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Processes controlling silicon isotopic fractionation in a forested tropical watershed: Mule Hole Critical Zone Observatory (Southern India)

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Abstract

Assessing the dynamics of the silica cycle in the critical zone remains challenging, particularly within the soil, where multiple processes are involved. To improve our understanding of this cycle in the Tropics, and more specifically the role played by vegetation, we combined elemental Si mass balance with the \(\delta^{30}\)Si signatures of the compartments involved in the water-plant-rock interactions of a tropical forested watershed, Mule Hole (Southern India). To accomplish this, we analysed (1) the \(\delta^{30}\)Si values of present-day litter phytoliths from tree leaves and grass, as well as soil amorphous silica (ASi); (2) the Si isotope fractionation induced by phytolith dissolution; (3) the silicon mass balance inferred from isotopes at the soil-plant scale; and (4) the consistency between water sources and the \(\delta^{30}\)Si signatures in the ephemeral stream.

The \(\delta^{30}\)Si values of present-day litter phytoliths and soil ASi vary within a narrow range of 1.10 to 1.40 ‰ for all samples, but two deep vertisol samples which likely trapped phytoliths from different vegetation growing under more humid conditions, as indicated by pollen analysis. A homogeneous signature of litter is a minimum condition for using \(\delta^{30}\)Si as a
proxy for the litter/phytolith source of Si. However, litter-ash dissolution experiments demonstrate that the incipient dissolution of phytoliths fractionates Si isotopes, with the preferential dissolution of $^{28}$Si over $^{30}$Si yielding $\delta^{30}$Si values as low as -1.41‰. Values close to the whole-sample signatures, i.e., above 1‰, were recovered in the solution after a few hours of water-ash interaction. At the soil-plant scale, the average $\delta^{30}$Si value of soil-infiltrating solutions is slightly lighter than the average phytolith signature, which suggests phytoliths as the source of soil dissolved Si. The isotopic budget of dissolved Si within the soil layer, which was obtained based on previous elemental fluxes, is imbalanced. Equilibrating the isotopic budget would imply that up to 4100 mol ha$^{-1}$ yr$^{-1}$ of silica is taken up by vegetation, which is almost twice as large as that initially estimated from the elemental budget. The additional Si flux taken up, and likely stored in woody stems, was estimated assuming that Si isotopes followed a steady-state model for the whole Si plant uptake and then followed a Rayleigh model once in the plants. The $\delta^{30}$Si value of the additional Si flux taken up should be close to 0‰, i.e., enriched in light Si isotopes compared to the litter. If steady-state conditions apply, the source could correspond to soil ASi dissolution or deep (saprolite) root uptake. At the outlet of the watershed, the stream exhibits low $\delta^{30}$Si values (0.28 to 0.71‰) during peak flows and high $\delta^{30}$Si values (1.29 to 1.61‰) during the recessions at the end of the rainy season. Heavy $\delta^{30}$Si signatures are consistent with the expected domination of seepage at the end of floods. The light $\delta^{30}$Si values during peak flow are slightly lower than the overland flow signature and reflect either a sampling bias of overland flow or a minor but significant contribution of another Si source within the stream, possibly the partial dissolution of phytoliths from the suspended load, with slight isotopic fractionation. This study confirms that vegetation controls the silicon cycle in this dry tropical forest. It also shows that silicon isotopes yield a better grasp of the mass balance and sources and potential mechanisms involved than the consideration of only silicon concentrations. However, this proxy still relies on working hypotheses, notably steady-state and/or Rayleigh fractionation models, which need to be confirmed in further studies.

1 Introduction

The biological cycle of silicon has received increasing attention in the last decade because it mobilizes large amounts of silica, particularly in the Tropics, where rivers contribute 74% of the dissolved flux to the ocean (Martin & Meybec, 1979; Tréguer and De La Rocha, 2013).
For example, in the Amazonian forest, the silica flux brought to the soil by litterfall (as phytoliths) is four times more important than the silica flux exported by the river (Lucas et al., 1993). Similarly, at Reunion Island, Meunier et al. (2001) found that the silica annually recycled by litter was similar to the chemical weathering flux estimated by Louvat & Allègre (1997). The high solubilities of phytoliths and other amorphous silica particles in soils make them potentially significant contributors to the solute silica budget within the soil (Alexandre et al., 1997; Fraysse et al., 2010; Cornelis and Delvaux, 2016) and possibly to rivers. The biological control of silica fluxes in river systems can be observed in some streams, such as in Hawaii, using the Ge/Si ratio (Derry et al., 2005); more recently, this was observed in the experimental watershed of Mule Hole (Southern India) using combined solute mass balance at the soil-plant scale and hydrological modelling (Riotte et al., 2014a).

Because silicon isotopes are fractionated during interactions between water, soil and plants, the isotopic signature ($\delta^{30}$Si values) of these compartments has the potential to trace the origin of the silica fluxes exported from watersheds (e.g., Cardinal et al., 2010; Opfergelt and Delmelle, 2012; Hughes et al., 2013). The fractionation of silicon isotopes is significant in natural environments, particularly in the various compartments of the Critical Zone (i.e., Earth’s surface zone, encompassing groundwater, soils, rivers and vegetation), with values up to 12‰ (see reviews in Opfergelt and Delmelle, 2012, Frings et al., 2016). The lightest Si isotopic compositions are found in the silicification zones of aquifers ($\delta^{30}$Si = -5.7‰, Basile-Doelsch et al., 2005), while the heaviest ones are observed in some vegetation components ($\delta^{30}$Si = +6.1‰, Ding et al., 2005).

The use of $\delta^{30}$Si as a tracer of biogeochemical cycling in watersheds is complex. Both the formation of clay minerals during weathering and biological uptake result in the preferential incorporation of the light $^{28}$Si isotope relative to the heavier $^{29}$Si and $^{30}$Si isotopes (see review in Poitrasson, 2017). The silicon isotope fractionation factors (expressed as $\Delta^{30}$Si rather than using the $^{30}$Si notation, see Poitrasson, 2017) between parent and authigenic minerals are therefore negative. They range from -1‰ for the adsorption of Si onto ferrihydrite (Delstanche et al., 2009) to -1.5 and -2‰ for the incorporation of Si into allophane and kaolinite (Ziegler et al., 2005; Georg et al., 2007). Some of the mechanisms controlling this fractionation factor during water-mineral interactions were recently identified through laboratory experiments. During the abiotic precipitation of amorphous silica (further referred to as ASi), Geilert et al. (2014) observed that the isotopic fractionation between precipitated and dissolved Si decreased with increasing temperature from -2.1‰ at 10°C to almost 0‰
beyond 50°C. Geilert et al. (2014) suggested that the effect of temperature on isotopic fractionation is “system-dependent”, i.e., indirect and primarily linked to the saturation level of the solution, changes in reactive surface area and the flow regime. Although the silica saturation levels of soil pore water can be easily estimated in natural systems, assessing soil reactive surface areas and flow regimes remains challenging.

The use of $\delta^{30}$Si as a tracer for studying the silica cycle in vegetation is also not straightforward for two reasons. First, the range of $\delta^{30}$Si values in worldwide plants, as well as in the phytoliths they produce, exceeds 8‰ (Frings et al., 2016), which means that cycling through plants does not induce a specific isotopic signature. However, if the $\delta^{30}$Si signatures of all plants in a given ecosystem are “system-dependent”, they should vary within a narrow range, as observed in a boreal forest (Engström et al. 2008). In this case, $\delta^{30}$Si signatures could be used to trace the contribution of vegetation to the silica cycle in ecosystems and river systems, providing that the “vegetation signature” is distinct from the signature induced by silicate weathering (i.e., the dissolution of primary minerals and the precipitation of clay minerals) or adsorption/desorption processes. Second, significant isotopic fractionation also occurs within plant compartments, with large inter-specific variations. For instance, Ding et al. (2005) reported a large increase of $\delta^{30}$Si within rice plants, from -1‰ in roots to +6‰ in grain, while Opfergelt et al. (2006) observed a limited increase of 0.26‰ from pseudostems to leaves in banana plants. According to these authors, the magnitude of isotopic fractionation within a plant is related to the intensity of evapotranspiration, which, to some extent, is also a “system-dependent” parameter.

In the Mule Hole watershed (which has been monitored as a Critical Zone Observatory (CZO) since 2003), South India, the prominent role of vegetation in the water cycle and the biogeochemical cycle has previously been demonstrated (Ruiz et al., 2010; Riotte et al., 2014a, b). For instance, evapotranspiration accounts for at least 80% of the water balance and controls the decennial fluctuation of the water stock in the vadose zone, as well as runoff and groundwater discharge (Maréchal et al., 2009; Ruiz et al., 2010). In addition, the silica flux annually recycled through the litter compartment represents 10 to 15 times the dissolved silica flux exported by the stream and dominates the solute mass balance at the soil-plant scale. However, assessing the respective fluxes of plant uptake and silicate weathering within the soil layer requires more constraints than solute mass balance at the soil-plant scale.

By taking advantage of the decennial hydro-geochemical monitoring of the Mule Hole watershed and subsequent recent elemental budgeting (Riotte et al., 2014a, b), the objective of
the present study is to describe the processes affecting silica during water-soil-plant interactions and the impact of these processes on Si mass balance using the silicon isotopic signatures ($\delta^{30}$Si) of the dissolved and solid water-soil-plant compartments. This study also represents an opportunity to assess the potential and limitations of $\delta^{30}$Si as a tracer of silica sources at both the soil-plant and watershed scales. First, the morphotypes and isotopic compositions of amorphous silica particles (ASi) (i.e., phytoliths) are characterized and compared with those of fresh litter, soils and stream suspended loads. Second, the $\delta^{30}$Si signature of ASi dissolution is assessed from environmental samples (i.e., overland flow water, which gathers water from canopy and forest floor interactions) and batch leaching experiments. Third, solute isotopic budgets are carried out at the soil-plant and watershed scales to assess the identified processes and sources controlling the silica cycle in this forested watershed, including the hourly and seasonal $\delta^{30}$Si fluctuations in the stream.

