How big is the influence of biogenic silicon pools on short-term changes of water soluble silicon in soils? Implications from a study of a ten-year-old plant-soil-system

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Abstract

The significance of biogenic silicon (BSi) pools as a key factor for the control of Si fluxes from terrestrial to aquatic ecosystems has been recognized since decades. However, while most research has been focused on phytogenic Si pools, knowledge on other BSi pools is still limited. We hypothesized different BSi pools to influence short-term changes of the water soluble Si fraction in soils to different extents. To test our hypothesis we took plant \textit{(Calamagrostis epigejos, Phragmites australis)} and soil samples in an artificial catchment in a post-mining landscape in the state of Brandenburg, Germany. We quantified phytogenic
(phytoliths), protistic (diatom frustules and testate amoeba shells) and zoogenic (sponge spicules) Si pools as well as Tiron extractable and water soluble Si fractions in soils at the beginning (t₀) and after ten years (t₁₀) of ecosystem development. As expected the results of Tiron extraction showed, that there are no consistent changes of the amorphous Si pool at ‘Chicken Creek’ as early as after ten years. In contrast, compared to t₀ we found increased water soluble Si and BSi pools at t₁₀, thus we concluded BSi pools to be the main driver of short-term changes of water soluble Si. However, because total BSi represents only small proportions of water soluble Si at t₀ (<2 %) and t₁₀ (2.8-4.3 %) we further concluded smaller (<5 µm) and/or fragile phytogenic Si structures to have the biggest impact on short-term changes of water soluble Si. In this context, extracted phytoliths (>5 µm) only amounted to about 16 % of total Si contents of plant materials of C. epigejos and P. australis at t₁₀, thus about 84 % of small-scale and/or fragile phytogenic Si are not quantified by the used phytolith extraction method. Analyses of small-scale and fragile phytogenic Si structures are urgently needed in future work as they seem to represent the biggest and most reactive Si pool in soils, thus the most important driver of Si cycling in terrestrial biogeosystems.

**Keywords**

biosilicification, initial biogeosystem, phytogenic Si, protistic Si, zoogenic Si
1. Introduction

Various prokaryotes and eukaryotes are able to synthesize hydrated amorphous silica (\(\text{SiO}_2 \cdot n\text{H}_2\text{O}\)) structures from monomeric silicic acid (\(\text{H}_4\text{SiO}_4\)), a process called biosilicification (Ehrlich et al. 2010). In terrestrial biogeochemical systems biogenic silicon (BSi) synthesized by bacteria and fungi, plants, diatoms, testate amoebae and sponges can be found forming corresponding microbial, phytogenic, protophytic, protozoic and zoogenic BSi pools, respectively (Puppe et al. 2015, Sommer et al. 2006). BSi has been recognized as a key factor for the control of Si fluxes from terrestrial to aquatic ecosystems as it is in general more soluble compared to silicate minerals (e.g., Fraysse et al. 2006, 2009). These fluxes influence marine diatom production on a global scale (Dürr et al. 2011, Sommer et al. 2006, Struyf & Conley 2012). Marine diatoms in turn can fix large quantities of carbon dioxide via photosynthesis because up to 54% of the biomass in the oceans is represented by diatoms, thus diatoms have an important influence on climate change (Tréguer & De La Rocha 2013, Tréguer & Pondaven 2000).

While the importance of phytogenic Si pools for global Si fluxes has been recognized for three decades (e.g., Bartoli 1983, Meunier et al. 1999, Street-Perrott & Barker 2008), information on the other BSi pools is comparably rare (Clarke 2003). However, in recent publications the potential importance of diatoms, testate amoebae and sponge spicules in soils for Si cycling has been highlighted (Aoki et al. 2007, Creevy et al. 2016, Puppe et al. 2014, 2015, 2016). Furthermore, evidence arises that BSi pools are in disequilibrium at decadal time scales due to disturbances and perturbations by humans, e.g., by changes in forest management or farming practices (Barão et al. 2014, Keller et al. 2012, Vandevenne et al. 2015). In consequence, BSi accumulation and BSi dissolution are not balanced, which influences Si cycling in terrestrial biogeochemical systems not only on decadal but also on millennial...
Sommer et al. (2013), for example, found the successive dissolving of a relict phytogenic Si pool to be the main source of dissolved Si in soils of a forested biogeosystem. Due to the fact that the continuous decomposition of this relict phytogenic Si pool is not compensated by an equivalent buildup by recent vegetation the authors concluded a BSi disequilibrium on a decadal scale. On a millennial scale Clymans et al. (2011) estimated the total amorphous Si storage in temperate soils to be decreased by approximately 10% since the onset of agricultural development about 5,000 years ago. This decrease not only has consequences for land-ocean Si fluxes but also influences agricultural used landscapes because Si is a beneficial element for many crops (e.g., Epstein 2009, Ma & Yamaji 2008).

For a better understanding of BSi dynamics, chronosequence studies are well suited, because they allow us to analyze time-related changes of BSi pools during biogeosystem development. In the present study we analyzed various BSi pools in differently aged soils of an initial artificial catchment (‘Chicken Creek’) in a post-mining landscape in NE Germany. Chicken Creek represents a study site with defined initial conditions and offers the rare opportunity to monitor BSi dynamics from the very beginning. Former studies at this site revealed i) a formation of protophytic (diatom frustules), protozoic (testate amoeba shells) and zoogenic (sponge spicules) Si pools within a short time (<10 years) and ii) a strong relation of spatiotemporal changes of protistic (diatoms and testate amoebae) BSi pools to the vegetation, because plants provide, e.g., rhizospheric micro-habitats including enhanced food supply (Puppe et al. 2014, 2016). From these results it can be concluded that especially vegetated spots at initial biogeosystem sites represent hot spots of BSi accumulation of various origin (compare Wanner & Elmer 2009). Furthermore, construction work with large machines resulted in differently structured sections of Chicken Creek with slight differences.
in abiotic conditions (for details see subsection 2.1.) (Gerwin et al. 2010). These differences in turn lead to section-specific vegetation dynamics at Chicken Creek (Zaplata et al. 2010).

Knowledge about BSi accumulation dynamics is crucial for the understanding of Si cycling in terrestrial biogeoecosystems. We regard water extractable Si as an useful proxy for desilication and biological uptake (plants, testate amoebae etc.). In addition, we used an alkaline extractant (Tiron) to detect eventual short-term changes of the amorphous Si fraction. We hypothesized i) BSi pools to influence short-term changes of water soluble Si in initial soils, but no short-term changes in amorphous Si fractions, ii) the phytogenic Si pool to be the most prominent one in size, thus the biggest driver of short-term changes of water soluble Si, and iii) BSi pool changes to be section-specific, i.e., related to vegetation dynamics. The aims of the present study were i) to quantify various BSi pools, i.e., protophytic, protozoic, zoogenic and phytogenic Si pools, during initial soil and ecosystem development, (ii) to analyze potential section-specific short-term changes of these BSi pools after a decade of ecosystem development, and iii) to evaluate the influence of different BSi pools on water soluble Si in these soils.

