New highlights of phytolith structure and occluded carbon location: 3-D X-ray microscopy and NanoSIMS results

A. Alexandre¹, I. Basile-Doelsch¹, T. Delhaye², D. Borshneck¹, J. C. Mazur¹, P. Reyerson³, and G. M. Santos⁴

1Centre Européen de Recherche et d’Enseignement des Géosciences de l’Environnement (UMR 7330), CNRS, Aix-Marseille Université, Europôle méditerranéen de l’Arbois BP 80, 13545 Aix en Provence CEDEX 04, France
2Plateforme NanoSIMS, OSUR, Université de Rennes 1, Campus de Beaulieu, 35042 Rennes CEDEX, France
3Department of Geography, University of Wisconsin-Madison, 550 North Park Street, Madison, WI 53706, USA
4Department of Earth System Science, University of California, Irvine, B321 Croul Hall, Irvine, CA 92697-3100, USA

Correspondence to: A. Alexandre (alexandre@cerege.fr)

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Abstract. Phytoliths contain occluded organic compounds called phytC. Recently, phytC content, nature, origin, paleoenvironmental meaning and impact in the global C cycle have been the subject of increasing debate. Inconsistencies were fed by the scarcity of in situ characterizations of phytC in phytoliths. Here we reconstructed at high spatial resolution the 3-D structure of harvested grass short cell (GSC) phytoliths using 3-D X-ray microscopy. While this technique has been widely used for 3-D reconstruction of biological systems it has never been applied in high-resolution mode to silica particles. Simultaneously, we investigated the location of phytC using nanoscale secondary ion mass spectrometry (NanoSIMS). Our data evidenced that the silica structure contains micrometric internal cavities. These internal cavities were sometimes observed isolated from the outside. Their opening may be an original feature or may result from the beginning of dissolution of silica during the chemical extraction procedure, mimicking the progressive dissolution process that can happen in natural environments. The phytC that may originally occupy the cavities is thus susceptible to rapid oxidation. It was not detected by the NanoSIMS technique. However, another pool of phytC, continuously distributed in and protected by the silica structure, was observed. Its N/C ratio (0.27) is in agreement with the presence of amino acids. These findings constitute a basis to further characterize the origin, occlusion process, nature and accessibility of phytC, as a prerequisite for assessing its significance in the global C cycle.

1 Introduction
When absorbing nutrients in the soil, plants roots also uptake a significant amount of silicon (Si). The Si fluxes recycled by plants are substantial; for example, Si take up by tropical forests or grasslands can reach 2 to 10 times Si fluxes generated from the dissolution of soil silicates that are exported to stream water (e.g., Blecker et al., 2006; Struyf and Conley, 2009; Cornelis et al., 2011; Alexandre et al., 2011). Inside the plant, Si is transported in the sap and deposited in the cells, in the cell walls and in extracellular spaces of stems and leaves as micrometric hydrous amorphous silica particles called phytoliths. Upon plant decay, part of the phytolith production can be incorporated into soils or sediments and preserved for as long as millions of years (Alexandre et al., 2011; Miller et al., 2012; Strömberg et al., 2013). These fossil phytolith assemblages can be used for reconstructing past vegetation and climate conditions via their morphological and geochemical signatures (Piperno, 2006; Alexandre et al., 2012). Phytoliths occlude small amounts of organic compounds, first evidenced by the production of carbon (C) and nitrogen (N) during dry ashing (Jones and Beavers, 1963). Later on, scanning transmission electron microscopy (STEM) and energy dispersive X-ray (EDX) analyses of phytoliths in the plant tissues confirmed that the occluded organic compounds contained C, N and phosphorus (P) (Laue et al., 2007). By extension, these occluded compounds are here called phytC. PhytC, which is assumed to be...
protected from natural oxidation by the siliceous structure, has been the subject of increasing attention and debate.