2 Environmental settings

The experimental watershed of Mule Hole, which has an area of 4.1 km$^2$, is located at 11°430N–76°260E within the Cauvery Basin, on the Deccan Plateau, in the subhumid zone of the climatic gradient induced by the Western Ghats. The bedrock is composed of 85% Precambrian peninsular gneiss (metasediments, Shadakshara Swamy et al., 1995) containing quartz, oligoclase, sericite, biotite and chlorite as major minerals. In Mule Hole, the average regolith consists of 15-m-thick immature saprolite and 2-m-thick soils (Braun et al., 2009). The soil comprises a ferralsol-vertisol system (Barbiero et al. 2007, 2010). The alternation of subhumid and semi-arid conditions at Mule Hole during at least the Quaternary (Violette et al., 2010a) limited the intensity of chemical weathering and allowed the preservation of few primary minerals, such as Na-plagioclase and sericite, in soils. The secondary mineral assemblages in the ferralsol profile comprise kaolinite and Fe-oxyhydroxides, and those in the vertisol areas comprise a mix of kaolinite, smectite, Fe-oxyhydroxides and relics of pedogenic carbonates (Barbiero et al., 2010, Violette et al., 2010a). Four associations of soils and bedrocks were found and defined (Fig. 1): (A) ferralsol on gneiss, which is the most abundant; (B) vertisol on amphibolite; (C) shallow ferralsol on gneiss; and (D) vertisol on gneiss in valley bottoms.

The annual rainfall at Mule Hole has varied from 800 to 1500 mm/yr over the last 35 years, with an average value of 1100 mm/yr. The hydrological modelling of the watershed (Violette et al., 2010b) and End-Member Mixing Analysis (Riotte et al., 2014a; Audry et al., 2014)
indicate that the stream flow is mostly composed of overland flow (> 90%), i.e., rainwater having interacted with the canopy and then at the ground surface with decaying litter. Soil pore water contributes the last 10% during flood recessions, when soils are saturated at the end of monsoon season.

The vegetation cover is a dry deciduous forest (Barbiero et al., 2007), with the Anogeissus – Tectona - Terminalia (ATT) facies (70% of the watershed area) developed principally on ferralsols and less frequently on vertisols. Shallow ferralsols are also covered by Shorea trees (Shorea Roxb. ex C.F.Gaertn), and vertisols are covered by Ceriscoides turgida (Hook.f.) Tirven. Litter fall occurs within a short period of time after monsoon season, typically from December to February.

The isotopic budgets of the present study rely on the solute mass balances recently performed at two spatial scales in order to understand the biogeochemical processes and their impact on the solute fluxes exported from the watershed: (1) those at the soil-plant profile scale, assessed using the hydrological modelling output for water fluxes and the chemical compositions of rainfall, throughfall, overland flow and soil pore water samples (Riotte et al. 2014a; Braun et al., 2017); and (2) those at the watershed scale, assessed using the hydrological modelling output for water fluxes and the chemical compositions of the stream (Violette et al., 2010b, Maréchal et al. 2011).

3 Materials and methods

3.1 Water, soil, bedrock and vegetation sampling

The water samples analysed in this study for their $\delta^{30}$Si values, namely, stream water, overland flow, soil porewater and groundwater, were analysed during the broader, long-term monitoring of the Mule Hole watershed, which is an environmental observatory (BVET, see https://mtropics.obs-mip.fr/experimental-tropical-watersheds-2/) (Fig. 1). The sampling and storage protocols are described in Riotte et al. (2014a). Stream water was sampled on an hourly basis, and groundwater was sampled on a monthly basis. The stream samples selected for $\delta^{30}$Si measurements (n=11) were chosen in order to cover both hourly and seasonal fluctuations in the stream composition in 2004 and 2005 (see Electronic Annex EA-1). In March 2004, the litter was burnt by a forest fire during the dry season. The two first short storms that occurred a few days later (i.e., in April and May, each with a 3-hour duration) exhibited particularly high silica concentrations (Audry et al., 2014) and were thus selected to
assess the signal of litter ash leaching. One major storm (with a 48-hour duration) that occurred in the middle of the 2004 monsoon season, and is thus more representative of the hydrological functioning of the watershed, was also analysed for its $\delta^{30}$Si composition. In 2005, stream storm samples were collected during the first short storm in the summer season (April) and at the end of the monsoon season, when the soil layer was saturated (November). Overland flow samples (n=7) were collected during the 2009 monsoon season on the ferralsol and vertisol/ferralsol transition (see below and Riotte et al., 2014a for sampling setup); each sample integrated overland flow events for two to three weeks. Deep groundwater samples were collected at 4 locations in 2005: 2 were collected close to a transect crossing the outlet, and 2 others were collected along the crest line (P5 in gneiss, P6 in amphibolite). Soil pore water samples were collected in vertisol seepage within the streambed in 2004 (n=5) and in ferralsol/vertisol transition seepage in 2009 and 2010 (n=4). All water samples were filtrated in the laboratory soon after collection (Sartorius® cellulose acetate, 0.22-µm membrane).

Fig. 1: The Mule Hole watershed with soil and vegetation map, showing the sampling locations of composite soils (plots A, B, C, D; see text for description), soil profiles, overland flow and ground water (modified from Riotte et al., 2014a). ATT (medium green): Anogeissus – Tectona – Terminalia facies. Dark green: swamp facies; light green: Shorea facies.
Soil samples were selected from two well-documented soil profiles, one in ferralsol (5 depths) and one in vertisol (4 depths). Their mineralogy, granulometry, cation exchange capacity and major and trace element contents are summarized in Electronic Annex EA-2 and given in detail in Violette et al. (2010a and 2010b). Composite surface soils are composed of 36 to 42 samples collected between depths of 0 and 10 cm within a one-hectare plot of each soil and bedrock association (Fig. 1): (A) ferralsol on gneiss, (B) vertisol on amphibolite, (C) shallow ferralsol on gneiss and (D) vertisol on gneiss. Approximately 100 g of each soil sample was sieved with water to sizes of 2 mm and 50 µm. The < 2 mm fraction was considered to represent the bulk soil. The < 2 µm fraction was recovered from the < 50 µm fraction after the decantation of the coarse particles according to Stokes’ law. Bulk soils and the < 2 µm fraction were further dried and powdered with an agate mortar. A fresh bedrock sample was collected at a depth of 50 m as drilling cuttings when the “P5” piezometer was drilled. Its chemical composition corresponds to the average gneiss composition estimated by Braun et al. (2009).

Leaf and herbaceous litters were collected in large bags during the dry seasons of 2009 and 2010 from the dominant soil/vegetation configuration in the watershed, i.e., ATT vegetation developed on ferralsol. Litter samples consisted of A. latifolia, T. grandis and T. crenulata leaves and T. triandra grass. The litter of T. grandis was picked up as dry leaves on the ground in January and March 2009 and 2010. The leaves of A. latifolia and T. crenulata, which are more difficult to identify once they have fallen, were harvested on trees during their senescence in January. T. triandra was also cut in January, prior to senescence. All litter samples were dried at 70°C and powdered in an agate mortar (using the protocol detailed in Riotte et al., 2014a). The silica content of A. latifolia leaves was very low, i.e., 0.04 % Si dry weight (DW), which is close to the detection limit (Riotte et al., 2014a). Because this species was unlikely to contribute to silica cycling, δ³⁰Si analyses were restricted to the three species containing measurable silica, i.e., T. triandra grass (1.88 % Si DW), T. grandis (1.97 % Si DW) and T. crenulata (0.4 % Si DW) leaves.

3.2 Assessment of δ³⁰Si during ASi dissolution

Assessing the δ³⁰Si value of ASi dissolution is of primary importance for tracing the origin of silica in rivers and assessing the consistency between elemental and isotopic mass balances. Two approaches were used in this study, namely, one onsite approach and one experimental approach.
The onsite approach relied on the dominant contribution of ASi dissolution to overland flow (Riotte et al., 2014a). Overland flow (n=7) samples were collected close to the streambed at the ground surface using a PVC pipe connected to a buried can during the 2009 and 2010 monsoon seasons. Each collector drained an area of approximately 100 m² and integrated several weeks of intermittent flow. The chemical compositions of overland flow and soil pore water are given in Riotte et al. (2014a). One advantage of using overland flow is that each sample integrates a significant area (~100 m²) and several ephemeral flow events. The limit of this approach is the long interaction, i.e., up to two weeks, between the finest particles (likely phytoliths) and the solution collected in the sampling can, which can prevent the detection of possible kinetic isotopic fractionation occurring during ASi dissolution. The possible contribution of dust was extensively discussed in Riotte et al. (2014). It was considered to be negligible (3% of the Si concentration in overland flow) because the throughfall contained very low Si contents. Also, the ASi origin of Si in overland flow is supported by the good relationship between its Si/Na, DOC/Na, Ca/Na ratios, which all increase during the monsoon season as litter decays, as well as by the oversaturation of overland flow solutions with respect to clays (Riotte et al., 2014a).

The experimental approach, as described in detail in Audry et al. (2014), was performed on litter ash from the ATT vegetation species. It consisted of 2-step batch experiments (with a solid/solution ratio of 2 g L⁻¹, i.e., close to field conditions) and was designed to assess the impact of forest fires on the Mule Hole stream composition. In the first step, a mixture of ash/deionized water was filtered after 5 minutes of interaction to quantify and remove the easily dissolving salts present in the ash that would have dissolved during the first rainfall event. In the second step, the deionized water was renewed with the same solid/solution ratio. The leaching progress was monitored by periodically sampling the suspension without stopping the stirring in order to maintain a constant solid/solution ratio during the full duration of the experiment. The suspension aliquots were filtered, and the filtrates were analysed for their major dissolved species, including silica. The δ³⁰Si signatures were determined for the filtrates retrieved from the first step of the experiment and the filtrates retrieved after 100 hours of interaction for the three species containing measurable silica, i.e., T. triandra grass, T. grandis and T. crenulata leaves. After 100 hours of interaction, the Si concentration in the leachate had reached its maximum value, i.e., 2.83 mmol L⁻¹ for T. grandis and 3.18 mmol L⁻¹ for T. triandra (see Electronic Annex EA-3, data from Audry et al., 2014) beyond saturation with amorphous silica (~2 10³ µmol L⁻¹, log K = -2.71; Parkhurst and Appelo, 1999). The amount of silica released by T. crenulata from the first step was too
small to allow the determination of $\delta^{30}\text{Si}$; thus, for this species, $\delta^{30}\text{Si}$ could only be measured in the filtrate after 100 hours.