2. Material and methods

2.1. Study site

The study site Chicken Creek (51°36′18″ N, 14°15′58″ E) represents an artificial catchment in a post-mining landscape located in the active mining area ‘Welzow-South’ (lignite open-cast mining, 150 km south-east of Berlin) in the state of Brandenburg, Germany (Kendzia et al. 2008, Russell et al. 2010). Climate at Chicken Creek is characterized by an average air temperature of 9.6°C and an annual precipitation of 568 mm comprising data from 1981 to 2010 (Meteorological Station Cottbus, German Weather Service).
For construction of the about 6 ha sized catchment an 1-3 m thick base layer (aquiclude) of Tertiary clay was covered by a 2-3 m thick sandy, lignite- and pyrite-free Quaternary sediment serving as water storage layer (aquifer) (Gerwin et al. 2010, Kendzia et al. 2008). Quaternary material was taken from a depth of 20-30 m during lignite mining process and its texture is classified as sand to loamy sand (Table 1) with low contents of carbonate (Gerwin et al. 2009, 2010, Russell et al. 2010). Dumping of material and construction work with large machines (e.g., stackers and bulldozers) resulted in differently structured sections of Chicken Creek. Generally, the catchment area can be divided into four sections: i) an eastern part (ca. 1.8 ha), ii) a western part (ca. 1.6 ha), iii) a central trench (ca. 0.9 ha) separating the eastern from the western part and iv) a southern part (ca. 1.5 ha) with a pond at the lowest point (Fig. 1). Construction work was completed in September 2005 (time zero, t₀). Analyses subsequent to catchment completion indicated slight differences in abiotic conditions (soil pH, conductivity, skeleton content (soil particle diameter >2 mm), proportions of sand, silt and clay, concentration of organic and inorganic carbon) between the eastern and the western part (Gerwin et al. 2010). The primary mineral component in all particle size fractions at t₀ was quartz (only small amounts of K-feldspar, plagioclase). Calcite comprised 0.5-4.5 % of the initial sediment, dolomite was only detectable in few samples with contents of 0.5 % and magnesite (MgCO₃) was not detectable by mineralogical analysis (W. Schaaf, pers. comm., 2011). For detailed information on site construction and initial ecosystem development see Gerwin et al. (2010) and Schaaf et al. (2010), respectively.

2.2. Soil sampling

We took samples shortly after construction of Chicken Creek (2005, t₀) and after an ecosystem development period of about ten years (2015, t₁₀). For t₀ (no vegetation
detectable) we assumed only very few biogenic siliceous structures homogenously distributed on the whole area of Chicken Creek, i.e., no section-specific distribution of BSi (BSi_0 east ≈ BSi_0 west ≈ BSi_0 south) at the beginning of ecosystem development (Puppe et al. 2016). This is why we did not sample all different sections of the catchment, but took soil samples in six field replicates to quantify BSi pools at t_0. However, for t_{10} we hypothesized section-specific differences in BSi pool quantities related to section-specific vegetation dynamics. To evaluate these differences after a decade of ecosystem development and to cover the biggest possible BSi accumulation in soil we focused on spots where Si accumulating plant species, i.e., *Calamagrostis epigejos* and *Phragmites australis* became dominant (Zaplata et al. 2010). Thus we took samples in the eastern (C. *epigejos* dominant) and western (mainly C. *epigejos* dominant, one spot with *P. australis*) and southern section (P. *australis* dominant) of Chicken Creek.

For an accurate description of changes of abiotic soil conditions and related phytogenic Si in every section we took soil and plant samples in eastern, western and southern sections at t_0 as well as t_{10}. Erosion and deposition processes were clearly evident in the Chicken Creek catchment during the first years without plant cover. Substantial surface changes resulted from rill erosion as aerial photographs (rill network) and a comparison of photogrammetry-based digital elevation models showed (Schneider et al. 2013). Interrill erosion did not lead to surface changes larger than about 20 cm during the first five years. Afterwards the establishment of an area-wide plant cover substantially reduces interrill erosion. Because all soil data at t_0 referred to a depth increment of 30 cm we reasonably assumed the same soil conditions for the sampled t_0-spots during the first years. Furthermore, we carefully selected sampling points at t_{10} to be not influenced by erosion, i.e., at spots with low surface roughness and outside rills. Soil samples for the determination of soil properties and plant
samples were taken in five (western and southern section) and six (eastern section) field replicates at \( t_0 \) and \( t_{10} \) (Fig. 1). At every sampling point three undisturbed soil cores where taken with a core cutter (diameter = 3.4 cm, depth = 5 cm) and transferred into plastic bags. Bulk densities were calculated from dividing weight of dried (105°C) soil samples by corresponding volume.

2.3. Determination of basic soil properties

Soil samples were air dried and sieved and the fine earth fraction (<2 mm) was used for laboratory analyses. Soil pH was measured based on the DIN ISO Method 10390 (1997) in 0.01 M \( \text{CaCl}_2 \) suspensions at a soil to solution ratio of 1:5 (w/v) after a 60 minute equilibration period using a glass electrode. The total carbon content was analyzed by dry combustion using an elemental analyzer (Vario EL, Elementar Analysensysteme, Hanau, Germany). Carbonate (\( \text{CaCO}_3 \)) was determined conductometrically using the Scheibler apparatus (Schlichting et al. 1995). Organic carbon (\( C_{\text{org}} \)) was computed as the difference between total carbon and carbonate carbon. Analyses of basic soil properties were performed in two lab replicates per sample.

2.3.1. Water Extractable Si (\( Si_{H2O} \))

Water extractable Si was determined based on a method developed by Schachtschabel & Heinemann (1967). Ten grams of dry soil (<2 mm) were weighed into 80 mL centrifuge tubes and 50 mL distilled water added together with three drops of a 0.1% \( \text{NaN}_3 \)-solution to prevent microbial activity. Total extraction time was seven days in which tubes were shaken by hand twice a day for twenty seconds. Mechanical (constant) shaking by using, e.g., a roll mixer, was avoided to prevent abrasion of mineral particles colliding during shaking.
(McKeague & Cline 1963). The solutions were centrifuged (4000 rpm, 20 min), filtrated (0.45 µm polyamide membrane filters) and Si was measured by ICP-OES, (ICP-iCAP 6300 DUO, Thermo Fisher SCIENTIFIC GmbH). Analyses of water extractable Si were performed in two lab replicates per sample.