Based on the assumption that phytC originated from the photosynthesis of atmospheric CO$_2$ in the host plant, several studies used phytC$^{14}$C and $\delta^{13}$C signatures respectively as a dating tool (Piperno and Becker, 1996; Piperno and Stothert, 2003; McMichael et al., 2012) and a paleoenvironmental proxy (Kelly et al., 1991; Smith and White, 2004; Carter, 2009; Webb and Longstaffe, 2010; McInerney et al., 2011). However, very recently, $^{14}$C-AMS measurements of phytC samples from modern grasses yielded ages of several thousand years, which suggested that phytoliths may incorporate a substantial amount of old carbon, potentially from the soil (Santos et al., 2010, 2012). Amino acids from soils have been shown to be taken up by plants, and transported in small proportions to roots, stems and shoots (Paungfoo-Lonhienne et al., 2008; Whiteside et al., 2009, 2012; Gao et al., 2010; Warren, 2012). Thus it is not inconsistent to assume that C and N derived from these soil amino acids have been trapped in phytoliths. Although the hypothesis still needs to be verified, it raises the question of the molecular nature of phytC. Several techniques such as high-performance liquid chromatography (HPLC), gas chromatography mass spectrometry (GC-MS), protein staining, micro-Raman analysis and X-ray photoelectron spectroscopy (XPS) have been used to characterize phytC and led to contradictory results, especially regarding the presence or not of amino acids (Harrison, 1996; Pironon et al., 2001; Smith and Anderson, 2001; Elbaum et al., 2009; Watling et al., 2011). The problem is that these methods were applied on phytolith concentrates that were not proven to be completely devoid of extraneous organic remains. Chemical extractions leading to high-purity phytolith concentrates are indeed difficult to implement. Although the absence of organic particles can be checked by scanning electron microscopy (SEM) coupled with EDX (Corbineau et al., 2013), the presence of extraneous organic remains on the phytolith surface cannot be accurately detected.

Differences in the efficiency of phytolith extraction protocols may also explain the inconsistencies in phytC quantification. Accurately quantifying the phytC is important for the assessment of its significance in the terrestrial C cycle. Multiple studies have recently claimed that phytC may play a role in atmospheric CO$_2$ sequestration and climate change mitigation (Parr and Sullivan, 2005; Parr et al., 2010; Song et al., 2014; Huang et al., 2014; Li et al., 2014; Zuo et al., 2014), although the fluxes of phytC from vegetation to soils and the residence time of phytC in soils are still largely unknown. PhytC content as high as 20 % dry weight was obtained when using a phytolith extraction method based on microwave digestion (Parr and Sullivan, 2014). This value was more than 20 to 200 times higher than the values obtained using a chemical method verified to be 100 % efficient for removing extraneous organic particles (from 0.1 to 1 % dry weight; Smith and White, 2004). The difference was somewhat justified by partial dissolution of phytC when using aggressive protocols. The assumption that phytC may be located at different sites in the silica structure, with different accessibility to oxidation, has been put forward (Parr and Sullivan, 2014). This assumption supplemented a previous one, widely found in the literature, that micrometric opaque areas observed by natural light (NL) microscopy on some phytoliths were holes containing the phytC (Prychid et al., 2003; Piperno, 2006; Carter et al., 2009; Song et al., 2012; Parr and Sullivan, 2014). No measurements were however performed to support any of these hypotheses.

Finally, the debates on content, location, nature, origin and paleoenvironmental meaning of phytC have been fed by the scarcity of in situ characterizations of phytC in phytoliths, despite few seminal works (Harrison, 1996; Laue et al., 2007). Here we reconstructed, at high spatial resolution, the 3-D structure of grass phytoliths using 3-D X-ray microscopy. Simultaneously, we characterized the location of phytC using nanoscale secondary ion mass spectrometry (NanoSIMS).