3.3 Amorphous Si and Dissolved Si measurements

The ASi pool was separated from the soil (5 g aliquot) and suspended load (0.6 to 3 g, based on sample availability) matrices using heavy liquid flotation. This method allows the quantification and identification of phytolith morphotypes and diatom frustules by optical microscopy (Kelly, 1990). The procedure includes overnight carbonate removal with 37% HCl, the oxidation of organic matter (OM) with 33% H$_2$O$_2$ at 50–70 °C, and clay removal by decantation with sodium hexametaphosphate to ease deflocculation. To remove heavy minerals, we used ZnBr$_2$ heavy liquid ($d=2.3$ g cm$^{-3}$). After observing under the microscope that ASi is mainly composed of phytoliths, we calculated the concentration of Si from ASi/phytoliths assuming a mean water content in opal of 10% (Bartoli & Wilding, 1980), which is equivalent to 0.37 moles of H$_2$O for 2 moles of SiO$_2$ (ASi = 0.42 × mass of phytolith). This is referred herein as ASi$_{zb}$. This extraction protocol may lead to the underestimation of ASi due to phytolith loss during extraction or to the overestimation of ASi when the final fraction is not pure and contains organic compounds or mineralogical residues (Meunier et al., 2014). Because mineralogical residues (typically clays) could impact the estimation of ASi and alter the $\delta^{30}\text{Si}$ signatures of phytoliths, we used a purity index based on the chemical composition of the ASi$_{zb}$ fraction. The chemical composition was determined after the alkaline digestion of ASi with NaOH (see section 3.4) with an ICP-OES Horiba Ultima Expert at GET. The purity of the ASi$_{zb}$ fractions was determined using the relation

$$\text{ASi}_{zb} \text{ purity} = \frac{(\text{Si}-\Sigma(\text{Al, Fe, Ca, Mg, Mn, K}))/\text{Si}}{\text{Si}}$$

where concentrations are expressed in mol g$^{-1}$. Using this index (dimensionless), pure amorphous silica would exhibit an ASi$_{zb}$ purity of 1 and clay would exhibit a value close to 0. For each sample, the mass of ASi/phytoliths that was extracted was expressed as a percentage of the sample dry weight (% DW).

The ASi$_{zb}$ contents were measured along the two vertisol and ferralsol reference profiles, on three suspended loads of streams and overland flow and on four composite 0-20 cm soils corresponding to the four vegetation-soil-bedrock occurrences in the watershed: (A) ferralsol developed on gneiss with ATT vegetation facies (dominant configuration), (B) vertisol developed on amphibolite with ATT vegetation facies, (C) shallow and sandy ferralsol...
developed on gneiss and covered with Shorea and (D) vertisol developed on gneiss and covered with Ceriscoides. The ASi zb data from the 5 vertisols and one ferralsol reported in Table 2 are from Meunier et al. (2014). The dissolved Si concentrations (DSi) of stream, overland flow, soil pore water and groundwater samples were determined with a Hach® spectrophotometer at the Indo-French Cell for Water Sciences (IISc, Bangalore) using the molybdate blue method, with a precision of better than 10%. The analytical quality was assessed using Perade-09 certified reference material.

Plant samples were first digested using a mixture of concentrated H₂O₂ and HNO₃. The residual silica fraction, which was mainly composed of phytoliths, was dissolved separately using concentrated NaOH. Samples were then homogenized, and their silica concentrations were determined with an ICP-OES Horiba Ultima Expert at the GET lab (Toulouse, France). Soil samples from vertisol and ferralsol profiles were analysed for their major (including Si) and trace elements by ICP-OES at the SARM (Nancy, France) after digestion by borate fusion. The Si concentrations of ash samples were determined by ICP-OES after alkaline NaOH digestion at GET. The overall precision obtained for the concentration measurements in solids was 5 to 10%.

3.4 δ³⁰Si measurements

Solid samples (soils, litter and ASi zb) dedicated to δ³⁰Si measurements were digested and purified following the procedure described in Zambardi and Poitrasson (2011), which was itself modified from Georg et al. (2006). Soil and bedrock samples were powdered in an agate mortar, whereas litter samples were calcined at 900°C for 3 hours. Approximately 5 mg of powdered sample was melted with an ~200 mg NaOH pellet in a silver crucible and then dissolved in water and diluted HCl. The silicon fraction was separated from cations by column chromatography using Bio-Rad AG50W-X12 cationic resin. Si recovery was very close to 100%. Procedural blanks were lower than 50 ng Si, i.e., negligible compared to the amount of Si in each sample. Prior to isotopic measurements, the purified Si fraction was diluted to 2 or 3 mg L⁻¹ based on the initial amount of Si; the HCl concentration was adjusted to 0.05 mol L⁻¹; and 1 mg L⁻¹ of Mg was added as an internal standard for correcting online mass bias variations during the analysis. Isotopic compositions were measured using a Neptune (Thermo®) multi-collector ICP-MS in dynamic multicollection mode using medium- or high-resolution slits following
the methodology of Zambardi and Poitrasson (2011). The reference material NBS 28, which was also spiked with 1 mg L\(^{-1}\) of Mg, was measured between each sample (sample-standard bracketing). Most sample measurements were replicated 3 to 6 times. Isotopic data were normalized to NBS 28 and expressed as:

\[
\delta^{30}\text{Si} \ (\text{‰}) = \left( \frac{^{30}\text{Si}}{^{28}\text{Si}} \right)_{\text{sample}} \times 1000 - 1
\]

The uncertainty associated with the \(\delta^{30}\text{Si}\) data (2SE) corresponds to two standard errors obtained from the 3 to 6 replicates. The reproducibility was assessed by regular measurements of the reference material BHVO-2 (basaltic rock). The average \(\delta^{30}\text{Si}\) value obtained here for BHVO, i.e., -0.27 ± 0.06 ‰ (2SD, n=24), matches those obtained by Abraham et al. (2008) (-0.29 ± 0.11 ‰), Zambardi and Poitrasson (2011) (-0.27 ± 0.08 ‰), and Savage et al. (2012) (-0.30 ± 0.09 ‰). The standard deviation of the BHVO-2 measurements (±0.06 ‰) is considered here to represent the best achievable uncertainty of our \(\delta^{30}\text{Si}\) data.

3.5 Phytolith morphotypes and pollen analysis

Phytoliths extracted from the soil and suspended load samples by heavy liquid flotation were analysed under the microscope to evaluate the relative contributions of various plant types to the pool of ASi in the Mule Hole soils. Slides were mounted using Canada balsam. Phytolith morphotypes were described and counted at x400 and x1000 magnifications. They were classified based on their 3D shape and surface textures (Madella et al. 2005). They were then assigned their most likely botanical origin (grass, sedge, forest indicator, or non-diagnostic) according to e.g., Krishnan et al. (2000), Piperno (2006), Collura and Neumann (2016). Pollen was extracted from eight soil samples – 4 from each composite surface soil from the one-hectare plots and 4 from the vertisol profile (site D, Fig. 1) following standard protocols (Faegri et al., 1991 and Moore et al., 1991). Only seven of these samples could be analysed microscopically, as one contained too little pollen. Pollen taxa were counted under x400 and x1000 magnifications and identified using regional pollen atlases (Huang, 1972; Vasanthy, 1976; Nayar, 1990 and Tissot et al., 1994) as well as the Thanikaimoni pollen slide collection at the IFP as identification tools. The ecological attribution of the pollen taxa used here refers to Barboni and Bonnefille (2001), Barboni et al. (2003) and Anupama et al. (2014).
4 Results

4.1 $\delta^{30}\text{Si}$ values of bedrock, soils and suspended load

The $\delta^{30}\text{Si}$ value of the gneissic bedrock is $-0.34 \pm 0.08$‰ (Table 1), which falls within the lowest part of the range defined for igneous rocks (Savage et al., 2012). The gneiss signature is consistent with the light signatures observed in metasediment-derived igneous bedrock (Savage et al., 2012, Poitrasson & Zambardi, 2015). The isotopic signatures of the bulk ferralsol samples are very close to those of the bedrock composition. However, they slightly decrease with depth from $-0.26 \pm 0.08$‰ at 0-20 cm to $-0.42 \pm 0.27$‰ at 100-120 cm. These signatures are heavier than most of the values reported for tropical soils (e.g., Opfergelt et al., 2009; 2012, Bern et al., 2010; Ziegler et al., 2005).

The clay-size fraction of ferralsol samples, which mostly comprises kaolinite, yielded a $\delta^{30}\text{Si}$ value of $-1.30$‰ (n=3), which is homogeneous in the soil profile. The fractionation factor between kaolinite and bedrock is thus close to $-1$‰, which is less than the values that have been observed in other soils to date (Opfergelt and Delmelle, 2012 and references therein), which are closer to $-2$‰.

The suspended load of the overland flow yielded a $\delta^{30}\text{Si}$ signature of $-0.90 \pm 0.14$‰, which falls in between those of bulk soil and kaolinite, thus reflecting the mobilization of the fine grain size fractions from topsoil during rainy events.

4.2 Contribution to amorphous Si in soils: ASi content, phytolith morphotypes, pollen analysis, $\delta^{30}\text{Si}$ of ASi

4.2.1 ASi content

The amorphous silica (ASi$_{ab}$) concentrations in the soil profiles range from 0.05 to 0.24 % for ferralsol and from 0.06 to 0.86 % for vertisol (Table 2). The degree of ASi$_{ab}$ purity was elevated (>0.85) in surface soils and suspended load samples but decreased with ASi$_{ab}$ content and depth in soil profiles, down to 0.42 for the ferralsol sample collected at a depth of 200-220 cm (Table 2). The ASi$_{ab}$ concentrations in soils are comparable to those in other forest soils, both in humid tropics (0.02 % to 0.25 %, Congo, Alexandre et al., 1997; 0.3 to 2 %, Cameroon, Cary et al., 2005) and temperate climates (0.3-1.4 %, France, Cornelis et al., 2010). Their concentrations decreased with depth, as it is commonly the case in pristine tropical forests (Alexandre et al., 1997; Alexandre et al., 2011; Blecker et al., 2006; Saccone
et al., 2007; Saccone et al., 2008). This distribution reflects a balance between litterfall input and the dissolution/preservation of phytoliths in the soil profile. The ASi stocks in ferralsol and vertisol, assessed by extrapolating the measured values to the 2-m-thick profiles, should then range from 4 kg m\(^{-2}\) Si to 8 kg m\(^{-2}\) Si, respectively. These values are comparable to those observed in other mature tropical forests in Amazonia (Lucas, 2001) and the Congo (Alexandre et al. 1997) and are higher than those observed in savannas (Alexandre et al., 2011). The ASi\(_{ib}\) values of the four composite surface soils ranged from 0.4 to 0.65 %. This narrow range suggests a homogeneous phytolith content at the soil surface of Mule Hole. The corresponding stock for the 15-cm topsoil layer would be 1 kg m\(^{-2}\) ASi, which is equivalent to approximately 150 years of litter production (Riotte et al., 2014a) and 1000 to 2000 years of litter production for the whole soil layer.