2.3.2. Tiron extractable Si ($Si_{Tiron}$), aluminum ($Al_{Tiron}$) and iron ($Fe_{Tiron}$)

The Tiron ($C_6H_4Na_2O_8S_2\cdotH_2O$) extraction followed the method developed by Biermans & Baert (1977), modified by Kodama & Ross (1991). It has been used to quantify amorphous biogenic and pedogenic Si (Kendrick & Graham 2004), although a partial dissolution of primary minerals is well known (Kodama & Ross 1991, Sauer et al. 2006). The extraction solution was produced by dilution of 31.42 g Tiron with 800 mL of distilled water, followed by addition of 100 mL sodium carbonate solution (5.3 g Na$_2$CO$_3$ + 100 mL distilled water) under constant stirring. The final pH of 10.5 was reached by adding small volumes of a 4M NaOH-solution. For the extraction 30 mg of dry soil were weighed into 80 mL centrifuge tubes and a 30 mL aliquot of the Tiron solution was added. The tubes were then heated at 80°C in a water bath for 1h. The extracted solutions were centrifuged at 4000 rpm for 30 min, filtrated (0.45 µm polyamide membrane filters, Whatman NL 17) and Si, Al and Fe measured by ICP-OES. Analyses of Tiron extractable Si, Al and Fe were performed in three lab replicates per sample.

2.4. Microscopical analyses of diatoms, sponge spicules and testate amoebae

Fresh soil samples were homogenized by gentle turning of the plastic bags before air drying. Afterwards 2 g of fresh soil were taken per sample and stored in 8 mL of formalin (4 %). Subsequently, biogenic siliceous structures, i.e., diatom frustules, testate amoeba shells and
sponge spicules (Fig. 2A-D), were enumerated in soil suspensions (125 mg fresh mass (FM)) received from serial dilution (1000-125 mg soil in 8 mL of water each) using an inverted microscope (OPTIKA XDS-2, objectives 20:1 and 40:1, equipped with a digital camera OPTIKAM B9).

2.5. Determination of phytoliths in soil samples

10 g of dry soil material (<2 mm) were processed in four steps (adapted from Alexandre et al. 1997). First organic matter was oxidized using H₂O₂ (30 Vol. %), HNO₃ (65 Vol. %) and HClO₄ (70 Vol. %) at 80°C until reaction subsides. Secondly, carbonates and Fe oxides were dissolved by boiling the sample in HCl (10 Vol. %) for 30 min. Thirdly, the <2 μm granulometric fraction was removed by dispersion of the remaining solid phase of step 2 with 2 Vol. % sodium hexametaphosphate solution (6–12 h), centrifugation at 1000 rpm for 2–3 min, and subsequent decantation. Finally, the phytoliths were separated by shaking the remaining solid phase of step 3 with 30 mL of sodium polytungstate (Na₆(H₂W₁₂O₄₀)·H₂O) with a density of 2.3 g cm⁻³ and subsequent centrifugation at 3000 rpm for 10 min. Afterwards, the supernatant was carefully pipetted and filtered using 5 μm teflon filters. This step was repeated three times. The filter residue was washed with water, bulked, dried at 105°C, and weighted.

2.6. Quantification of biogenic Si pools

In general, biogenic siliceous structures consist of hydrated amorphous silica (SiO₂·nH₂O). We assumed an average water content of about 10 % for these structures to avoid an overestimation of BSi pools (Mortlock & Froelich 1989).
Protophytic Si pools (represented by diatom frustules) were quantified by multiplication of Si contents per frustule with corresponding individual numbers (see Puppe et al. 2016). Protozoic Si pools (represented by testate amoebae) were quantified by multiplication of silica contents of diverse testate amoeba taxa (Aoki et al. 2007) with corresponding individual numbers (living plus dead individuals, for details see Puppe et al. 2014, 2015). Zoogenic Si pools (represented by sponge spicule fragments) were calculated by multiplying volumes (µm$^3$) of the found spicule fragments with the density of biogenic Si (2.35 g cm$^3$) and summing up the results. Volume measurements were conducted using a laser scanning microscope (Keyence VK-X110, magnification 200-2.000x) (details in Puppe et al. 2016). For laser scanning microscopy spicule fragments were taken from soil suspensions by micromanipulation, washed in dist. H$_2$O and placed on clean object slides. After air drying images of spicule fragments were acquired (software Keyence VK-H1XVD) and analyzed (software Keyence VK-H1XAD).

Phytogenic Si pools were estimated by multiplying the numbers of found phytoliths with corresponding mean volumes (µm$^3$) of phytoliths, multiplying these results with the density of biogenic Si (2.35 g cm$^3$) and summing up the results. Volume measurements with the laser scanning microscope of 30 typical elongate (Fig. 2E) and 30 typical bilobate phytoliths (Fig. 2F) resulted in mean volumes of 3765 µm$^3$ and 707 µm$^3$, respectively. For laser scanning microscopy extracted phytoliths were placed on clean object slides and images were acquired and analyzed analogous to sponge spicules. For bilobate phytoliths we measured the upper half per phytolith and doubled the result to obtain the corresponding total volume, thus we assumed bilobate phytoliths to be symmetric. We assumed phytoliths to consist of 95 % SiO$_2$ and 5 % other elements, i.e., carbon (Song et al. 2012) and other elements like iron, aluminum or calcium (Buján 2013).
BSi pools (mg m$^{-2}$) were calculated considering bulk density (g cm$^{-3}$), thickness (5 cm) and – for protistic and zoogenic Si pools – water content (% of fresh mass) per soil sample. Silica (M = 60.08 g mol$^{-1}$) pools were converted to Si (M = 28.085 g mol$^{-1}$) pools by multiplication with 28/60 (details in Puppe et al. 2014, 2015, 2016).

2.7. Plant analyses

Plant and litter samples of *C. epigejos* and *P. australis* were collected in the summer of 2015. In general, monomeric silicic acid (H$_4$SiO$_4$) enters the plant via its roots and is carried in the transpiration stream towards transpiration termini. When water evaporates, silicic acid becomes supersaturated and is precipitated as hydrated silica in the form of phytoliths. The vast majority of Si in plants is located at the transpiration termini (e.g., leaves) in the aerial plant parts, while considerably less Si can be found in other plant portions like stems, roots and rhizomes. Sangster (1983), for example, found no significant Si depositions in rhizomes of *P. australis*. Consequently, we only analyzed the aboveground vegetation (including transpiration termini and stems). The collected plant material was washed with distilled water to remove adhering soil minerals and oven-dried at 45°C for 48 hours.