## 2 Material and methods

Grasses are among the main producers of phytoliths. The leaves of *Triticum durum* wheat (TD-F-L), were harvested in 2012 at the Genomics Research Centre in Fiorenzuola d’Arda (Italy). Hundreds of grams were made available to us for phytC investigation. Phytoliths were extracted from 50 g of dry leaves using a wet chemical protocol recently set up for geochemical analysis of phytC. The protocol was described in detail in Corbineau et al. (2013). The organic matter was oxidized with H$_2$SO$_4$, H$_2$O$_2$, HNO$_3$ and KClO$_3$, and potential remains on the phytolith surface were dissolved using KOH (pH of 11). The absence of residual extraneous organic particles was checked using SEM-EDX (Corbineau et al., 2013). Dominant phytolith types were recognized according to Madella et al. (2005) using NL microscopy at 600× and 1000× magnifications. As expected, the grass small cell group (GSC) and the bulliform cell group dominated the assemblage. These groups, which form in all grass epidermis, also dominate phytolith assemblages produced by grasslands and recovered from soils (e.g., Alexandre et al., 2011). Several NL microscopy and SEM pictures, illustrating the composition of the TD-F-L phytolith assemblage were taken. For the purpose of morphological comparison, pictures of fossil GSC and bulliform phytoliths from available soil assemblages described in previous papers, were additionally taken.

The 3-D structure of the GSC phytoliths was reconstructed by X-ray imaging at the micro-scale, using a 3-D X-ray microscope Zeiss Ultra XRM L 200. A few phytoliths, randomly selected from the bulk sample, were deposited on the inner surface of a bevel-cut Kapton tube of 50 µm internal diameter. Five individual GSC phytoliths were recognized by optical microscopy at 200× magnification and their position...
located for 3-D visualization. The principle of the 3-D X-ray microscopy technique is based on focusing the X-ray beam on a rotating sample using an optical lens; then transmitted X-rays are diffracted by a Fresnel zone plate on a scintillator in front of an optical device to produce a 200× magnified image of the phytolith captured by a charge-coupled device (CCD) image sensor. Using a 1K × 1K detector, it leads to a voxel size of 63 nm. The X-ray beam path is continuously flushed with helium to minimize the absorption of X-rays by air, the sample and the optics excepted. While this technique has been widely used for 3-D reconstruction of biological systems it has never been applied in high-resolution mode to silica particles. Analysis of the phytoliths proceeded at 150 nm resolution for a 65 µm field of view, in conventional absorption contrast imaging mode at 8 keV (copper rotating anode; power set at 40 kV and 30 mA). Using this mode, the contrast was generated both from the different X-ray attenuation coefficients of the chemical elements composing the sample and from the density. A total of 901 X-ray projections were recorded between −90 and +90° at an angle step of 0.2° and an exposure time of 80 s per view. After 20 h of analysis, reconstruction of the phytolith volume was performed using XMReconstructor (Zeiss Xradia software). The resulting stack of 2-D grayscale slices was then exported to Avizo Fire (FEI group) for further image processing.