4.2.2 Phytolith morphotypes

Phytolith counts are given in Electronic Annex EA-4 and summarized in Fig. 2. Under the microscope, the ASi appears to mostly comprise phytoliths. Diatoms represent less than 4% of all samples, including the stream and overland flow samples. The non-diagnostic elongate and blocky phytoliths are the dominant morphotypes (Fig. 2, micrographs in Electronic Annex AE-5) in all samples but the surface soil sample located in the Shorea forest, which contains abundant forest indicator phytoliths (Fig. 2). Grass/sedge indicators are more dominant than forest ones, but the abundance of grass/sedge indicators decreases with soil depth. The proportion of pitted phytoliths in ferralsol and vertisol profiles increases with depth, revealing an “aging effect” on deep phytoliths (Fig. 2).

4.2.3 Pollen analysis

Pollen counts are given in Electronic Annex EA-6. The analysis of the vertisol profile shows that more taxa with an affinity for evergreen forests are represented in the bottom of the vertisol profile (100-160 cm) compared to the top, which is characterized by pollen taxa with an affinity to moist deciduous forests (Electronic Annex AE-7). Changes associated with more seasonality and/or less annual precipitation are likely responsible for these changes in the pollen taxa composition through the vertisol profile at Mule Hole.
4.2.4 $\delta^{30}Si$ of ASi

The raw $\delta^{30}Si$ values in soil ASi range from -0.36‰ (ferralsol at a depth of 100-120 cm) to 1.56‰ (vertisol at a depth of 140-160 cm), which is a particularly wide range. However, the comparison between raw $\delta^{30}Si$ and ASi$_{zb}$ purity clearly shows that this variability is mostly due to impurities. The five ferralsol samples define a linear relationship ($R^2 = 0.997$) between the fresh litter (leaves & grass; ASi$_{zb}$ purity assumed to be 1 and an average measured $\delta^{30}Si$ value of 1.25‰) and the clay end-member (ASi$_{zb}$ purity = 0 and $\delta^{30}Si = -1.30‰$; Fig. 3).

**Fig. 2: Phytolith morphotypes and diatom contents in suspended load and soils of Mule Hole. ATT = Anogeissus – Tectona – Terminalia facies. ND = non-diagnostic.**

**Table: Phytolith morphotypes and diatom contents in suspended load and soils of Mule Hole. ATT = Anogeissus – Tectona – Terminalia facies. ND = non-diagnostic.**

<table>
<thead>
<tr>
<th>Suspended load</th>
<th>% Forest Indicators</th>
<th>% Grass/Sedge</th>
<th>% ND-Elongate</th>
<th>% ND-Blocky all</th>
<th>% Diatoms</th>
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<th>% Grass/Sedge</th>
<th>% ND-Elongate</th>
<th>% ND-Blocky all</th>
<th>% Diatoms</th>
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<td>(ATT) Vertisol/Amph.</td>
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<td>(Shorea) Sapro/Gneiss</td>
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<td>(Swamp) Vertisol/Gneiss</td>
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<th>% Grass/Sedge</th>
<th>% ND-Elongate</th>
<th>% ND-Blocky all</th>
<th>% Diatoms</th>
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<th>% ND-Elongate</th>
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Fig. 3: Influence of purity on $\delta^{30}\text{Si}$ signatures in the ASi$_{zb}$ fractions of surface soils, soil profiles and suspended load in Mule Hole. Purity index is defined as $(\text{Si}-(\text{Al}+\text{Fe}+\text{Ca}+\text{Mg}+\text{Mn}, \ K))/\text{Si}$. Purity index values of fresh litter and clays are theoretical. OF = overland flow. This diagram shows that despite being apparently pure under microscope, the ASi fraction separated by dense liquor can still contain significant amounts of impurities that can impact the $\delta^{30}\text{Si}$ signature of this fraction. Correcting for impurities allows us to recover the $\delta^{30}\text{Si}$ value of the pure ASi fraction.

The ASi$_{zb}$ content of the stream suspended load (0.37-0.52 %, Table 2) reflects the values of the overland flow (0.56 %) and the average topsoil composition of the watershed, as more than 90% of the runoff originates from overland flow (Maréchal et al., 2009). The $\delta^{30}\text{Si}$ signature of ASi$_{zb}$ in the suspended loads of overland flow and streams lies on the $\delta^{30}\text{Si}$ vs ASi$_{zb}$ purity relationship defined by ferralsol, making the “pure” $\delta^{30}\text{Si}$ value of the suspended load similar to the litter and surface soil signatures. In contrast to humid tropics, where the ASi of large rivers is dominated by diatoms (e.g., Hughes et al., 2011), our microscope observations indicate that in headwater streams such as Mule Hole or Mengong, Cameroon (Cary et al., 2005), ASi is almost exclusively composed of phytoliths. At Mule Hole, the ephemeral nature of the stream prevents diatom ingrowth. The similarities between the phytolith morphotypes of the stream, overland flow and ferralsol surface (Fig. 2) are consistent with the dominant contribution of overland flow on ferralsol to the ASi flux exported by the stream (Riotte et al., 2014a).
4.3 $\delta^{30}$Si of natural solutions

The $\delta^{30}$Si values of DSi in overland flow ($n=7$) ranged from 0.44 ± 0.16 ‰ to 2.22 ± 0.69‰, with a cluster around 1.08 ± 0.11‰ ($n=5$; Table 3). No shift in $\delta^{30}$Si was noticed during the rainy season, in contrast to the Si/Na and DOC/Na molar ratios, which increased as litter decayed (Riotte et al., 2014a). The $\delta^{30}$Si values of DSi in soil pore water was homogeneous whatever the soil type, 1.64 ± 0.11‰ ($n=9$; Table 3). Apart from the first two storms of 2004, during which litter-ash leaching released highly above-average Si fluxes (Audry et al., 2014) with specific $\delta^{30}$Si values, the $\delta^{30}$Si values of the stream water varied based on the water sources, with the lightest $\delta^{30}$Si values (0.28 to 0.74‰) occurring during flood peaks and the heaviest ones occurring during peak recession (1.04 to 1.61‰) (Table 3). Similar patterns were reported in tropical (Hughes et al., 2013) or temperate to boreal rivers (Georg et al., 2006; Engström et al. 2010; Pokrovsky et al., 2013). The range of $\delta^{30}$Si values in the Mule Hole stream of 1.3‰ is as wide as those observed in some large rivers, such as the Solimoes River from Amazonia (Hughes et al., 2013), which undergoes contrasted weathering regimes from the Andes to the Amazon plain, or Siberian rivers (1 to 1.5‰; Pokrovsky et al., 2013).

4.4 $\delta^{30}$Si values of ash leaching experiments

Ash leaching batches display large isotopic fractionation during ASi dissolution, especially at short interaction times. Incipient ash leaching (5 minutes), which coincides with the dissolution of the most soluble and alkaline salts, yielded extremely light $\delta^{30}$Si values, i.e., -0.77 ± 0.04 ‰ for T. triandra ash and down to -1.41 ± 0.02 ‰ for T. grandis ash. The solutions produced by long-term ash leaching (5 days) were isotopically heavier and fell within the range of the $\delta^{30}$Si values of phytoliths for T. triandra and T. crenulata, i.e., 1.45 ± 0.07 ‰ and 1.17 ± 0.06 ‰, respectively. The long-term ash leaching of T. grandis, yielded a $\delta^{30}$Si value of 2.29 ± 0.02‰, which is significantly higher than the bulk signature and may have resulted from the small-scale heterogeneity of this sample. The $\delta^{30}$Si values of all three leaching experiments increased with the proportion of dissolved ASi, calculated as the amount of Si in solution divided by the initial amount of Si in the litter ash; with the notable exception of the long-term ash leaching of T. grandis, they all defined a broad trend matching that of another leaching experiment performed on Hawaiian phytoliths (Ziegler et al., 2005) (Fig 4).
Fig. 4: Evolution of $\delta^{30}\text{Si}$ in the dissolved phases of ash leaching experiments with the percentage of dissolved litter-Si (fraction of Si release divided by the total Si content in the bulk litter). Grey area: range of phytolith compositions found in this study. Open circles: phytolith dissolution experiments from Ziegler et al. (2005), with dotted line representing the isotopic composition of the phytolith.

5 Discussion

5.1 Silicon pools in the soil and variability of $\delta^{30}\text{Si}$ -ASi in the watershed

When compared to the weathering degree estimated from the Chemical Index of Alteration (Nesbitt and Young, 1982) or the Weathering Index of Parker (Parker, 1970), the Mule Hole soils (CIA 80 to 84, WIP 14 to 15) plot slightly away from the trend defined by worldwide soil data (Opfergelt & Delmelle, 2012): for a given weathering index, the $\delta^{30}\text{Si}$ values of the Mule Hole soils are heavier than those in most other locations. The narrow range of Si isotopic fractionation between the bedrock and soils of Mule Hole can be explained by the mineralogical composition of the soils and particularly by the partitioning of silicon between primary and secondary phases. Approximately 86 to 95% of soil silicon is still located in primary minerals (e.g., quartz, plagioclase), whereas only 4 to 13% is located in clay minerals, mostly kaolinite from ferralsols (Barbiero et al., 2007; Braun et al. 2009). Such clay mineral contents are low compared to those of 35 to 85% found, for instance, in Guadeloupe soils (Opfergelt et al., 2012). Quartz accounts for 70 to 79% of the total Si of the soil (Table 1) and therefore dominates the soil isotopic signature. The fact that quartz abundance is not considered in the calculation of weathering indices could explain why the Mule Hole data plot outside the trends of worldwide soil $\delta^{30}\text{Si}$ vs weathering indexes (Opfergelt & Delmelle, 2012), as well as the scattering of these trends. Our data confirm the conclusion of Opfergelt
and Delmelle (2012) that the $\delta^{30}\text{Si}$ values of bulk soils depend on the ratio of clay to primary minerals. The relatively high $\delta^{30}\text{Si}$ value of the clay fraction could possibly be explained by a slight contribution of quartz to this fraction, which would slightly increase the $\delta^{30}\text{Si}$ value, as was recently proposed by Ameijeiras-Mariño et al. (2017). Alternatively, some minerals of the bedrock could exhibit slightly heavier $\delta^{30}\text{Si}$ values than the bulk rock, which could account for the limited apparent fractionation factor. However, the narrow range of $\delta^{30}\text{Si}$ values observed for separate silicate minerals (Savage et al., 2014) or modelled between quartz and albite at 600 °C (<0.1 ‰, Méheut et al., 2014) makes this assumption unlikely.