2.7.1. Total Si content in plant materials

Plant samples were milled using a knife mill (Grindomix GM 200, Retsch) in two steps: 4,000 rpm for 1 min and then 10,000 rpm for 3 min. Sample aliquots of approximately 100 mg were digested under pressure in PFA digestion vessels using a mixture of 4 mL distilled water, 5 mL nitric acid (65 %), and 1 mL hydrofluoric acid (40 %) at 190°C using a microwave digestion system (Mars 6, CEM). A second digestion step was used to neutralize the hydrofluoric acid with 10 mL of a 4 %-boric acid solution at 150°C. Silicon was measured by
ICP-OES (ICP-iCAP 6300 Duo, Thermo Fisher Scientific GmbH) with an internal standard. To avoid contamination, plastic equipment was used during the complete procedure. Analyses of total Si content were performed in three lab replicates per sample.

2.7.2. Determination of phytoliths in plants and litter

Plant material was washed with distilled water and oven-dried at 45°C for 48 hours. Removal of organic matter was conducted by burning the samples in a muffle furnace at 450°C for 12 hours. Next, the material was subject to additional oxidation using 30 % H₂O₂ for 12 hours. The obtained material was filtered through a teflon filter with a mesh size of 5 µm. The isolated phytoliths and siliceous cast (>5 µm) were subject to analysis via polarized light microscopy (Nikon ECLIPSE LV100 microscope) for full characteristics. We used laser scanning microscopy for measurements of the surface-area (µm²) of the 30 typical bilobate and 30 typical elongated phytoliths used for volume measurements (see 2.6) and calculated corresponding surface-area-to-volume ratios (A/V ratios) as an indicator for the resistibility of these siliceous structures against dissolution. Higher A/V ratios indicate a bigger surface-area available for dissolution processes.

2.8. Statistical analyses

Correlations were analyzed using Spearman’s rank correlation (rₜ). Significances in two-sample (n = 2) cases were verified with the Mann-Whitney U-test. For k-sample (n > 2) cases the Kruskal-Wallis analysis of variance (ANOVA) was used followed by pairwise multiple comparisons (Dunn’s post hoc test). Statistical analyses were performed using software package SPSS Statistics (version 19.0.0.1, IBM Corp.).
3. Results

3.1. Basic soil parameters

Soils at the initial state (t_0) showed in the upper 5 cm organic carbon contents (C_{org}) between 1.1 g kg^{-1} and 4.4 g kg^{-1} in the western section, 0.8 g kg^{-1} and 1.8 g kg^{-1} in the eastern section and 0.2 g kg^{-1} and 3.3 g kg^{-1} in the southern section. This corresponded to mean carbon stocks of 237 g m^{-2} (west), 123 g m^{-2} (east) and 160 g m^{-2} (south, Table 2). After 10 years (t_{10}) of ecosystem development the C_{org} stocks increased up to a factor of 3 (396-556 g m^{-2} in the upper 5 cm) compared to corresponding values at t_0. This resulted in a surprisingly high mean annual CO_{2}-C sequestration rate of 27-32 g m^{-2} (upper 5 cm). Hereby the largest C_{org} stock changes were found in the western section of the area followed by the eastern section and the southern section (Table 2).

The carbonate contents (CaCO_3) at t_0 varied between means of 1.0 g kg^{-1} (west), 0.9 g kg^{-1} (east) and 1.8 g kg^{-1} (south). The corresponding stocks were 88 g m^{-2} (west), 91 g m^{-2} (east) and 174 g m^{-2} (south, Table 2). The carbonate pools in the western and eastern section were very similar, while the high carbonate values in the southern section were due to the original soil properties. At t_{10} the distribution of carbonate was as follows: in the western section there was an increase of about 17 % (from 88 g m^{-2} to 101 g m^{-2}), in the eastern part a distinct decrease of about 67 % (from 91 g m^{-2} to 30 g m^{-2}) was detected and in the southern section again a decrease of about 28 % (from 174 g m^{-2} to 126 g m^{-2}) was identified.

At t_0 the pH values of the soils showed a range between 7.9 and 8.3 (Table 2) with relatively low variation between the different sections. After 10 years the pH values decreased to 7.1-7.4 in all sections.
3.2. Water and Tiron extractions

The mean water soluble Si (Si$_{\text{H}_2\text{O}}$) contents in the upper 5 cm showed low variation between the different sections at $t_0$: 7.3 mg kg$^{-1}$ (west), 7.2 mg kg$^{-1}$ (east) and 8.6 mg kg$^{-1}$ (south). The corresponding stock values were 0.7 g m$^{-2}$ (west), 0.87 g m$^{-2}$ (east) and 0.84 g m$^{-2}$ (south) for all sections at $t_0$ (Table 2). After 10 years ($t_{10}$) an overall significant increase of Si$_{\text{H}_2\text{O}}$ in each of the different sections compared to $t_0$ was found. The corresponding stock values were 1.7 g m$^{-2}$ (west), 1.5 g m$^{-2}$ (east) and 2.2 g m$^{-2}$ (south, Table 2).

At $t_0$ the mean Tiron extractable Si contents in the upper 5 cm varied between 5.5 g kg$^{-1}$ (west), 5.2 g kg$^{-1}$ (east) and 4.1 g kg$^{-1}$ (south). The related stock values were 524 g m$^{-2}$ (west), 503 g m$^{-2}$ (east) and 399 g m$^{-2}$ (south, Table 2). After 10 years ($t_{10}$) the Tiron extractable Si content showed a slight increase in the western section to 6.5 g kg$^{-1}$ (552 g m$^{-2}$), while the concentration in the eastern section decreased significantly to 2.6 g kg$^{-1}$ (196 g m$^{-2}$, Table 2).

In the southern section only a slight decrease to 3.8 g kg$^{-1}$ (317 g m$^{-2}$) was found. The Al and Fe extractable Tiron contents followed the distribution of the Si concentrations with one exception in the western section, where contrary to Si the Al and the Fe contents slightly increased at $t_{10}$ (Table 2). Si/Al ratios ranged between 1.6 and 2.2 at Chicken Creek. Tiron extractable Si and Al fractions as well as Tiron extractable Al and Fe fractions were strongly correlated (Table 3).

