, 0.64, 0.85, 1.06, 1.29, 1.49 and 1.80 mM Si. The ic solutions were prepared by using MilliQ water and pure NaNO$_3$. A background electrolyte concentration of 1 mM NaNO$_3$ was used throughout to maintain ionic strength constant. Contact between ic Si solution and Fe oxide was made in HDPE bottles, on a reciprocating shaker in a dark room at 20°C. A suspension of 5 g of Fe oxide was transferred to a dialysis-membrane (SPECTRA/POR 4; MWCO 12–14,000; Ø = 16mm) and plunged in the Si solution of given ic. The initial solution volume was 1000 ml, as gravimetrically determined (1 g:250 ml solid:liquid ratio). The pH was adjusted to the target pH 5.5 ± 0.2 by addition of 1 M NaOH or 0.7 M HNO$_3$. The target pH value is common for natural waters in equilib-rium with atmospheric CO$_2$. The adsorption experiments were done in triplicates. Series 1 and 2 provided solution aliquots to determine the bulk concentrations of Si determined by ICP–AES (Jarrell Ash Iris Advantage, detection limit <0.7 μM Si). Adsorbed Si was computed as the difference between the Si solution concentrations before and after contact with Fe oxide. Series 3 was specifically devoted to the determination of Si isotopes. Solution aliquots (10 ml) were sampled, respectively, after 6, 12, 24, 48, 72, 96, 192, 288, 408 and 504 h of contact time between Fe oxide and Si solution, and kept in HDPE bottles. The aliquots from each solution were transferred to polyethylene scintillation vials, then acidified by adding 50 μl of 7 M HNO$_3$ and stored in the dark at 4°C prior to further analysis. During the experiment, pH was regularly checked and adjusted to the target when necessary, and systematically checked on a single series one hour prior to sampling solutions.

2.3. Isotopic composition of solutions at given contact times

Two ic solutions (0.64 and 1.06 mM Si) and 20 solution aliquots from series 3 were selected on the basis of adsorption data with respect to detection limit required for Si isotope measurement. Dissolved Si in the chosen samples was purified by triethylamine molybdate co-precipitation and...
combustion in covered Pt crucibles at 1000 °C (De La Ro-cha et al., 1996), and dissolved in dilute suprapur HF–HCl mixture (Cardinal et al., 2003). Si isotopes were measured using a Nu Plasma MC–ICP–MS operating in dry plasma mode, with an external Mg doping to correct mass bias (Cardinal et al., 2003). Due to unresolved isobaric interference on 30Si, only 29Si and 28Si isotopes were measured accurately (Cardinal et al., 2003). The analytical method was supported by an inter-laboratory comparison exercise and proved to be accurate on secondary reference materials (Reynolds et al., 2007). The average precision on δ29Si was ±0.08 (±2σSEM). The results are presented as δ29Si (‰), expressing the 29Si/28Si ratios of our samples relative to the NBS28 silica sand standard (National Institute of Standard and Technology Reference Material #8546) for silicon isotopes (Carignan et al., 2004):

δ29Si(‰) = \left( \frac{\delta_{\text{samples}}}{\delta_{\text{NBS28}}} \right)_{\text{Si}} - 1 \times 1000  

Very small concentrations (below 0.5% of the total Si budget per experiment) of initial contaminating Si from mineral synthesis should not affect the isotopic composition of our solutions. Indeed, a significant contamination of ±0.08‰ (±2σSEM) should be only induced by such contaminating Si bearing an isotopic composition lighter than −48‰ or heavier than +49‰, which is very unlikely regarding Si-isotopic variations on Earth (Basile-Doelsch, 2006).

3. RESULTS AND DISCUSSION

Mineralogical and chemical characterizations show that the separate synthesized products were genuine ferrihydrite and goethite minerals.