The low purity index of ASi$_{zb}$ in deep ferralsol samples attests to the difficulty of recovering the soil ASi fraction, despite using an elaborate purification protocol (see section 3.2) and further microscope quality control. In contrast to the microscope control, which is qualitative, the chemical control, as assessed through the purity index, is sensitive and quantitative. It allows us to quantify contamination and correct the $\delta^{30}\text{Si}$ value of ASi when necessary. In the case of deep ferralsol samples, the clay mineral contribution was significant and impacted the $\delta^{30}\text{Si}$ of ASi$_{zb}$, which thus must be corrected. The alignment of the clay and fresh litter compositions of 5 ASi$_{zb}$ ferralsol samples (Fig. 3) indicates that the $\delta^{30}\text{Si}$ of ASi$_{zb}$ is constant along the ferralsol profile and corresponds to the average signature of fresh litter. It is remarkable that the ASi$_{zb}$ $\delta^{30}\text{Si}$ values of the four composite surface soils, the shallow vertisol sample and the suspended load of the overland flow and stream also lie close to the relationship defined by ferralsol samples, implying that the ASi$_{zb}$ $\delta^{30}\text{Si}$ can be corrected by extrapolation to an ASi$_{zb}$ purity of 1. Then, the $\delta^{30}\text{Si}_{\text{corr}}$ values of all these ASi$_{zb}$ samples (Table 2) fall within a narrow range, between 1.10 and 1.30 ‰, corresponding to the present-day litter phytolith composition.

In contrast to the ferralsol profile and composite surface soils, the ASi fractions of the vertisol profile exhibit scattered ASi$_{zb}$ $\delta^{30}\text{Si}$ compositions, i.e., from 0.52 to 1.56 ‰ (uncorrected from clay impurities), although the contribution of impurities does not exceed 20%. If the clay signature in this soil profile is the same as that of the ferralsol, then the ASi$_{zb}$ $\delta^{30}\text{Si}_{\text{corr}}$ values of deeper vertisol samples should vary from 0.62 to 2.10 ‰. The presence of smectite in vertisols may induce a heavier $\delta^{30}\text{Si}$ value of the clay pool, as was observed for another clay mineral, illite (-0.68 ‰, Steinhofel et al., 2011). However, because of the purity level of the vertisol ASi fraction, the clay signature only marginally impacts the $\delta^{30}\text{Si}_{\text{corr}}$ value; if the clays have a $\delta^{30}\text{Si}$ value of -0.5 ‰, the range of the $\delta^{30}\text{Si}_{\text{corr}}$ values of deeper vertisol samples would be still 0.5 to 2.0 ‰. Rather, the scattered signatures could result from the impervious
properties of vertisols (Kurtzman et al., 2016) that could host preserved phytoliths from past vegetation/climate conditions or have induced opal precipitation due to the intense evaporation of soil pore water. Pedogenic carbonates dated to the Last Glacial Maximum, when semi-arid conditions prevailed (Violette et al., 2010a), are present in the vertisols of Mule Hole. Given the low permeability of vertisols, phytoliths from this period could also have been preserved, which would also explain their higher abundance in vertisols compared to ferralsols. Indeed, the phytolith morphotypes at the vertisol surface (Fig. 2; Supplementary table 1), which are dominated by elongate shapes, differ from those of deeper horizons, where blocky shapes are dominant relative to elongate shapes. The antiquity of these phytoliths is indicated by an increase in the proportions of pitted phytoliths with depth, probably due to weathering over time, whereas the absence of cultivation pollen markers below a depth of 100 cm suggests that those soils were preserved from significant human impacts. A spike in certain ubiquitous pollen taxa in tandem with the spike in evergreen taxa below this depth suggests a mosaic landscape of patches or islands of wet/humid forests separated by openings, which is certainly different from the present landscape in terms of its species composition (Electronic Annex AE-7). Then, the large variability of ASi$_{zb}$ $\delta^{30}$Si values in deeper vertisol horizons could come from phytoliths of a vegetation cover different than the present one. One cannot exclude that ageing effects also impacted ASi$_{zb}$ $\delta^{30}$Si. Alternatively, it is known that the seasonal swelling/cracking properties of vertisols favour pore water evaporation and then salinization (e.g., Baram et al., 2013). Deep soil pore water could then reach saturation with opal and allow opal precipitation at this depth. If this process is not completely reversible over a monsoon-dry season cycle, small amounts of opal could have accumulated over time in the soil and then contributed to the ASi$_{zb}$ fraction of deep vertisol samples, possibly also impacting the $\delta^{30}$Si signature of ASi$_{zb}$.

To summarize, apart from two depths in the vertisol in which phytoliths likely represent fossils of another vegetation cover, the $\delta^{30}$Si value of ASi remains remarkably constant throughout the watershed, which confirms that the $\delta^{30}$Si values of vegetation are “system-dependent” rather than “vegetation species-dependent” (Geilert et al., 2014). However, it remains unclear why two vertisol samples exhibit a wide range of ASi $\delta^{30}$Si values, i.e., from 0.6 to 2.1 ‰.
5.2 Silicon isotopic solute mass balance at the soil-plant scale

5.2.1 Initial elemental Si mass balance

The Si isotopic mass balance at the soil-plant scale (Fig. 5) relies on the initial elemental mass balance performed by Riotte et al. (2014a). At the ground surface, the solute Si flux infiltrating the soil layer (i.e., the soil input flux), assuming an overland flow composition, is 1850 mol ha\(^{-1}\) yr\(^{-1}\) of DSi, which is similar to that brought to the ground by litter fall, i.e., 2200 ± 200 mol Si ha\(^{-1}\) yr\(^{-1}\). The origin of the Si infiltrating the soil was assigned to ASi dissolution from litter, based on the good relationship between the Si/Na, DOC/Na, and Ca/Na ratios that all increase during the monsoon season as litter decays. Since the Na concentration in overland flow was identical to that of rainfall and throughfall, the contribution of primary minerals, such as Na-plagioclase, to the overland flow solution was considered to be negligible, while its oversaturation with respect to clay minerals indicated that clays were not a potential source. Within the soil, the Si solute mass balance was performed using both Si and Na budgets (Fig. 5, left side, from Riotte et al., 2014a). With a DSi output of only 704 ± 150 mol ha\(^{-1}\) yr\(^{-1}\), the input-output budget indicates an apparent loss of 1150 mol DSi ha\(^{-1}\) yr\(^{-1}\) from the soil layer. However, the net soil output of 650 mol Na ha\(^{-1}\) yr\(^{-1}\), which could only originate from the weathering of Na-plagioclase, likely induced the release of an additional 1300 mol DSi ha\(^{-1}\) yr\(^{-1}\) if kaolinite is the weathering product, according to the following weathering reaction (Riotte et al., 2014a):

\[
2\left(\text{Na}_{0.92}\text{Ca}_{0.08}\right)\left(\text{Al}_{1.07}\text{Si}_{2.93}\right)\text{O}_8 + 2\text{H}^+ + 9\text{H}_2\text{O} \rightarrow 1.07\text{Si}_2\text{Al}_2\text{O}_5(\text{OH})_4 + 1.84\text{Na}^+ + 0.16\text{Ca}^{2+} + 3.72\text{H}_4\text{SiO}_4 \quad (3)
\]

Then, according to this Na-Si solute budget, the DSi loss from the soil layer would be at least 2450 mol ha\(^{-1}\) yr\(^{-1}\); this value is similar to the annual litter flux (tree leaves and grass). The loss of Si from soil pore water was attributed to vegetation uptake.

5.2.2 Si mass balance inferred from isotopes

The Si mass balance inferred from isotopes consists of multiplying the elemental Si fluxes of section 5.2.1 by their isotopic signature (\(\delta^{30}\text{Si}\)). When \(\delta^{30}\text{Si}\) values were obtained from several samples of a given compartment, the variability between these data was taken into account using two standard deviations between samples.

The elemental mass balance inferred from isotopes first provides the opportunity to test the assumption of the biogenic origin of Si in overland flow proposed by Riotte et al. (2014a).
Accordingly, the three potential sources of Si, namely, fresh litter (1.25 ± 0.14 ‰), primary minerals (-0.34 ‰) and clay minerals (-1.30 ‰), exhibit contrasting δ³⁰Si values. The average δ³⁰Si of overland flow DSi (1.15 ± 0.50 ‰) is much heavier than the signature of the clay-size fraction (-1.30 ‰), thus ruling out any contribution of clay mineral dissolution to the silica content in overland flow. It is indeed slightly lower than the fresh litter average, 1.25 ± 0.14 ‰, which is consistent with the fractionation factor Δ³⁰Si_ASi-DSi of -0.26 ‰ estimated for ASi dissolution in the delta of Okavango (Frings et al., 2014a) and in the range observed during diatom dissolution, i.e., approximately 0 ‰ (Wetzel et al., 2014) to -0.55 ‰ (Demarest et al. 2009). The similarity between the δ³⁰Si values of overland flow and fresh litter leaves no doubt about the biogenic origin (phytolith dissolution) of DSi in overland flow. By extension, the biogenic origin of Si in solutions infiltrating the soil (Riotte et al., 2014a) is therefore confirmed.

The isotopic mass balance then allows us to evaluate whether the plant uptake of Si is limited to cycling through the litter or involves a larger flux with storage in the woody biomass. Tropical trees are known to accumulate silica in their woody stems; at least one of the most abundant tree species of Mule Hole, T. grandis, contains 0.27 to 0.66 % of silica in its wood (Kjaer et al., 1999). If the Si stored in a plant were concentrated in the litter, δ³⁰Si_uptake should be equal to δ³⁰Si_litter, whereas if Si is partly stored in the woody biomass, δ³⁰Si_uptake should differ from δ³⁰Si_litter because Si isotopes fractionate within the plants (Opfergelt, 2006; Ding et al. 2005). The δ³⁰Si_uptake can first be estimated from the solute isotopic budget within the soil layer, which requires an assessment of the δ³⁰Si value of the silica released during plagioclase weathering. If plagioclase crystals break down into kaolinite with the release of dissolved Na and silica (a process referred to in Fig. 5 as plagioclase weathering and described by the above weathering reaction), the δ³⁰Si value of the silica released can be estimated according to a mass balance equation (Bouchez et al., 2013):

\[
\delta^{30}\text{Si}_{\text{released}} = \frac{FSi_{\text{plagioclase}}\delta^{30}\text{Si}_{\text{plagioclase}} - FSi_{\text{kaolinite}}\delta^{30}\text{Si}_{\text{kaolinite}}}{FSi_{\text{released}}} \tag{4}
\]

Using the reaction stoichiometry (FSi values) given in the weathering reaction (3) (from Riotte et al., 2014a) and the δ³⁰Si_{kaolinite} values measured in ferralsol clays (-1.30 ± 0.03‰) and assuming that the δ³⁰Si_{plagioclase} value is similar to the bedrock signature (-0.34 ± 0.08‰), the estimated δ³⁰Si_{released} value produced by Na-plagioclase weathering would be
+0.21 ± 0.14‰. Adding the DSi flux infiltrated from the surface (1850 mol ha⁻¹ yr⁻¹ with δ³⁰Si of 1.15 ± 0.50‰) to the Si released by Na-plagioclase weathering indicates that the minimum DSi flux to the soil would have an average δ³⁰Si value of +0.76 ± 0.35‰. Since the soil output of 842 mol DSi ha⁻¹ yr⁻¹ (Riotte et al., 2014a) has an average δ³⁰Si value of +1.64 ± 0.11‰, the silica removed from soil pore water and assigned to plant uptake should exhibit a δ³⁰Si value of +0.52±0.30‰. This signature is 0.7‰ lower than the average δ³⁰Si_litter value, which means that plants likely take up more silica than expected from the elemental mass balance. Therefore, the latter must be re-evaluated and isotopically equilibrated.