3.3. Biogenic Si pools in soils

In general, total biogenic Si pools increased in every section after ten years of ecosystem development with statistically significant differences between $t_0$ (11.6 ± 6.5 mg Si m$^{-2}$) and the southern section at $t_{10}$ (96.0 ± 87.2 mg Si m$^{-2}$) (Fig. 3). Total BSi showed strong positive and statistically significant correlations to water soluble Si (Table 3). Phytogenic (phytoliths
>5 µm) Si pools ranged from 0-18 mg m\(^{-2}\) (mean: 6.6 mg m\(^{-2}\)) at \(t_0\) and significantly increased to means of 20.7 mg m\(^{-2}\) (range: 7-52 mg m\(^{-2}\)) and 12.9 mg m\(^{-2}\) (range: 14-15 mg m\(^{-2}\)) at the eastern and southern section during 10 years, respectively (Fig. 4A). Protophytic Si pools (diatom frustules) ranged from 0-7 mg m\(^{-2}\) (mean: 2.6 mg m\(^{-2}\)) at \(t_0\) and increased up to a mean of 47.4 mg m\(^{-2}\) (range: 0.1-162 mg m\(^{-2}\)) at \(t_{10}\) (southern section) (Fig. 4B). At \(t_0\) no sponge spicules were found with one exception representing an extreme value (12.7 mg m\(^{-2}\)). After one decade of ecosystem development zoogenic Si pools increased to a maximum of 46 mg m\(^{-2}\) at the southern section (\(t_{10}\)) (Fig. 4C). Protozoic Si pools were zero at \(t_0\) with one exception representing an extreme value (1.8 mg m\(^{-2}\)) and significantly increased to 4.6 mg m\(^{-2}\) (range: 1-11 mg m\(^{-2}\)) and 11.5 mg m\(^{-2}\) (range: 2-36 mg m\(^{-2}\)) in the eastern and the southern section at \(t_{10}\), respectively (Fig. 4D).

At \(t_0\) most BSi (>50 %) is represented by phytoliths >5 µm followed by diatom frustules, sponge spicules and testate amoeba shells (Fig. 5). After ten years of ecosystem development the proportion of the different BSi pools to total BSi changed. While the proportion of protozoic Si pools increased in all sections at \(t_{10}\), the other BSi pools showed more variable changes over time. The proportion of phytogenic Si pools either increased (western section) or decreased (eastern and southern sections). In contrast, the proportion of protophytic Si pools decreased at the western section and increased in the eastern and southern sections. The proportion of zoogenic Si pools decreased in the western and eastern sections, but increased slightly in the southern section at \(t_{10}\).

### 3.4. Phytoliths and total Si content in plant materials

The total content of Si was determined for two Si accumulating plant species *Calamagrostis epigejos* and *Phragmites australis* dominating distinct catchment sections. For *C. epigejos*...
the mean total content of Si was 2.25 % (range: 1.8-3.1 %), whereas for *P. australis* a mean total Si content of 2.70 % (range: 2.0-3.2 %) was determined (Fig. 6A, B). For litter we found mean total Si contents of 3.1 % (range: 2.8-3.3 %) and 2.9 % (range: 1.7-3.2 %) for *C. epigejos* and *P. australis*, respectively.

Phytoliths >5 µm were also isolated from both plants; for *C. epigejos* the mean extracted phytolith content was 0.37 % (range: 0.31-0.46 %), whereas for *P. australis* a mean phytolith content of 0.43 % (range: 0.37-0.50 %) was determined (Fig. 6A, B), i.e., related to the total Si content of plants 16.4 % and 15.9 % of phytogenic Si were represented by extracted phytoliths >5 µm in *C. epigejos* and *P. australis*, respectively. Thus, small-scale (<5 µm) and/or fragile (siliceous structures mostly thinner than 5 µm, but up to several hundred micrometers long, Fig. 7) phytogenic Si represented 83.6 % and 84.1 % of total phytogenic Si in *C. epigejos* and *P. australis*, respectively. Mean extracted phytolith contents in plant litter were 0.47 % (range: 0.35-0.70 %) and 0.51 % (range: 0.41-0.59 %) for *C. epigejos* and *P. australis*, respectively.

Surface-areas of 30 typical bilobate and 30 typical elongate phytoliths were in a range of 216 µm² to 3,730 µm² and 2,302 µm² to 22,203 µm², respectively (Table 4). The corresponding volumes of bilobate and elongate phytoliths were in a range of 36 µm³ to 2,046 µm³ and 390 µm³ to 14,649 µm³, respectively. Surface-to-volume ratios of bilobate and elongate phytoliths were in a range of 0.7 to 9.8 and 0.6 to 5.9 with means of 2.8 and 2.6, respectively.

3.5. **BSi and Si fractions under Calamagrostis epigejos and Phragmites australis**

Water soluble Si fractions increased by 99 % and 163 %, total BSi by 281 % and 660 % after ten years of ecosystem development in soils under *C. epigejos* and *P. australis*, respectively.
In contrast, Si\textsubscript{Tiron} decreased by 42\% and 1.4\% from t\textsubscript{0} to t\textsubscript{10} in soils under \textit{C. epigejos} and \textit{P. australis}, respectively. If we assume mean dry biomasses of 115 g m\textsuperscript{-2} and 186 g m\textsuperscript{-2} for \textit{C. epigejos} and \textit{P. australis} (M. Wehrhan, pers. comm., 2017) about 2.6 g Si m\textsuperscript{-2} and 5.0 g Si m\textsuperscript{-2} are stored in the aboveground biomass at Chicken Creek at t\textsubscript{10}, respectively. For litter of \textit{C. epigejos} and \textit{P. australis} (mean dry biomasses of 59 g m\textsuperscript{-2} and 94 g m\textsuperscript{-2} at t\textsubscript{10}, M. Wehrhan, pers. comm., 2017) we calculated corresponding pools of about 1.8 g Si m\textsuperscript{-2} and 2.7 g Si m\textsuperscript{-2} at t\textsubscript{10}, respectively.