NanoSIMS analyses were performed on cross sections of TD-F-L phytoliths embedded in epoxy resin. One milligram of phytoliths was deposited on polytetrafluoroethylene (PTFE) filters (9 mm i.d.) stuck onto double face tape. Polypropylene (PP) tubes (10 mm i.d. and 15 mm long) were placed on the tape, encircling the phytoliths. Epoxy resin (Araldite 100/Hardener 16) was slipped into the tubes up to 3 mm height and left to dry for 3 h at 40 °C. Resin of 7 mm height was added and left to dry for 48 h at 40 °C. Those two steps prevented the resin from leaking from the base of the tube. Embedded samples were taken off the tubes and polished with diamond paste up to 0.1 µm, until the PTFE filter was completely removed and cross sections of phytoliths were visible in NL microscopy. Samples were sawn into 4 mm thick blocks. Dozens of GSC phytoliths cross sections to be analyzed with the NanoSIMS were located by SEM. The NanoSIMS technique is based upon the sputtering of a few atomic layers from the surface of a sample induced by a primary ion bombardment. The primary ion impact triggers a cascade of atomic collision. Atoms and atomic clusters are spontaneously ionized. These secondary ions are characteristic of the composition of the analyzed area. They are separated according to their mass and an image of the intensity of the secondary ion beam is made for a selected mass (http://www.cameca.com/instruments-for-research/sims.aspx. Over the past few years, the NanoSIMS technique has increasingly been used in geosciences to investigate the elemental and isotopic composition of organic and inorganic materials (Herrmann et al., 2007; Hatton et al., 2012; Mueller et al., 2012, 2013). The NanoSIMS technique however has only scarcely been used for measuring secondary ion emission from amorphous silica. One study showed NanoSIMS images of a thin section of a giant siliceous sponge spicule (several millimeters in diameter). A micrometric proteinaceous scaffold, which averaged 2 % C dry weight, could be detected in the siliceous structure (Müller et al., 2010). The NanoSIMS technique was also used for identifying silification sites in rice roots (Moore et al., 2011). Here, we analyzed the intensities of [28Si]−, [16O]−, [24C2]− and [26CN]− ions produced by selected areas of the GSC phytoliths polished cross sections using a Cameca NanoSIMS 50. The section was coated with 25 nm gold and introduced in the NanoSIMS. A [Cs]+ primary ion probe with 16 kV primary ion impact energy and a 8 kV secondary ion extraction voltage was used. The best adjustment for obtaining secondary ion images of [28Si]−, [16O]−, [12C]− and [26CN]− was the following: the selected phytolith surface were first pre-sputtered with a de-focused primary beam on a 60 µm × 60 µm area for 3 min. Then 256 × 256 pixel images were made using a 2.2 pA primary ion current (primary diaphragm diameter of 300 µm) and a counting time of 10 ms per pixel for areas of 30 µm × 30 µm. Analyses with longer counting time or larger primary diaphragm/higher primary beam intensity were also tested. Secondary ion images of [28Si], [16O]−, [12C]− and [26CN]− were processed using the ImageJ software (http://imagej.nih.gov/ij). Colors were assigned to different signal intensities, increasing from black to red. Images of the [26CN]/[12C] ratio were also created. Line scans were drawn across the analyzed surfaces and ion intensity vs. distance along the line were plotted.

For comparison with the NanoSIMS results, the C and N contents of the bulk TD-F-L phytolith sample were measured by chemiluminescence after combustion at 1350 °C (for C) and 1000 °C (for N). The C and N contents of the epoxy resin were measured with an elemental analyzer (EA) after combustion at 1350 °C.

3 Results

Three morphological categories of phytoliths, commonly found in grasses, constituted the bulk sample. SEM pictures of phytoliths placed on the aluminum mount illustrate these categories in Fig. 1. SEM pictures of cross sections of the same categories are shown in Fig. 2. For each category, the mode of silica deposition is specified below when it has been previously evidenced in SEM, TEM, fluorescence microscopy or NanoSIMS images of plant cross sections (Sanger and Parry, 1969; Sowers and Thurston, 1979; Harrison, 1996; Currie and Perry, 2007; Law and Exley, 2011; Moore et al., 2011). The first phytolith category is constituted by thin fragments of multi-cellular silica sheets, several tenths of a micrometer long and wide but less than a few micrometers thick (Figs. 1a, b, 2a, b). These silica “skeletons” (Sang-
SEM images of the thin section of the TD-F-L wheat phytolith types including silica sheets (a, b), and GSC phytoliths of the rondel type (c, d, e). GSC types show micrometric internal cavities (IC).
Two examples of reconstructed 3-D X-ray microscopy volumes are presented in Figs. 4 and 5. The observed patterns were common to the five analyzed GSC particles. The siliceous structure appeared porous at the sub-micrometer scale (Figs. 4a and 5a). Inside the structure, areas of a few micrometers in diameter, with significantly lower X-ray absorption than the surrounding, were observed (Fig. 4a). 2-D planes of the reconstructed volumes evidenced that these heterogeneities were the cavities several micrometers wide previously identified on the cross sections by SEM. The cavities were interconnected (Figs. 4b, 5b). Some particles showed cavities isolated from the phytolith surface by a few micrometer thick silica wall (Fig. 4b). Other particles showed cavities connected to the phytolith surface by small holes of 0.1 µm diameter only (Fig. 5b). These cavities appeared to be filled with air (no X-ray absorption), although the high contrast in X-ray absorbance between silica and air may have masked the presence of organic compounds.