5.2.3 Equilibration of the Si isotopic mass balance

In this section, we propose a three-step methodology for equilibrating the Si isotopic mass balance, which consists of (1) estimating the δ³⁰Si value of the bulk Si taken up by plants, (2) estimating the corresponding flux of the bulk Si uptake and (3) identifying the potential additional Si sources to plants:

(1) The Si isotopes taken up by plants could theoretically follow either a Rayleigh model, implying that the Si source is limited (i.e., a closed system), or a steady-state model, implying a non-limited Si source and open system. The Rayleigh model could apply, for instance, to a pot experiment with no other Si supply than the initial hydroponic solution. In the case of Mule Hole, the intensity of soil mineral weathering (for instance, ASi or Na-plagioclase) varies seasonally based on soil moisture fluctuations. However, soil moisture hardly falls below 20% in the ferralsol and 25% in the vertisol (Parate et al., 2011), which is sufficient to sustain soil-Si release throughout the year. This soil-Si flux complements the Si flux from litter decay that regularly infiltrates the ground surface during the monsoon season. The continuous (but obviously temporally variable) supply of Si makes the soil a relatively open system with a non-limited source of Si. As a consequence, a steady-state model would be more appropriate than the Rayleigh model to describe the silicon isotopic fractionation induced by plant uptake. Accordingly, the δ³⁰Si_{uptake} value would be linked to the soil pore water composition by the following relation (De La Rocha et al., 1997):

\[ \delta^{30} \text{Si}_{\text{uptake}} = \delta^{30} \text{Si}_{\text{soil porewater}} + \Delta^{30} \text{Si} \] (5)
With $\Delta^{30}\text{Si}$ being the isotopic fractionation factor between plants and solution. Although this parameter cannot be measured for a forest, it was measured for a few plants, such as rice (Ding et al. 2005) and banana (Opfergelt et al. 2006). Both plants exhibited the same $\Delta^{30}\text{Si}_{\text{plant}}$ value of -1‰. In this study, we assumed that this value applies to the silicon uptake of the forest. Taking into account the average $\delta^{30}\text{Si}_{\text{soil porewater}}$ value measured in the Mule Hole soil pore water, 1.64 ± 0.11‰ (n=9), the $\delta^{30}\text{Si}_{\text{uptake}}$ deduced from Eq. 3 would then be 0.64 ± 0.11‰. This value is robust because the silicon isotopic composition of the deep soil pore water is homogeneous over space and time (Table 3). This confirms that the total Si uptake of the vegetation should be larger than the flux measured in the litter and that its $\delta^{30}\text{Si}$ value should be enriched in light isotopes.

(2) Both the proportion of Si annually stored in the stem and its $\delta^{30}\text{Si}$ value can be deduced from $\delta^{30}\text{Si}_{\text{uptake}}$ and $\delta^{30}\text{Si}_{\text{litter}}$ assuming that silicon isotopes, once taken up by trees, follow a classical Rayleigh fractionation model, as has been observed for some plants, such as rice (Ding et al., 2005), banana (Opfergelt et al., 2006), bamboo (Ding et al., 2008) and maize (Sun et al., 2017). Silica precipitation in the woody stem can be considered to be the cumulative product while the remaining fraction, which ends up in the leaves, can be considered to be the reactant. The $\delta^{30}\text{Si}$ values of leaves are assimilated into the measured litter signature $\delta^{30}\text{Si}_{\text{litter}}$. Then, the fraction of Si annually stored in the stem can be expressed as:

$$f_{\text{woody stem}} = 1 - f_{\text{litter}} = 1 - e^{-\left(\frac{\delta^{30}\text{Si}_{\text{litter}} - \delta^{30}\text{Si}_{\text{uptake}}}{\Delta^{30}\text{Si}_{\text{Si}}^t}\right)}$$  \hspace{1cm} (6)

Considering the abovementioned $\delta^{30}\text{Si}_{\text{uptake}}$ and $\delta^{30}\text{Si}_{\text{litter}}$, the fraction of Si that is annually stored in woody stem would be 0.46 ± 0.02, which accounts for almost as much as the litter flux. As a consequence, the overall Si taken up by vegetation would reach 4100 mol ha⁻¹ yr⁻¹, which is twice the value initially estimated from the solute elemental budget (Riotte et al., 2014a). The $\delta^{30}\text{Si}_{\text{woody stem}}$ value can be deduced by rearranging the classical Rayleigh relationship:

$$\delta^{30}\text{Si}_{\text{woody stem}} = \delta^{30}\text{Si}_{\text{uptake}} - \Delta^{30}\text{Si} \cdot f_{\text{litter}} \cdot \frac{\ln(f_{\text{litter}})}{1-f_{\text{litter}} - f_{\text{woody stem}}}$$  \hspace{1cm} (7)
Here, the $\delta^{30}\text{Si}_{\text{woody stem}}$ value would be $-0.08 \pm 0.14\%o$. The fact that the Si accumulated in woody stems is isotopically lighter than that in the leaves is consistent with the progressive enrichment in $^{30}\text{Si}$ from the lowest parts of the plant (roots) to the highest ones (leaves) observed by Ding et al. (2005) and Opfergelt (2006).

(3) In the absence of additional proxies of silicate weathering fluxes or Si sources, one can only suggest hypotheses to explain the origin of the Si flux stored in the woody stem and its isotopic signature of $-0.08 \pm 0.14\%o$. If the Si source(s) consist(s) of a dissolved compartment, based on the above assumptions, its uptake should follow steady-state conditions, with a $\Delta^{30}\text{Si}_{\text{plant}}$ value of $-1 \%o$. The Si source should exhibit a $\delta^{30}\text{Si}$ value of approximately $+1 \%o$, which corresponds to the soil ASi ($1.25 \%o$ in ferralsols) and groundwater ($1.20 \pm 0.3\%o$; Table 3) signatures. The current stock of 4 kg ASi m$^{-2}$ in the ferralsol could indeed sustain the additional flux of 1900 mol Si ha$^{-1}$ yr$^{-1}$ for 600 years (twice as long for vertisols). This pool was also recently identified as the principal Si source for rice (Riotte et al., 2017) and may thus sustain many Si-accumulator plants. If plants directly take up Si from the soil compartments, with limited or no isotopic fractionation, then the $\delta^{30}\text{Si}$ value of $-0.08 \pm 0.14\%o$ in the stems could originate from the congruent dissolution of primary silicate minerals. The best rock-forming mineral candidates would be Al-poor phases, such as the ubiquitous Mg-hornblende and quartz, because their weathering would not induce the formation of clay minerals and thus result in limited to no silicon isotopic fractionation. In all cases, the contribution of clay minerals as a source of Si to plants is unlikely. The isotopic solute budget of silica at the soil-plant scale is summarized in Fig. 5.
Fig. 5: Isotopic budget at the soil-plant scale in the Mule Hole watershed, with the annual Si flux (bold; in mol ha⁻¹ yr⁻¹) and its δ³⁰Si value (italic; in ‰) for each compartment. Values in black were measured; those in red were inferred from elemental and/or isotopic mass balance. This budget was performed in three steps: (1) primary isotopic budget based on the elemental Si budget from Riotte et al. (2014a), (2) deduction of woody stem storage that balances the plant budget of part 1, (3) additional fluxes from the soil layer needed to equilibrate the whole isotopic soil-plant budget. Blue text: DSi fluxes related to water fluxes. Grey text: DSi flux released from minerals (independent from water budget). Uncertainties given by Riotte et al. (2014a) for computed water fluxes (20%) were used for the dissolved chemical fluxes. Uncertainty associated with the average measured litter flux is considered as the variability between the two measured years 2009 and 2010 (Riotte et al., 2014a).

To summarize the Si isotopic budget at the soil-plant scale, the δ³⁰Si signatures of overland flow, soil pore water and litter confirm the biogenic origin of silica in the soil-percolating solutes and add a subsequent constraint to the elemental budget; the Si uptake by vegetation should be twice as large as the previous estimation, which was based on the elemental budget alone (Riotte et al., 2014a).
5.3 Controls of $\delta^{30}$Si in stream dissolved and suspended loads

The few studies that have reported time-related fluctuations of $\delta^{30}$Si in rivers have proposed various explanations. Hughes et al. (2013) observed that for several streams of the Amazon River system, the highest $\delta^{30}$Si values corresponded to base flow conditions and low Si/cation molar ratios. These compositions were interpreted to reflect the greater contribution of the chemical weathering of primary minerals due to a greater contribution of groundwater. The chemical compositions of these waters were compatible with either smectite (bisiallitization) or kaolinite (monosiallitization) production. In Siberian watersheds, the highest $\delta^{30}$Si occurred during the summer-fall season, when biomass and the active soil layer thickness increased. Two explanations were proposed to explain these high signatures, i.e., the formation of secondary minerals (clays and amorphous phases) or Si uptake by vegetation, although it was not possible to determine the respective contribution of each process (Pokrovsky et al., 2013). According to the hydrological model of the Mule Hole watershed (Violette et al., 2010b), the ephemeral stream is mostly fed by overland flow on ferralsol ($53 \text{ mm yr}^{-1}$) and vertisol ($29 \text{ mm yr}^{-1}$). Soil seepage contributes slightly to the runoff at the end of the rainy season, when soils are saturated ($12 \text{ mm yr}^{-1}$). The relative contribution of each source is consistent with the evolution of the chemical composition during a given storm as well as during the rainy season (Riotte et al., 2014a).