4. Discussion

4.1. Drivers of short-term changes of water soluble Si at Chicken Creek

In general, weathering of silicates represents the ultimate source of Si(OH)\textsubscript{4} in terrestrial biogeoecosystems in the long term (Berner 2003). In this context, the long-term accumulation of BSi can influence the total amorphous (Tiron extractable) Si as it is known from forested catchments or old chronosequence soils (Conley et al. 2008, Kendrick & Graham 2004, Saccone et al. 2008). Contrary, short-term changes of BSi pools likely do not influence Tiron extractable Si in initial soils (total BSi represents only 0.002-0.03 \% of Tiron extractable Si at Chicken Creek). Thus, the major proportion of Tiron extractable Si at Chicken Creek seems to be of pedogenic origin (e.g., Si included in Al/Fe oxides/hydroxides). This is supported by relatively low Si/Al ratios (<5) indicating a minerogenic origin of Tiron extractable Si instead of BSi as a source of Si\textsubscript{Tiron} (Bartoli & Wilding 1980). We further exclude changes of Tiron extractable Si as the main driver of water soluble Si at Chicken Creek in the short term, because i) Si\textsubscript{Tiron} and Si\textsubscript{H2O} showed no statistical relationship at all and ii) a significant change of the Tiron extractable Si fraction occurred only in the eastern section, whereas in the western and southern section Si\textsubscript{Tiron} did not change significantly over time. We assume that
these changes of $S_{Tiron}$ in the eastern section are related to abiotic conditions (soil pH, conductivity, skeleton content, proportions of sand, silt and clay, concentration of organic and inorganic carbon), which were slightly different to the conditions of the western section already at $t_0$ (Gerwin et al. 2010). Furthermore, we excluded atmospheric inputs as potential drivers of short-term changes of water soluble Si at Chicken Creek. On the one hand, dust depositions (dry deposition) at Chicken Creek are very low (73-230 mg m$^{-2}$ d$^{-1}$) and only slightly above the annual average (70-90 mg m$^{-2}$ d$^{-1}$) measured in the state of Brandenburg (Wanner et al. 2015). On the other hand, the total input of Si (as a lithogenic element) by precipitation (wet deposition) is negligible as well (<1 kg Si ha$^{-1}$ yr$^{-1}$, Sommer et al. 2013).

Our results indicate a strong relationship between water soluble Si and total BSi. In this context, two different causal chains can be discussed: Either $SiO_2$-synthesizing organisms are drivers of the amount of $Si(OH)_4$ in the soil or – *vice versa* – the amount of water soluble Si in the soils is the main driver of $SiO_2$-synthesizing organisms as biosilicification is limited by $Si(OH)_4$. Laboratory studies, for example, revealed that $SiO_2$-synthesizing organisms, i.e., testate amoebae, can deplete the amount of $Si(OH)_4$ in culture media due to biosilicification (Aoki et al. 2007, Wanner et al. 2016). However, Wanner et al. (2016) also showed that culture growth of $SiO_2$-synthesizing testate amoebae was dependent on Si concentration in the culture media. Furthermore, *in situ* analyses showed that marine diatom blooms can deplete $Si(OH)_4$ concentrations in the oceans (Hildebrand 2008). In forested biogeosystems Puppe et al. (2015) found high individual numbers of $SiO_2$-synthesizing testate amoebae at study sites with low amounts of $Si(OH)_4$ and *vice versa*. However, it is unlikely that testate amoebae depleted amounts of $Si(OH)_4$ at these sites, because corresponding protozoic Si pools are relatively small compared to phytogenic ones (Puppe et al. 2015, Sommer et al. 2013). Regarding vegetation and corresponding phytogenic Si pools their influence on the
amount of Si(OH)$_4$ in soils has been shown in several studies (e.g., Bartoli 1983, Farmer et al. 2005, Sommer et al. 2013). On the other hand, phytolith production is probably more influenced by the phylogenetic position of a plant than by environmental factors like temperature or Si availability (Hodson et al. 2005, Cooke & Leishman 2012).

From our results and the discussion above we conclude short-term changes of water soluble Si to be mainly driven by BSi. However, total BSi represents only small proportions of water soluble Si at $t_0$ (<2 %) and $t_{10}$ (<4.5 %). From this result the question arises, where does the major part of the increase in water soluble Si at Chicken Creek come from? We will discuss this question in the subsection (4.2.) below.

### 4.2. Sources of water soluble Si at Chicken Creek

From former results of BSi analyses in forested biogeosystems, we assumed the phytogenic Si pool to be the most prominent in size. In this context, results of Sommer et al. (2013) and Puppe et al. (2015) showed that phytogenic Si pools in soils of forested biogeosystems were up to several hundred times larger than protozoic Si pools. However, phytogenic Si pools in soils are surprisingly small compared to other BSi pools at Chicken Creek. Our findings can be attributed to at least two reasons. Firstly, phytogenic Si is stored in a developing organic litter layer where it is temporarily protected against dissolution and secondly, the used methods were not able to accurately quantify the total phytogenic Si pool, but only the larger (>5 µm) and stable part.

Total Si and phytolith contents of litter samples at Chicken Creek did not differentiate from total Si and phytolith contents of plants. This fact indicates that litter decomposition and related Si release into the subjacent soil are relatively slow processes and we interpret our findings as a hint for a developing compartment of dead plant tissue above the mineral soil.
surface. Esperschütz et al. (2013) showed in a field experiment in initial soils near Chicken Creek that after 30 weeks only 50 % of the litter of C. epigejos were degraded, whereby degradation rates were highest in the first four weeks. Estimations of biomasses of C. epigejos and P. australis at Chicken Creek via remote sensing with an unmanned aerial system showed that the relation between phytogenic Si pools plant biomass and litter biomass is almost the same for both plant species (factor about 1.5, based on the total area of Chicken Creek), i.e., Si in the plants was about one third higher than in litter (M. Wehrhan, pers. comm., 2017, manuscript in preparation). At the sampling points about 1.8 g Si m$^{-2}$ and 2.7 g Si m$^{-2}$ were stored in the litter of C. epigejos and P. australis at $t_{10}$, respectively, which is in the range of published data for annual Si input through litterfall in a short grass steppe (2.2-2.6 g Si m$^{-2}$ per year, Blecker et al. 2006).

Altogether, these results clearly underline our interpretation of a developing organic layer where litter accumulates and phytogenic Si is temporarily stored and protected against dissolution, thus Si release is delayed biologically controlled as it can be observed at forested biogeosystems (Sommer et al. 2013). The Si pools in the aboveground biomass of C. epigejos (2.6 g Si m$^{-2}$) and P. australis (5.0 g Si m$^{-2}$) at Chicken Creek at $t_{10}$ are comparable to reported values of Great Plains grasslands (2.2-6.7 g Si m$^{-2}$ in the aboveground biomass) (Blecker et al. 2006) and reach about 30 % (C. epigejos) or 59 % (P. australis) of published data for a beech forest (8.5 g Si m$^{-2}$ in the aboveground biomass of Fagus sylvatica trees) in northern Brandenburg, Germany (Sommer et al. 2013), after (only) ten years of ecosystem development.