3.1. Monomeric Si adsorption by ferrihydrite and goethite

Only the average values of the duplicates are presented in following text, tables and figures. The error bars are systematically presented in the figures, and represent the experimental reproducibility. Table 2 presents the average bulk concentrations of Si in each solution corresponding to ferrihydrite and goethite, and to the respective eight ic solutions and 10 contact times. For both Fe oxides and each ic solution, the concentration of aqueous Si readily decreases with increasing contact time, revealing net Si adsorption. Below pH 9, the solution speciation of H₄SiO₄⁰ is pH independent, meaning that only aqueous H₄SiO₄⁰ was present in significant concentrations (Hiemstra et al., 2007). The systematic adjustment at pH 5.5 ± 0.2 required a larger addition of NaOH for ferrihydrite, indicating a net proton release during adsorption; the addition of NaOH was very low for goethite. In our controlled pH conditions, the H⁺ release is likely caused by the specific interaction of H₄SiO₄⁰ with OH groups of oxide surface involving ligand exchange under formation of a Fe oxide-monosilicate surface complex (Fe₂O₂Si(OH)₂ bi-dendate complex) (Sigg and Stumm, 1981; Hansen et al., 1994a; Hiemstra et al., 2007). At Si
concentration above \( \approx 0.2 \text{ mM} \), a surface Si tetramer \( \text{Fe}_2\text{O}_3\text{SiOHOSi}_3\text{O}_3(\text{OH})_9 \) may have formed on goethite surface; at pH 5.5, the proportion of this surface tetramer would be below \( \approx 25\% \), and the monomer species should be largely dominant (Hiemstra et al., 2007). The maximum value of the mole ratio of aqueous Si to ferrihydrite-Fe was 0.03. In these conditions, the only significant surface bonding of \( \text{H}_4\text{SiO}_4^0 \) with ferrihydrite surface should be the surface complexation of monomeric \( \text{H}_4\text{SiO}_4^0 \) (Swedlund and Webster, 1999). This interpretation is consistent with the adsorption data performed at pH 3–6 by Hansen et al. (1994a).

3.2. Quantitative Si adsorption by ferrihydrite and goethite

In fixed conditions of pH, ionic strength, temperature and solid:liquid ratio, the Si amount adsorbed depends on reaction time and type of Fe oxide (Hansen et al., 1994a). Per oxide mass, ferrihydrite generally adsorbs more Si than goethite (Table 2). After 504 h of contact between Fe oxide and Si solution, the fraction of adsorbed Si ranges between 63% and 86% for ferrihydrite, and between 37% and 72% for goethite (Fig. 3). Few studies have shown a larger Si adsorption by ferrihydrite over goethite (Hansen et al., 1994a), or by amorphous over crystalline Fe oxide (Jones and Handreck, 1963). Surface reactivity for oxyanions, weak acids and water is well known to decrease with increasing Fe oxide crystallinity (Parfitt, 1978; Schwertmann et al., 1985; Schwertmann and Taylor, 1989; Cornell and Schwertmann, 1996). As crystallinity increases, oxide crystals become larger and surface area decreases (Schwertmann et al., 1985). As measured by EGME retention, the surface area was 338 m\(^2\) g\(^{-1}\) for ferrihydrite and 147 m\(^2\) g\(^{-1}\) for goethite (Section 2.1). As expected (Cornell and Schwertmann, 1996), these EGME values are generally above BET-N\(_2\) values previously measured for synthesized Fe oxides: 35–87 m\(^2\) g\(^{-1}\) for goethite (Hansen et al., 1994a; Waltham and Eick, 2002; Garman et al., 2004; Luxton et al., 2006), 269–380 m\(^2\) g\(^{-1}\) for ferrihydrite (Hansen et al., 1994a,b; Hofmann et al., 2004). In agreement with Cornell and Schwertmann (1996), we believe, however, that there is large uncertainty about the surface area measurement of hydrous hydroxyl-bearing Fe oxide because this measurement requires a prior anhydrous vacuum. This conditioning can, indeed, modify surface particle, particle size and porosity through particle microaggregation (Hofmann et al., 2004).

The adsorption data are illustrated at 504 h contact time in Fig. 4. The data could be fitted to Freundlich, Langmuir, Temkin and Redlich–Peterson adsorption isotherms (not shown). For goethite, the adsorption data \( (Q, C) \) best fit a Freundlich isotherm expressed as:

\[
Q = K_F C^{1/n}
\]

where \( Q \) is the Si adsorbed per unit mass of oxide (mmol Si g\(^{-1}\)), \( K_F \) and \( n \) are empirical constants and \( C \) the \( \text{H}_4\text{SiO}_4^0 \) concentration of the solution (mM Si). This empirical model has been considered to be thermodynamically consistent with sorption on heterogeneous surfaces that imply different sorption sites and affinities (Weber et al., 1991). However, this is also consistent with the fact that the positive charge of Fe oxide surface decreases with the increase in the surface Si loading (Hingston et al., 1972; Anderson and Benjamin, 1985; Hiemstra et al., 2007), since the net proton release leads to a shift in the isoelectric point of Fe oxide (Garman et al., 2004; Luxton et al., 2006). For ferrihydrite, the adsorption data best fit a Temkin isotherm expressed as

\[
Q = A + B \ln C
\]

\( n = 2 \) of adsorbed Si, expressed as a fraction of initial Si after 504 h of contact. Ferrihydrite: open triangle. Goethite: full circle. Plotted error bar corresponds to standard deviation between series 1 and 2.