Fig. 6: Silicon isotopic composition of DSi in the Mule Hole stream during storms (values in red, in per mil). Green dots represent Si/Na molar ratios and blue lines represent the stream discharge (modified from Riotte et al., 2014a).
Both the high δ\textsuperscript{30}Si values and chemical compositions during flood recession reflect the contributions of soil seepage (Fig. 6), which is consistent with the expected water contributions in the stream (Violette et al., 2010b; Riotte et al., 2014a). Based on the isotopic mass balance at the soil-plant scale, it is possible to confidently assign the enrichment in the heavy Si isotopes of soil pore water, and then of flood recession, to the combination of both primary silicate chemical weathering and plant uptake.

Various processes have been suggested to explain the low δ\textsuperscript{30}Si values in rivers. In the Amazon Basin, the lowest δ\textsuperscript{30}Si values were found in the plain (Rio Negro) and attributed to an increase in the relative silica mobility (f\textsubscript{Si}), defined by the ratio of the Si/(Na\textsuperscript{8+}+K\textsuperscript{+}) molar ratio of the dissolved phase (Na and K corrected from atmospheric inputs) divided by the same ratio of the bedrock (Hughes et al., 2013). According to these authors, the silicate bedrock would dissolve in a congruent way, i.e., with a limited fraction of isotopically light Si relocated to secondary clay minerals (kaolinite). Most of the δ\textsuperscript{30}Si values of the large rivers of the Amazonian basin were indeed inversely correlated to f\textsubscript{Si}, which was compatible with Rayleigh or steady-state fractionation models. Such findings are consistent with those of Frings et al. (2015), who used a similar proxy in the Ganga Basin, and Georg et al. (2007) in Iceland, who used the ratio between DSi and Na as a silica mobility index. In both cases, the role of vegetation in the δ\textsuperscript{30}Si values of rivers was considered to be unlikely. However, neglecting the role of vegetation in the case of the Amazon plain is surprising, as it is covered by a dense forest, in which silica recycling by litter accounts for up to 1500 mol ha\textsuperscript{-1} yr\textsuperscript{-1} (Cornu et al., 1998). According to our observations at Mule Hole, the high K\textsuperscript{+}/Na\textsuperscript{+} values associated with the low δ\textsuperscript{30}Si values in the Amazonian rivers (Hughes et al., 2013) could also result from the contribution of vegetation cycling. Very low δ\textsuperscript{30}Si values (close to 0‰) were also observed in the black rivers of the Congo Basin and interpreted to represent the dissolution of clay minerals by organic-rich waters (Cardinal et al., 2010). In Siberian watersheds, the low δ\textsuperscript{30}Si values during spring floods (0.8‰) were assigned to the partial dissolution of the silicate suspended load because of the high solid/solution ratio (Pokrovsky et al., 2013). During the flow peaks of Mule Hole, the stream δ\textsuperscript{30}Si is also low (0.47 ± 0.22‰) and surprisingly lighter than the overland flow signature (1.15 ± 0.53‰). This is unexpected since the stream is fed by overland flow during flow peaks. An additional DSi source to the stream is needed. As this DSi source should exhibit a low δ\textsuperscript{30}Si value, it could correspond to the dissolution of the suspended load, as proposed by Pokrovsky et al. (2013) for Siberian rivers. However, the average DSi concentration during these flow peaks
(128 ± 35 µmol L\(^{-1}\)) exceeds saturation with respect to kaolinite and smectite, thus making clay dissolution unlikely. Moreover, the circum-neutral pH and low DOC content of the stream do not favour clay mineral dissolution. An alternative explanation could be partial, short-term (i.e., hourly) phytolith dissolution within the stream which, according to ash leaching experiments (see section 4.3), could release light Si isotopes. If the \(\delta^{30}\text{Si}\) value of the additional DSi source were -1‰, it would contribute to up to 30% of stream DSi, which is unlikely because ash leaching experiments indicate that the kinetic isotopic fractionation that releases light Si isotopes is expected to occur during incipient ASi dissolution (see Fig. 4). Sampling bias thus remains the most likely cause of the difference between the \(\delta^{30}\text{Si}\) values of the overland flow and streams.

The ASi flux exported annually by the stream as suspended load was estimated using the 2003-2014 specific runoff of 100 mm yr\(^{-1}\), the average suspended load content of 0.5 g L\(^{-1}\) and the average suspended load ASi\(_{sb}\) concentration of 0.44%. The ASi specific flux exported accounts for 89 mol ha\(^{-1}\) yr\(^{-1}\), which represents only 4% of the annual flux produced by litter fall; this is in good agreement with the data from the Kaveri River, India (Meunier et al., 2015). This also confirms that the silica mobilized by the vegetation at Mule Hole is rarely exported as ASi but is rather accumulated or recycled within the watershed (Riotte et al., 2014a). However, the specific ASi flux exported by the stream is close to the DSi flux for the period 2005-2009 of 130 mol ha\(^{-1}\) yr\(^{-1}\) (Riotte et al., 2014a). Similarities in the exported ASi and DSi were also recently observed in the Ganga River (Frings et al., 2014b) and represent further evidence that biota play a significant role in the export of silica to the ocean.

6 Conclusion

The investigation of silicon isotopic fractionation during water-rock/soil-plant interactions in the forested watershed of Mule Hole provided new constraints on the biogeochemical cycling of silica in this dry tropical forest:

- The \(\delta^{30}\text{Si}\) composition of amorphous Si in litter (tree leaves and grass), surface soil horizons and the ferralsol profile is particularly homogeneous, with a range of only 1.1 to 1.4‰. The only exceptions are deeper horizons from the vertisol profile (in a marginal area of the watershed) that also contain pollen taxa typical of humid forests. Phytoliths from these horizons may thus be representative of past, more humid conditions. The homogeneity of the \(\delta^{30}\text{Si}\) values in ASi confirms that the silicon isotope signature of
vegetation is “system-dependent”. Accordingly, δ³⁰Si should be suitable for tracing the biogenic component of silica cycling at both the soil-plant and watershed scales.

- At the soil-plant scale, the δ³⁰Si value of solutes percolating into the soil is similar to the isotopic signature of phytoliths, which supports the biogenic origin of DSi in the soil input. Within the soil, the assignment of δ³⁰Si signatures to the solute elemental budget suggests an imbalance. The isotopic equilibration of the solute budget, considering a steady-state model for plant uptake and Rayleigh fractionation within the plants, implies a silica flux annually taken up by vegetation that is twice as large as that first estimated from the elemental budget (Riotte et al., 2014a). This additional Si flux, which may be stored in woody stems, should exhibit a δ³⁰Si value close to 0 ‰. Such a signature could be compatible with soil ASi dissolution or deep root uptake if a steady-state model applies. If vegetation takes up Si directly from soil particles without isotopic fractionation, then primary silicate or Si desorption from clays could constitute possible sources. We observed that deep groundwater exhibits the same signature as litter dissolution, meaning that biotic (cycling through vegetation) and abiotic (silicate chemical weathering) reactions may lead to comparable δ³⁰Si signatures.

- Ephemeral stream storms tend to display low δ³⁰Si values during peak flows, whereas high δ³⁰Si values are observed during recessions, but only at the end of the rainy season, when soils are saturated. This pattern has been observed in most previous river studies. The chemical and isotopic compositions of stream recessions tend consistently towards soil pore water compositions, in agreement with the hydrological functioning of such an ecosystem. However, a slight difference was observed between the δ³⁰Si values of overland flow, which feeds the stream during flood peaks, and those of the stream. The contribution of clay mineral dissolution from the suspended load is unlikely, given the DSi concentration in the stream. We suggest that this slight difference results from a bias in overland flow sampling.

As a broader conclusion, it appears that stable silicon isotope fractionation provides significant insights into the processes controlling the biogeochemical cycle of silica in ecosystems, but only if water and elemental mass balances are already well constrained.

Acknowledgments
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**Figure & tables captions**

**Figure 1:** The Mule Hole watershed with soil and vegetation map, showing the sampling location of composite soils (plots A, B, C, D; see text for description), soil profiles, overland flow and ground water (modified from Riotte et al., 2014a). ATT (medium green): Anogeissus – Tectona – Terminalia facies. Dark green: swamp facies; light green: Shorea facies.

**Figure 2:** Phytolith morphotypes and diatom contents in suspended load and soils of Mule Hole. ATT = Anogeissus – Tectona – Terminalia facies. ND = non-diagnostic.

**Figure 3:** Influence of purity on $\delta^{30}\text{Si}$ signatures in the ASi$_{zb}$ fractions of surface soils, soil profiles and suspended load in Mule Hole. Purity index is defined as $(\text{Si-(Al+Fe+Ca+Mg+Mn, K)})/\text{Si}$. Purity index values of fresh litter and clays are theoretical. OF = overland flow. This diagram shows that despite being apparently pure under microscope, the ASi fraction separated by dense liquor can still contain significant amounts of impurities that can impact the $\delta^{30}\text{Si}$ signature of this fraction. Correcting for impurities allows us to recover the $\delta^{30}\text{Si}$ value of the pure ASi fraction.

**Figure 4:** Evolution of $\delta^{30}\text{Si}$ in the dissolved phase of ash leaching experiments with the percentage of dissolved litter-Si (fraction of Si release divided by the total Si content in the bulk litter). Grey area: range of phytolith compositions found in this study. Open circles: phytolith dissolution experiments from Ziegler et al. (2005), with dotted line representing the isotopic composition of the phytolith.

**Figure 5:** Isotopic budget at the soil-plant scale in the Mule Hole watershed, with the annual Si flux (bold; in mol ha$^{-1}$ yr$^{-1}$) and its $\delta^{30}\text{Si}$ value (italic; in ‰) for each compartment. Values in black were measured, those in red were inferred from elemental and/or isotopic mass balance. This budget was performed in three steps: (1) primary isotopic budget based on the elemental Si budget from Riotte et al. (2014a), (2) deduction of woody stem storage that balances the plant budget of part 1, (3) additional fluxes from the soil layer needed to equilibrate the whole isotopic soil-plant budget. Blue text: DSi fluxes related to water fluxes. Grey text: DSi flux released from minerals (independent from water budget). Uncertainties given by Riotte et al. (2014a) for computed water fluxes (20%) were used for the dissolved chemical fluxes. Uncertainty associated with the average measured litter flux is considered as the variability between the two measured years 2009 and 2010 (Riotte et al., 2014a).