The NanoSIMS results, common to the dozens of analyzed phytolith thin sections, are illustrated in Figs. 6–8. Adjustments were done to find the pre-sputtering duration (3 min), the primary ion beam intensity (L1 = 2 kV), the primary diaphragm diameter (750 µm) and the duration of analyses (11 min) appropriate for obtaining sufficient total ion current (TIC) and avoid charging effects (Figs. 6a, 7a). When the primary ion beam intensity was increased to L1 = 4 kV (Fig. 8a), when the primary diaphragm diameter was decreased to 300 µm (Fig. 8b), or when a succession of analyses resulted in increasing the duration of sputtering (Fig. 8c), a zone devoid of secondary ion signal appeared at the center of the silica surface. This was probably due to charging (Mueller et al., 2012) and/or to topographic heterogeneity (Winterholler et al., 2008). As silica was more resistant to polishing than the epoxy, silica surfaces were often convex (Fig. 8). The tests conducted here emphasized the importance of looking for the most efficient adjustment (i.e., avoiding charging and topographic effects) before performing NanoSIMS analyses on silica surfaces.

[28Si]−, [16O]−, [12C]− and [26CN]− images clearly individualized phytoliths from the epoxy resin. The [28Si]− and [16O]− images and scan lines showed that phytoliths were made of a continuous silica structure (Figs. 6 and 7) sometimes interrupted by central micrometric areas devoid of silica (Fig. 7). This is again in concordance with the central cavities identified in SEM and 3-D X-ray imaging. Carbon was present in the cavities and in the silica structure itself. However when values of [12C]− intensity were similar in the cavities and in the epoxy resin, they were 10 to 20 times lower in the silica structure than in the epoxy resin (Figs. 6 and 7). N was also present in the silica structure and [26CN]− intensity was 3 to 4 times lower in the silica structure than in the cavities or the epoxy (Figs. 6 and 7). Interestingly, the [26CN]−/[12C]− ratio ranged between 20 and 30 in the silica structure and between 5 and 10 in the cavities and the epoxy. The silica structure was thus enriched in N by a factor of 4 to 8 relative to the surrounding epoxy. These features were reproducible from a particle to another. Bulk C and N contents in phytoliths, measured by chemiluminescence and EA (cf. Sect. 2), were 0.4 and 0.1 % dry weight for phytoliths, and 68.8 and 2.8 % dry weight for the epoxy resin, respectively. The N / C ratio was 0.27 for the phytoliths and 0.04 for the epoxy resin. The bulk phytolith sample was thus enriched in N relative to the epoxy resin by a factor of 6.8, in agreement with N enrichments calculated from the NanoSIMS data. This consistency strengthened the accuracy of the [12C]− and [26CN]− relative intensities measured with the NanoSIMS. Finally, [26CN]−/[12C]− NanoSIMS images clearly showed that organic compounds, with N content significantly higher than in the resin, were continuously distributed (at the sub-micrometer scale) in the silica structure. On the contrary, cavities appeared to be filled with the epoxy resin.