Fig. 3. Average values (n = 2) of adsorbed Si, expressed as a fraction of initial Si after 504 h of contact. Ferrihydrite: open triangle. Goethite: full circle. Plotted error bar corresponds to standard deviation between series 1 and 2.
where Q and C are defined as in Eq. (2), and A and B are constants determined through linear regression from experimental data. For both ferrihydrite and goethite, the Temkin equation implies the condition that the energy of adsorption decreases linearly with surface coverage (Parfitt, 1978). This is consistent with the decrease of charge density as proposed for goethite. As discussed above, the much larger adsorption of Si on ferrihydrite in the range 0.1–0.6 mM Si is attributed to the larger availability of reactive sites in ferrihydrite relatively to goethite.

3.3. Adsorption kinetics

Si adsorption increases with time, first rapidly, then slower (Table 2). The time dependent adsorption of Si by Fe oxide is illustrated for the ic solutions 0.64 and 1.06 mM Si (Fig. 5). Si adsorption requires days–weeks. The rate of adsorption is rather large during the first hours (~50 h for goethite, ~100 h for ferrihydrite), then decreases. After ~200 h for goethite and ~300 h for ferrihydrite, it seems to reach a constant value, which increases from goethite to ferrihydrite with increasing aqueous Si concentration. These observations are in very good agreement with previous results (Hansen et al., 1994a,b). The change in the rate of Si adsorption onto ferrihydrite is similar to that observed for phosphate (Lijklema, 1980; Bolan et al., 1985). The slow adsorption step for phosphate has been attributed to diffusion into interparticle pores (Willett et al., 1988), whereas arsenate adsorption onto ferrihydrite has been successfully predicted using a pore-space diffusion model and assuming rapid adsorption at external surfaces of aggregates (Fuller et al., 1993). Accordingly, and in agreement with Hansen et al. (1994b), the rate of Si adsorption observed here is probably controlled by rapid interaction of monosilic acid with external oxide surface sites, and by slower interparticle diffusion.

3.4. Si-isotopic fractionation during adsorption of monosilic acid

The selected solutions correspond to the ic solutions 0.64 and 1.06 mM Si at various contact times ranging, respectively, between 0 and 192 h, and 0 and 504 h. The values of Si concentration in the selected solutions at pH 5.5 range between 0.25 and 0.96 mM Si (Table 2). These conditions of pH and Si concentration imply a dominant adsorption of monomeric Si (Sigg and Stumm, 1981; Hansen et al., 1994a,b; Swedlund and Webster, 1999; Hiemstra et al., 2007), but do not exclude the occurrence of a surface Si tetramer on goethite surface. For Si solutions oversaturated with respect to quartz ($K_S > 0.12$ mM Si), the proportion of surface Si tetramer was estimated below 15% for Si concentration below 1 mM Si (Hiemstra et al., 2007).

The Si-isotopic compositions of the selected solutions are given in Table 3. Generally the $\delta^{29}$Si value significantly increases with increasing time, and thus with decreasing Si concentration of the solution at various contact times. This increase involves a gradual fractionation of stable Si isotopes, revealing a depletion of the aqueous phase in light Si isotopes which parallels the adsorption of monosilic acid onto Fe oxide. Measured against its initial composition (+0.01 ± 0.04‰ (+2σ)), the solution is indeed systemically enriched with the heavy isotope, reaching a maximum $\delta^{29}$Si value of +0.70 ± 0.07‰ for ferrihydrite and +0.50 ± 0.08‰ for goethite at ic 1.06 mM Si after 504 h of contact. For ferrihydrite at ic 0.64 mM Si, solution at 72 h displays lighter Si-isotopic compositions than solution at 48 h, unlike the general increase of $\delta^{29}$Si in solution with time in other treatments (Table 3). This shift down might be attributed to a possible abnormal variation of pH during the experiment at 72 h in this series (series 3 was used exclusively for isotopic measurements and therefore pH was not
The isotopic composition of the solution can be predicted not be considered in further calculations and interpretation. (Jones and Handreck, 1963). The two measurements at 72 and 192 h will thus adsorption of Si onto Fe oxide (Jones and Handreck, 1963). The two measurements at 72 and 192 h will thus not be considered in further calculations and interpretation. The isotopic composition of the solution can be predicted by following models, generally used to describe biologically driven Si-isotopic fractionation:

The Rayleigh model following:

\[ \delta^{29}\text{Si}_{\text{solution}} = \delta^{29}\text{Si}_{\text{initial}} + \delta^{29}\epsilon_{R} \ln f \]  
(4)

The steady state model following:

\[ \delta^{29}\text{Si}_{\text{solution}} = \delta^{29}\text{Si}_{\text{initial}} - \delta^{29}\epsilon_{S}(1 - f) \]  
(5)

where \( \delta^{29}\text{Si}_{\text{initial}} \) and \( \delta^{29}\text{Si}_{\text{solution}} \) are, respectively, measured in the ic solution and the solution at each contact time (Table 3), \( f \) is the fraction of Si remaining in solution at each contact time (Table 2), and \( \epsilon_{R} \) is the fractionation factor (R for Rayleigh, S for steady state).

In order to decipher between the two models, best fits curves have been estimated based on three isotopic measurements of the beginning of each time series experiments (\( f = 1, \sim 0.8 \) and \( \sim 0.7 \)) where both models are still not distinguishable compared to the analytical standard error (calculated \( \epsilon_{c} \) in Table 4A). The experimental data were plotted on those best fit curves (Fig. 6). This indicated a preferential Rayleigh process rather than a steady state model for ferrihydrite at 1.06 mM Si (Fig. 6b), and also though in a less extent a Rayleigh process for goethite at both 0.64 and 1.06 mM Si (Fig. 6c and d).

To further reduce the uncertainty on the fractionation factor \( \epsilon_{c} \), more estimates have then been calculated from Rayleigh Eq. (4) using all experimental data available (Table 4B). The errors associated to the use of Eq. (4) to derive a fractionation factor are discussed in Scott et al. (2004). In the case of our \( \delta^{29}\text{Si} \) values, these errors are however not significant due to analytical limitations. Therefore, the
uncertainty on $29\delta_e$ is calculated as the error on the slope ($1\sigma_{SD}$) of $\delta^{29}\text{Si}_{\text{solution}}$ data plotted against $\ln f$ for Rayleigh including propagated analytical uncertainty. It was calculated using the REG procedure of the SAS System (version 9.1 for Windows, SAS Institute, Cary, NC, USA). Following the Rayleigh model, $29\delta_{eR}$ ($\pm 1\sigma_{SD}$) ranges between $-0.54 \pm 0.03\%_{oo}$ and $-0.55 \pm 0.02\%_{oo}$ for ferrihydrite, and between $-0.84 \pm 0.13\%_{oo}$ and $-0.78 \pm 0.11\%_{oo}$ ($\pm 1\sigma_{SD}$) for goethite, for ic 0.64 and 1.06 mM, respectively (Table 4B). The fractionation factor $29\delta_{eR}$ does not differ between ic solutions for a given oxide, but is significantly larger for goethite than for ferrihydrite. Estimates from the steady state model are also given in Table 4B (with uncertainty as the error on the slope of $\delta^{29}\text{Si}_{\text{solution}}$ data plotted against $(1-f)$), from which we deduce that (i) the uncertainty on the Rayleigh $29\delta_{eR}$ fractionation factor is much smaller than the one reckoned for a steady state fractionation ($29\delta_{eS}$), confirming our deduction drawn from the first calculation, (ii) change of ic does not affect $29\delta_{eR}$ neither for ferrihydrite nor for goethite.

Averaging our two groups of experiments at both ic solutions, our best estimates of $29\delta_{eR}$ ($\pm 1\sigma_{SD}$) are computed...
Si-isotopic fractionation by adsorption onto Fe oxide

4. IMPLICATIONS

4.1. Mechanisms of isotopic fractionation

Our $\delta^{29}_{\text{Si}}$ values following Rayleigh model (Table 4B) show that the Si-isotopic fractionation induced by $\text{H}_2\text{SiO}_4^-$ adsorption onto Fe oxide is similar or slightly larger to the one generated by biological processes such as Si uptake by plants ($\delta^{29}_{\text{Si}} = -0.53 \pm 0.17\%_{\text{wo}}$, Ding et al., 2005; $-0.52 \pm 0.16\%_{\text{wo}}$, Ziegler et al., 2005a; $-0.40 \pm 0.11\%_{\text{wo}}$, Opfergelt et al., 2006a) and diatoms ($\delta^{29}_{\text{Si}} = -0.57 \pm 0.21\%_{\text{wo}}$, De La Rocha et al., 1997).