Regarding methodological shortcomings of the used phytolith extraction procedure there are several aspects to be discussed. Wilding & Drees (1971), for example, showed that about 72 % of leaf phytoliths of American beech (Fagus grandifolia) are smaller than 5 µm. This is
in accordance with our findings. Phytoliths >5 µm only amounted to about 16 % of total Si contents of plant materials of *C. epigejos* and *P. australis*, thus about 84 % of phytogenic Si (<5 µm and/or fragile phytogenic Si structures) are not quantified by the used phytolith extraction method. Watteau & Villemin (2001) found even smaller (5-80 nm) spherical grains of pure silica in leaf residues in topsoil samples of a forested biogeosystem. In addition, silica depositions can be found in intercellular spaces or in an extracellular (cuticular) layer (Sangster et al. 2001), whereat no recognizable phytoliths are formed. These structures might be too fragile for preservation in soils and are likely lost to a great extent in the used phytolith extraction procedure due to dissolution. Meunier et al. (2017) analyzed different phytolith morphotypes, e.g, silica bodies originating from cells of the upper epidermis, silica casts of trichomes or parenchyma/collenchyma cells, of durum wheat plant shoots. They found fragile sub-cuticular silica plates (2-4 µm thick, up to several hundred micrometers long and wide) to be the second most common phytolith morphotype. This is corroborated by our own findings as the biggest part (about 84 %) of total plant Si is represented by small-scale (<5 µm) and/or fragile phytogenic Si in *C. epigejos* and *P. australis*. If we assume that total Si contents of plants at Chicken Creek are one-to-one reflected by phytogenic Si pools in soils we can easily calculate these small-scale and fragile pools resulting in about 130 mg m⁻² and 100 mg m⁻² (84 % of total, i.e., 156 mg m⁻² and 119 mg m⁻², phytogenic Si each) under *C. epigejos* and *P. australis*, respectively. These calculated phytogenic Si pools are about 13 (diatom frustules), 38 (testate amoeba shells) and 45 (sponge spicules) or 3 (diatom frustules) and 10 (testate amoeba shells, sponge spicules) times bigger than the other BSi pools at *C. epigejos* and *P. australis* sampling points, respectively. If we further assume an input of this phytogenic Si for at least seven years (Zaplata et al. 2010) phytogenic Si might be the main driver of short-term changes of water soluble Si at Chicken Creek. This
is supported by relatively high surface-to-volume ratios of bilobate and elongate phytoliths. These ratios are about three times higher compared to ratios of other biogenic siliceous structures, i.e., testate amoeba shells, diatom frustules and sponge spicules. In addition, Si pools represented by single siliceous platelets of testate amoeba shells have to be considered as well as these platelets can be frequently found in freshwater sediments, for example (Douglas & Smol 1987, Pienitz et al. 1995). Unfortunately, there is no information on the quantity of such platelet pools in soils available, but it can be assumed that these platelets can be frequently found in soils as they are used by some testate amoeba genera (e.g., Schoenbornia, Heleopera) for shell construction (Meisterfeld 2002, Schönborn et al. 1987). In general, it can be assumed that phytogenic Si structures <5 µm and single testate amoeba platelets (about 3-12 µm in diameter, Douglas & Smol 1987) are highly reactive due to their relatively high surface/volume ratios. However, to the best of our knowledge there is no publication available dealing with corresponding physicochemical analyses or dissolution kinetics of these siliceous structures. In general, experiments with phytoliths (>5 µm) showed that surface-areas and related dissolution susceptibilities are, for example, age-related due to changes in specific surface areas and the presence of organic matter bound to the surface of phytoliths (Fraysse et al. 2006, 2009).

5. Conclusions

Decadal changes of water soluble Si at Chicken Creek are mainly driven by BSi, thus Si cycling is biologically controlled already at the very beginning of ecosystem development. In this context, especially phytogenic Si plays a prominent role. However, a developing organic layer (L horizon) at the soil surface temporarily protects phytogenic Si against dissolution, because phytogenic Si is still incorporated in plant structural elements (tissues).
delaying biogenic Si pool is built up and Si release into the soil is retarded. Furthermore, established phytolith extraction methods alone are not suitable to quantify total phytogenic Si pools as phytoliths >5 µm seem to be only a minor part of this pool (about 16 % in the current study). In general, information on small-scale (<5 µm) and/or fragile phytogenic Si structures are urgently needed as they seem to represent the biggest and most reactive Si pool in soils, thus the most important driver of Si cycling in terrestrial biogeochemical systems. Future work should focus on i) the quantification of this pool, ii) physicochemical analyses of its components, and (iii) their dissolution kinetics in lab experiments. The combination of modern microscopical (SEM-EDX, laser scanning microscopy) (this study, Puppe et al. 2016, Sommer et al. 2013) and spectroscopical (FTIR and micro-FTIR spectroscopy) (Liu et al. 2013, Loucaides et al. 2010, Rosén et al. 2010) methods might introduce new insights in this field.

Acknowledgements

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Historical land use change has lowered terrestrial silica mobilization. *Nature Communications* 1, 129.


Fig. 1. Map of Chicken Creek (W = western section, CT = central trench, E = eastern section, S = southern section with pond). Triangles indicate the sampling points used for BSi analyses at $t_0$ (n = 6). Circles indicate the sampling points used for measurements of soil parameters (at $t_0$ and $t_{10}$) and plant analyses (only at $t_{10}$) (W, n = 5; E, n = 6; S, n = 5). Empty and filled circles represent sampling points where *Calamagrostis epigejos* and *Phragmites australis* became dominant. Note that the size of sampling points is not to scale.
**Fig. 2.** Micrographs (light microscope) of biogenic silica structures found at Chicken Creek. A) pennate diatom (valve view), B) testate amoeba shell (*Euglypha cristata*), C) and D) sponge spicules (fragments), E) elongate phytolith and F) bilobate phytolith. All scale bars: 50 µm.
Fig. 3. Total biogenic Si pools in soils (means ± standard deviation, upper 5 cm) at Chicken Creek at the end of construction work ($t_0$) and after ten years of ecosystem development ($t_{10}$). Significant differences are indicated by different letters ($p < 0.05$, Kruskal-Wallis ANOVA with Dunn’s post hoc test).
Fig. 4. Boxplots (top, middle and bottom lines of the boxes show the 25th, 50th and 75th percentiles, respectively, and whiskers represent 1.5× the inter-quartile ranges) of biogenic Si pools in soils (upper 5 cm) at Chicken Creek at the end of construction work ($t_0$) and after ten years of ecosystem development (western, eastern and southern sections, $t_{10}$). A) Phytogenic Si pools (phytoliths), B) protophytic Si pools (diatom frustules), C) zoogenic Si pools (sponge spicules) and D) protozoic Si pools (testate amoeba shells). Significant differences are indicated by different letters ($p < 0.05$, Kruskal-Wallis ANOVA with Dunn’s post hoc test). Circles and asterisks indicate outliers and extreme values, respectively. Note different scales for diagrams A+B and C+D.
Fig. 5. Proportion of phytoliths (PHY), sponge spicules (SPO), diatom frustules (DIA) and testate amoeba shells (TA) to total BSi in soils (upper 5 cm) at Chicken Creek at \( t_0 \) and \( t_{10} \). Note that total BSi pools differ in size (see Fig. 3).
Fig. 6. Comparison of water soluble Si (Si$_{H2O}$) as well as amorphous Si (Si$_{Tiron}$) fractions and total BSi in soils (means ± standard deviation, upper 5 cm), where Calamagrostis epigejos (A) and Phragmites australis (B) became dominant. Data are given for t$_0$ (no vegetation) and t$_{10}$ (C. epigejos, P. australis). For t$_{10}$ total plant Si contents, extracted phytogenic Si (phytoliths) contents and Si pools for C. epigejos and P. australis (plants and litter) are stated in addition. Paintings from Cornelia Höhn, Müncheberg.