4 Discussion

4.1 PhytC locations in the silica structure of GSC phytoliths

SEM, 3-D X-ray microscopy and NanoSIMS images showed that the silica structure of GSC phytoliths was homogeneous at the micrometric scale and systematically contained central micrometric interconnected cavities. The fact that some particles contained cavities isolated from the outside suggests that the opening to the outside can be either original or result from dissolution after the phytolith formation. Phytoliths often contain up to a few percent by dry weight of aluminium (Al) by dry weight (Bartoli and Wilding, 1980; Carnelli et al., 2004) co-precipitating with silica (Hodson and Sangster, 1993). As Al dissolves in strong acids and in strong bases, the phytolith chemical extraction procedure that included HNO3 and H2SO4 steps may have initiated phytolith surficial dissolution and opened the few micrometer thick sil-
Figure 4. 3-D X-ray microscopy of a GSC phytolith from wheat (TD-F-L): (a) four views of the reconstructed volumes; internal cavities (IC) are distinguishable. (b) 2-D X-ray slices superimposed on the phytolith volume rendering, showing from front to back the internal cavity (IC). No connection to the surface was evidenced. The blue area corresponds to the thresholding of the phytolith grayscale values.

Figure 5. 3-D X-ray microscopy of a GSC phytolith from wheat (TD-F-L): (a) reconstructed volume. (b) 2-D X-ray images from back to front of the phytolith showing the internal cavity (IC) and its connection to the surface, forming holes (H). The blue area corresponds to the thresholding of the phytolith grayscale values.

ica wall between the cavities and the phytolith surface. The procedure also included a final alkaline step (KOH solution at pH 11) that may also have increased the dissolution features on the silica surfaces. As phytoliths were directly extracted from the plant, the surficial dissolution was revealed here at its beginning. It is expected to reach higher degrees over time in natural environments where multiple dissolution factors come into play (Iler, 1979). Large dissolution features were indeed often observed on fossil phytoliths and were quantified to assess the degree of weathering of soil phytolith assemblages (Alexandre et al., 1999; Oleschko et al., 2004). To illustrate this point, SEM and NL microscopy images of the entirety and cross sections of fossil monocellular phytoliths collected from soils are shown in Fig. 9. The phytolith types are characteristic of grass epidermis (GSC types and Cuneiform bulliform types; Madella et al., 2005) (Fig. 9a, b) and wood parenchyma (globular granulate type; Madella et al., 2005) (Fig. 9c). The dissolution of silica has made central depressions of several micrometers wide. The particles appear empty inside, which is consistent with dissolution starting from the silica walls located between the cavities and the phytolith surface, then slightly opening, or increasing the opening of the cavities to the outside, and then enlarging the cavities into dissolution depressions. Such dissolution depressions are not limited to GSC phytoliths. They were observed on many types of monocellular phytoliths from grasses and non-grasses extracted from soils and sediments as illustrated in Fig. 9a5 (Acicular type), 9b2 and 9b3 (globular granulate). This implies that the inner part of all these phytolith types was constituted by silica less dense than the outer part, either due to phytC occlusion or to a lack of...
dissolved Si available for precipitation during the phytolith formation.