According to Barling and Anbar (2004), the adequacy of the Rayleigh fit within our type of experimental design would indicate an irreversible adsorption process, which would suggest a kinetic isotope effect, and thus could be associated with (i) the variation of Si adsorption rate (Fig. 4 and Table 3), and (ii) the poor reversibility of Si adsorption through ligand exchange (Parfitt, 1978). However, we have no implemented an experimental design to test the irreversibility. Yet, in our scheme, isotopic fractionation can result from (i) an equilibrium fractionation between coexisting aqueous species coupled with a selective sorption of one of them (Siebert et al., 2003), (ii) the formation of inner sphere surface complexes (Lemarchand et al., 2007). In this respect, light boron isotopic enrichment on goethite is strongly dependent on pH and surface complex structure (Lemarchand et al., 2007). Here, the equilibrium fractionation process between coexisting aqueous species could be discarded because $\text{H}_2\text{SiO}_4^-$ is the only significant aqueous Si species in our experimental conditions (see Section 3.1). The Ge isotopic fractionation during Ge sorption onto goethite privileges the selective sorption of light Ge isotopes, and has been successfully modeled by surface complexation involving the interaction of monomeric Ge hydro-complexes with $>\text{FeOH}^\text{2+}$ and $>\text{FeO}^\text{2-}$ sites of goethite (Galy et al., 2002). These complexes have been further experimentally identified as Ge bi-dendate surface complexes composed of tetrahedrally coordinated Ge attached to the corners of two adjacent Fe octahedra (Pokrovsky et al., 2006). Here, we tentatively propose that the Si-isotopic fractionation induced by $\text{H}_2\text{SiO}_4^-$ sorption onto ferrhydrite and goethite is caused by the formation of Fe oxide-monosilicate bi-dendate inner surface complexes ($\equiv\text{Fe}_2\text{O}_3\text{Si(OH)}_3^\text{2-}$) (Sigg and Stumm, 1981; Hansen et al., 1994a; Dietzel, 2002; Hiemstra et al., 2007). However, surface Si polymerization, according to the definition of McBride (1994), cannot be disregarded, as the occurrence of Si tetramer has been predicted on goethite surface for Si solutions oversaturated with respect to quartz ($\sim0.12\text{ mM Si}$) (Hiemstra et al., 2007) (see Section 3.1). This would hypothetically contribute to increase Si isotope fractionation for goethite relatively to ferrhydrite, for which the only significant surface bonding of $\text{H}_2\text{SiO}_4^-$ should be the surface complexation of monomeric $\text{H}_2\text{SiO}_4^-$ (Hansen et al., 1994a,b; Swedlund and Webster, 1999). Such a hypothesis would fit with the enrichment of light Si isotopes observed in the clay-sized fraction of soils with increasing weathering (Ziegler et al., 2005a,b; Opfergelt et al., accepted for publication) suggesting that clay formation may privilege light Si isotopes. Indeed, such formation in soil environment requires Si polymerization at low pressure and temperature. These hypotheses need, however, both further in-depth field-based investigations and a theoretical evaluation of the size of the isotopic energy shifts between soluble $\text{H}_2\text{SiO}_4^-$ and potential Si adsorbed polymers.

We cannot rule out that desorption occurred during our experiment. Since the experimental device was not designed to assess the impact of Si desorption on Si-isotopic fractionation it would be hazardous to extrapolate our results to the desorption process. We believe that the impact of desorption would be minor on our isotopic data because (i) the adsorption curves showed very regular patterns in accordance with adsorption as the main process driving silicon content in the solution (Fig. 5) (Hansen et al., 1994a,b); (ii) most of the data were acquired for $f > 0.5$ which should favor adsorption over desorption before saturation of sites is reached (Hansen et al., 1994a,b); (iii) to impact significantly on our results, desorption should fractionate isotopes at least in the same extent as adsorption which is unlikely. Unfortunately, to the best of our knowledge, there is so far no experimental setting to study isotopic fractionation induced by the sole Si desorption.