<table>
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<th>Soil</th>
<th>Litter</th>
<th>Difference</th>
</tr>
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<td>Si$_{H2O}$: 0.80 ± 0.37 g m$^{-2}$</td>
<td>Si$_{H2O}$: 1.59 ± 0.46 g m$^{-2}$</td>
<td>+ 99 %</td>
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<td>Si$_{Tiron}$: 497 ± 212 g m$^{-2}$</td>
<td>Si$_{Tiron}$: 290 ± 154 g m$^{-2}$</td>
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<td>BSi: 11.6 ± 6.5 mg m$^{-2}$</td>
<td>BSi: 44.2 ± 26.2 mg m$^{-2}$</td>
<td>+ 281 %</td>
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<td>Si$_{H2O}$: 2.16 ± 0.35 g m$^{-2}$</td>
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<td>Si$_{Tiron}$: 443 ± 175 g m$^{-2}$</td>
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<td>BSi: 88.2 ± 80.3 mg m$^{-2}$</td>
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Fig. 7. Micrographs of fragile phytogenic Si structures (arrows) of *C. epigejos* (A) and *P. australis* (B).
Table 1. Contents of skeleton (>2 mm), fine earth (<2 mm), sand, silt and clay fractions (upper 30 cm) at the sampling points in western, eastern and southern sections at Chicken Creek (t₀, calculations based on data of Gerwin et al. 2010). Minimal (Min) as well as maximal (Max) values, means (\(\bar{x}\)) and standard deviations (SD) are given.

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<td>%</td>
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**Table 2.** Measured soil parameters (upper 5 cm, means ($\bar{x}$) with standard deviation (SD)) at the different sections of Chicken Creek. Significant differences between $t_0$ and $t_{10}$ are each stated in bold (Mann-Whitney U-test, $p < 0.05$) or marked with asterisks ($p < 0.1$) for the western, eastern and southern section.

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<th>Section</th>
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<th>$S_{\text{Tiron}}$</th>
<th>$A_{\text{Tiron}}$</th>
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<th>$\text{CaCO}_3$</th>
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<td>312</td>
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<td>62</td>
<td>57</td>
<td>258</td>
<td>40</td>
<td>0.1</td>
</tr>
</tbody>
</table>

**Table 3.** Spearman’s rank correlations between measured soil parameters and total BSi (upper 5 cm, $n = 6$) at Chicken Creek. Significant correlation coefficients are given in bold ($p < 0.05$).

<table>
<thead>
<tr>
<th></th>
<th>$S_{\text{H}_2\text{O}}$</th>
<th>$S_{\text{Tiron}}$</th>
<th>$A_{\text{Tiron}}$</th>
<th>$F_{\text{Tiron}}$</th>
<th>$C_{\text{org}}$</th>
<th>$\text{CaCO}_3$</th>
<th>pH</th>
<th>BSi</th>
</tr>
</thead>
<tbody>
<tr>
<td>$S_{\text{H}_2\text{O}}$</td>
<td>1.000</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>$S_{\text{Tiron}}$</td>
<td>-0.257</td>
<td>1.000</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>$A_{\text{Tiron}}$</td>
<td>-0.600</td>
<td>0.829</td>
<td>1.000</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>$F_{\text{Tiron}}$</td>
<td>-0.486</td>
<td>0.771</td>
<td>0.943</td>
<td>1.000</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>$C_{\text{org}}$</td>
<td>0.714</td>
<td>0.086</td>
<td>-0.371</td>
<td>-0.486</td>
<td>1.000</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>$\text{CaCO}_3$</td>
<td>0.200</td>
<td>0.086</td>
<td>-0.086</td>
<td>-0.029</td>
<td>0.029</td>
<td>1.000</td>
<td></td>
<td></td>
</tr>
<tr>
<td>pH</td>
<td>-0.600</td>
<td>0.200</td>
<td>0.486</td>
<td>0.543</td>
<td>-0.771</td>
<td>0.543</td>
<td>1.000</td>
<td></td>
</tr>
<tr>
<td>BSi</td>
<td>0.941</td>
<td>-0.213</td>
<td>-0.577</td>
<td>-0.577</td>
<td>0.880</td>
<td>0.152</td>
<td>-0.698</td>
<td>1.000</td>
</tr>
</tbody>
</table>
Table 4. Surface-areas, volumes and surface-to-volume ratios (A/V) of different biogenic siliceous structures found at Chicken Creek.

<table>
<thead>
<tr>
<th></th>
<th>Surface-area (µm$^2$)</th>
<th>Volume (µm$^3$)</th>
<th>A/V ratio</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Min.</td>
<td>Max.</td>
<td>Min.</td>
</tr>
<tr>
<td>Bilobate phytoliths</td>
<td>216</td>
<td>3,730</td>
<td>36</td>
</tr>
<tr>
<td>Elongate phytoliths</td>
<td>2,302</td>
<td>22,203</td>
<td>390</td>
</tr>
<tr>
<td>Diatom frustules*</td>
<td>351</td>
<td>9,901</td>
<td>347</td>
</tr>
<tr>
<td>TA shells*</td>
<td>1,229</td>
<td>5,085</td>
<td>900</td>
</tr>
<tr>
<td>Sponge spicules*</td>
<td>305</td>
<td>16,963</td>
<td>291</td>
</tr>
<tr>
<td>Spicule fragments*</td>
<td>2,828</td>
<td>17,268</td>
<td>5,255</td>
</tr>
</tbody>
</table>

* Data taken from Puppe et al. (2016).