**Figure 6:** Silicon isotopic composition of DSi in the Mule Hole stream during storms (values in red, in per mil). Green dots represent Si/Na molar ratio and blue lines represent the stream discharge (modified from Riotte et al., 2014a).
References


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**Figure 2**

- **Suspended load**
  - Stream (3212-24)
  - Stream (3316-28)
  - Overland flow (4056)
  - Overland flow (4080)

- **Surface composite soils**
  - (ATT) Ferral./ Gneiss
  - (ATT) Vertisol/ Amph.
  - (Shorea) Sapro/ Gneiss
  - (Swamp) Vertisol/ Gneiss

- **Vertisol profile**
  - 0-20
  - 40-60
  - 100-120
  - 140-160

- **Ferralsol profile**
  - 0-20
  - 40-60
  - 100-120
  - 140-160
  - 200-220

Legend:
- % Forest Indicators
- % Grass/Sedge
- % ND-Elongate all
- % ND-Blocky all
- % ND-Elongate
- % ND-Blocky
- Not pitted
- Pitted
- % Diatoms

- Suspended load:
  - Stream (3212-24)
  - Stream (3316-28)
  - Overland flow (4056)
  - Overland flow (4080)

- Surface composite soils:
  - (ATT) Ferral./ Gneiss
  - (ATT) Vertisol/ Amph.
  - (Shorea) Sapro/ Gneiss
  - (Swamp) Vertisol/ Gneiss

- Vertisol profile:
  - 0-20
  - 40-60
  - 100-120
  - 140-160

- Ferralsol profile:
  - 0-20
  - 40-60
  - 100-120
  - 140-160
  - 200-220
Figure 3

The graph shows a scatter plot with the Purity index on the y-axis and $\delta^{30}\text{Si}$ (‰) on the x-axis. A red dashed line indicates a strong linear correlation with $R^2 = 0.9572$. Different data points are represented by various symbols:

- Orange circles: Composite surface soils
- Black circles: Vertisol profile
- Red circles: Ferralsol profile
- Light blue squares: Suspended load stream
- Light blue diamonds: Suspended load OF
- Light gray diamonds: Clays
- Green triangles: Litter
Figure 4

The figure shows a scatter plot with the following data points:

- **Tectona Grandis**: Blue diamonds
- **Themeda Triandra**: Red squares
- **Terminalia**: Green triangles

The y-axis represents the isotopic ratio of $\delta^{30}\text{Si}$, expressed in per mil (%), while the x-axis represents the dissolution percentage. The plot includes a shaded region indicating the range of variation for the data points.
Figure 5

- **Litter production**: 2200 / 1.25\%
- **Storage woody stem**: 2200 / -0.11\%
- **Infiltration to soil**: 1850 / 1.15\%
- **Plagioclase weathering**: 1300 / 0.21\%
- **Soil leaching**: 704 / 1.60\%
- **Soil ASi dissolution**: ?
- **Primary minerals**: ?
- **Desorption**: ?
- **Deep root uptake**: 0 - 840 / 0.2\%
Figure 6

Litter ash leaching

Discharge (m$^3$/s$^{-1}$)

$\delta^{30}$Si (%)
Table 1: Repartition of primary and secondary Si pools in ferralsol and vertisol profiles of the Mule Hole watershed, with δ³⁰Si of the ferralsol profile (bulk and clay fractions). "% Si" refers to the percentage of total Si in a given pool.

<table>
<thead>
<tr>
<th></th>
<th>δ³⁰Si</th>
<th>2σ</th>
<th>n</th>
<th>clay content</th>
<th>% Si  &lt;2µm</th>
<th>2µm (*)</th>
<th>% Si</th>
<th>% Si</th>
<th>% Si</th>
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<tr>
<td>Fresh gneiss</td>
<td>-0.34</td>
<td>0.08</td>
<td>3</td>
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<td>8</td>
<td>1.29</td>
<td>0.16</td>
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<td>12</td>
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<td>3</td>
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<tr>
<td>200-220 cm</td>
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<td>0.09</td>
<td>3</td>
<td>42</td>
<td>9</td>
<td>1.1</td>
<td>0.34</td>
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</table>

(*): Estimation of Si located in the clay granulometric fraction, assuming this latter contains 21% of Si (data from Violette et al., 2010)

(**): Estimation of Si located in clays from the chemical composition of the soil, assuming Na is located in Na-plagioclase,

K in sericite and Al in Na-plagioclase, sericite and clays and Al<sub>clay</sub> = Al<sub>total</sub> - Alsericite - Al<sub>Na-plagioclase</sub>

(***): Siprim.min. = (Sitotal - Siclairs - ASi)/Sitotal

(#) : Siquartz = Siprim.min. - Sisericite - SiNa-plagioclase
Table 2: ASi contents, chemical composition and $\delta^{30}$Si of the ASi fraction in the soils and suspended matter of Mule Hole and $\delta^{30}$Si of the litter.

<table>
<thead>
<tr>
<th>Sample</th>
<th>ASi (wt %)</th>
<th>ASi composition (%)</th>
<th>purity index</th>
<th>ASi composition (%)</th>
<th>$\delta^{30}$Si (‰)</th>
<th>$\sigma$</th>
<th>$n$</th>
<th>$\delta^{30}$Si (‰) **</th>
</tr>
</thead>
<tbody>
<tr>
<td>composite Ferralsol on gneiss (0-15cm)</td>
<td>0.36</td>
<td>37</td>
<td>31</td>
<td>1.6</td>
<td>0.0</td>
<td>0.1</td>
<td>0.5</td>
<td>0.88</td>
</tr>
<tr>
<td>composite Vertisol on amphibolite (0-15cm)</td>
<td>0.58</td>
<td>40</td>
<td>1.3</td>
<td>1.4</td>
<td>0.0</td>
<td>0.5</td>
<td>0.4</td>
<td>0.93</td>
</tr>
<tr>
<td>composite shallow Ferralsol on gneiss (0-15cm)</td>
<td>0.45</td>
<td>45</td>
<td>1.9</td>
<td>0.5</td>
<td>0.0</td>
<td>0.0</td>
<td>0.6</td>
<td>0.94</td>
</tr>
<tr>
<td>composite Vertisol on gneiss (0-15cm)</td>
<td>0.55 (†)</td>
<td>48</td>
<td>1.5</td>
<td>0.6</td>
<td>0.1</td>
<td>0.1</td>
<td>0.5</td>
<td>0.95</td>
</tr>
<tr>
<td>Vertisol (0-20cm)</td>
<td>0.86 (†)</td>
<td>40</td>
<td>1.2</td>
<td>0.3</td>
<td>0.2</td>
<td>0.1</td>
<td>0.4</td>
<td>0.95</td>
</tr>
<tr>
<td>Vertisol (40-60cm)</td>
<td>0.42 (†)</td>
<td>49</td>
<td>2.1</td>
<td>0.3</td>
<td>0.3</td>
<td>0.1</td>
<td>0.5</td>
<td>0.94</td>
</tr>
<tr>
<td>Vertisol (100-120cm)</td>
<td>0.06 (†)</td>
<td>37</td>
<td>5.0</td>
<td>2.0</td>
<td>0.3</td>
<td>0.4</td>
<td>0.9</td>
<td>0.80</td>
</tr>
<tr>
<td>Vertisol (140-160cm)</td>
<td>0.14 (†)</td>
<td>28</td>
<td>2.9</td>
<td>1.8</td>
<td>0.1</td>
<td>0.2</td>
<td>0.5</td>
<td>0.84</td>
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<tr>
<td>Ferralsol (0-20cm)</td>
<td>0.24 (†)</td>
<td>33</td>
<td>2.9</td>
<td>1.3</td>
<td>0.0</td>
<td>0.0</td>
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<td>Ferralsol (40-60cm)</td>
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<td>1.8</td>
<td>0.8</td>
<td>0.0</td>
<td>0.0</td>
<td>0.6</td>
<td>0.92</td>
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<td>Ferralsol (100-120cm)</td>
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<td>23</td>
<td>10.4</td>
<td>6.0</td>
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<td>0.3</td>
<td>1.1</td>
<td>0.36</td>
</tr>
<tr>
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<td>0.12</td>
<td>26</td>
<td>8.3</td>
<td>4.9</td>
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<td>0.2</td>
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<td>Ferralsol (200-220cm)</td>
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<td>0.2</td>
<td>1.1</td>
<td>0.42</td>
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<tr>
<td>Suspended matter stream 3212-3224</td>
<td>0.52</td>
<td>36</td>
<td>2.1</td>
<td>1.0</td>
<td>0.1</td>
<td>0.1</td>
<td>0.4</td>
<td>0.91</td>
</tr>
<tr>
<td>Suspended matter stream 3316-3328</td>
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<td>36</td>
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<td>1.5</td>
<td>0.1</td>
<td>0.2</td>
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<tr>
<td>Suspended matter overlandflow 4056</td>
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<td>0.0</td>
<td>0.9</td>
<td>0.88</td>
</tr>
</tbody>
</table>

**Sample**
- Themeda Triandra (from phytoliths)
- Themeda Triandra (from total digestion)
- Terminalia Alata (from total digestion)
- Tectona Grandis (from total digestion)

(*) ASi purity = (Si-(Al+Fe+Ca+Mg+Mn+K))/Si
(†) data from Meunier et al. (2014)

(litter)

<table>
<thead>
<tr>
<th>Litter Sample</th>
<th>ASi (wt %)</th>
<th>ASi composition (%)</th>
<th>purity index</th>
<th>ASi composition (%)</th>
<th>$\delta^{30}$Si (‰)</th>
<th>$\sigma$</th>
<th>$n$</th>
<th>$\delta^{30}$Si (‰) **</th>
</tr>
</thead>
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<tr>
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<td>Themeda Triandra</td>
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<td>Tectona Grandis</td>
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</table>
Table 3: Silica content and $\delta^{30}$Si in the water compartments of Mule Hole.

<table>
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<tr>
<th>sampling date</th>
<th>discharge</th>
<th>Si</th>
<th>$\delta^{30}$Si</th>
<th>2$\sigma$</th>
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<td><strong>Stream</strong></td>
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<td>2004SS1</td>
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<td>1.20</td>
<td>270</td>
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<td><strong>Overland flow</strong></td>
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<td>on Ferralsol</td>
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<td>193</td>
<td>1.03</td>
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<tr>
<td>on Ferralsol</td>
<td>05/10/09</td>
<td>164</td>
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<td>16/10/09</td>
<td>223</td>
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<td>21/11/09</td>
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<td><strong>Soil porewater</strong></td>
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<td>Vertisol 1</td>
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<td>Vertisol 2</td>
<td>15/06/04</td>
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<td>15/06/04</td>
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<td>1.61</td>
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<tr>
<td><strong>Groundwater</strong></td>
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