4.2. Environmental significance

Our data provide unequivocal answers to the four questions of critical environmental significance (see Section 1). Si isotope fractionation (1) occurs during Si adsorption onto Fe oxide at common pH for soil solutions, (2) is similar or larger to the one generated by Si uptake by biota, and can thus contribute (3) to the depletion of light Si isotopes in river waters, and (4) to the relative concentration of light Si isotopes in soil clay fractions through Si sorption on pedogenic Fe oxides. Our data further suggest that the concentration of light Si isotopes in soil clay-sized fractions (Ziegler et al., 2005a,b; Opfergelt et al., accepted for publication) can be at least partly due to $\text{H}_2\text{SiO}_4^-$ sorption onto secondary iron oxides.

Silicon adsorption by soil Fe oxides is thus a process which may partly control the depletion of light Si isotope in river waters, in addition to clay formation (Ziegler et al., 2005b; Georg et al., 2006, 2007), Si uptake by plants (Ding et al., 2005; Opfergelt et al., 2006a) and diatoms (Allman et al., 2005). However, impact of Si adsorption on the Si-isotopic budget will be strongly limited if oxide surfaces are Si-saturated as shown experimentally with $^{32}$Si (Ziegler et al., 2005a). Ferrhydrite and goethite are sparingly soluble constituents of weathered rocks and soils, with solubility products in the range $10^{-38}$ to $10^{-46}$ M (Schwertmann and Taylor, 1989). In well drained conditions, Fe released from weathered parent material is thus poorly mobile and accumulates in soils as secondary Fe(III) minerals: Fe oxides rapidly precipitate as discrete solid phases from the weathering solution of decomposed primary silicates. As inferred from studies on the adsorption of dissolved organic matter, juvenile oxide surfaces are very effective to adsorb...
solutes with which they specifically interact (Guggenberger and Kaiser, 2003). Rock weathering and soil development lead to the formation of both clay minerals and secondary oxides. These processes may thus add their mutual isotopic effects to significantly impact the Si-isotopic signature of drained waters. Weathered soil and saprolite layers thus acquired a high propensity to impact soil solutions and exported waters through Si adsorption. As an example, Si adsorption could have impacted isotopic signatures of Siberian rivers draining iron-rich swamp zones and peat soils of the permafrost landscape, which are found to be strongly enriched in heavy Si isotopes compared to basaltic bedrock (Reynolds et al., 2006).

Si stable isotopes thus constitute a promising tracer with respect to the processes involved in the weathering environment: biological fractionation during plant phytolith formation (Ding et al., 2005, 2008; Opfergelt et al., 2006a), sequestration of Si in soil clay-sized minerals (Ziegler et al., 2006a; Opfergelt et al., accepted for publication), and adsorption of Si by pedogenic iron oxyhydroxides (this study). A similar statement concerns the Ge/Si ratio as a weathering tracer (Mortlock and Froelich, 1987; Scribner et al., 2006; Derry et al., 2005). The impact of weathering stage, soil development and free iron oxide availability on the Si-isotopic composition of source and river waters thus deserves further field-based studies to progress in the appraisal of the silicon continental cycle.

5. CONCLUSION

The adsorption of H$_2$SiO$_4$\(^+\) by ferrihydrite and goethite at pH 5.5 strongly fractionates Si isotopes by selectively adsorbing light isotopes and leaving a companion solution enriched with heavy Si isotopes. The isotopic fractionation of silicon is similar to or slightly larger than that generated by Si uptake by plants and diatoms. We suggest that the concentration of light Si isotopes in soil clay-sized fractions is at least partly due to H$_2$SiO$_4$\(^+\) sorption onto secondary iron oxides. We conclude that rock weathering and soil development could impact the Si-isotopic signature of natural waters drained to streams through Fe oxide synthesis, as oxide surfaces specifically interact with aqueous monosilicic acid. Although occurring at very different time scales (from the rainy event to the soil formation), the processes of pore water drainage, clay-sized Fe oxide formation and Si adsorption would significantly impact the Si-isotopic signal to oceans, as continental runoff water contributes to more than 80% of the Si input to the marine Si budget (Tréguer et al., 1995).

In the future, the Si-isotopic fractionation induced by adsorption onto Al-oxides and in natural soils should also be investigated along with the impact of desorption.

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