Inside the internal cavities, no original organic compounds could be detected by NanoSIMS. If initially present, they may have been squeezed out and replaced by the epoxy resin during the polishing step. On the contrary, the $[26\text{CN}]^-$/$[12\text{C}]^-$ images clearly evidenced the presence of organic compounds rich in N continuously distributed in the silica structure and clearly differentiated from the epoxy resin. The absolute composition of $[26\text{CN}]^-$ and $[12\text{C}]^-$ was not calculated. This would have required including standard materials with known composition in the analyzed section. However, the consistency of N enrichment of the organic compound in the silica structure (measured by NanoSIMS) with N enrichment of the bulk phytC (measured by chemiluminescence/EA) supports the claim that the organic com-
Evidence of the continuous distribution of phytC in the silica structure, at the sub-micrometric scale, suggests that it had been occluded since the early stage of silicification. SEM, environmental scanning electron microscope (ESEM) and TEM-EDX analyses showed that silica first precipitates in the inner cell wall, probably triggered by the presence of callose or lignin (Laue et al., 2007; Law and Exeley, 2011; Zhang et al., 2013). Silica nanospheres are then organized in a variety of structural motifs such as sheet-like, globular and fibrillar bundles that, from the cell wall, infill the cell lumen in a centripetal way (e.g., Kaufman et al., 1981; Sangster and Parry, 1981; Perry et al., 1987; Laue et al., 2007; Zhang et al., 2013), until most of the cell becomes silicified (Motomura, 1996; Laue et al., 2007). As previously noted, an organic template may participate to the silica formation Harrison, 1996; Laue et al., 2007). This organic template, progressively trapped in the silica structure, may constitute the phytC evidenced by NanoSIMS in the phytoliths. Its N / C value (0.27) is in the range of N / C values characteristic of amino acids. Amino acids may originate either from the cell itself or from the extra-cellular space. Different families of transporters have been identified for their import into plant cells (Tegeder, 2012). At the same time, amino acids entering the cell simultaneously to silica thanks to an invagination/vesicle formation mechanism previously evidenced (Neumann and De Figueiredo, 2002) may occur.

At the end of the cell silicification, residual cell organic compounds that were not already occluded may gather in any remaining space and delimitate the micrometric central cavities. This second pool of phytC should be rapidly oxidized when phytoliths start to dissolve after their deposition in litter, soil or sediment (Fig. 9). This suggests that this phytC pool participates to a limited extent in long-term atmospheric CO₂ sequestration. These considerations rise the need to further estimate the respective contributions to C contents measured from bulk phytolith concentrates of (i) phytC in the silica structure, (ii) phytC in the central cavities and (iii) extraneous C that may remain on porous phytolith surfaces. This is a prerequisite for reliable assessments of the significance of phytC in atmospheric CO₂ sequestration. For that purpose, phytC contents measured from phytolith assem-

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Figure 8. SEM images of the polished section showing convex silica surfaces (Si) in the epoxy resin (r). Associated NanoSIMS $^{28}\text{Si}^-$ images showing central areas devoid of secondary ion signal. (a) $[\text{Cs}]^+$ primary beam with $L_1 = 4 \text{kV}$, D 1-1 primary diaphragm (750 mm), 11 min; (b) $[\text{Cs}]^+$ primary beam with $L_1 = 2 \text{kV}$, D 1-2 primary diaphragm (300 mm), 11 min; (c) $[\text{Cs}]^+$ primary beam with $L_1 = 2 \text{kV}$, D1-1 primary diaphragm (750 mm), 3 min analyses for 1, 2 and 3 successively.

Figure 9. NL microscopy and SEM images of dissolution depressions (DD) affecting fossil phytoliths from soils. (a) Grass epidermis monocellular phytoliths (cuneiform bulliform types and acicular type) from Mascareignite (MSG 70, La Réunion, France) (Crespin et al., 2008); NL microscopy phytolith surface (1, 2), SEM phytolith volume (3) and polished section (4, 5). (b) Grass epidermis monocellular phytoliths from a ferrugineous soil (Salitre, Brazil) (Alexandre et al., 1999); NL microscopy phytolith. (c) Phytoliths from palms and trees from a ferrallitic soil (Dimonika, RDA) (Alexandre et al., 1997); SEM globular granulate type volumes (1, 2) and polished section (3). (d) Opaque areas observed with NL microscopy on bulliform cell phytoliths from MSG 70 (1, 2) and Salitre (3, 4). Scale bars: 10 µm.
blages characterized by 3-D X-ray microscopy as dominated by phytoliths with closed internal cavities or by phytoliths with open cavities should be compared.

4.3 Reassessment of NL microscopy observations

Several studies have speculated that opaque areas observed by NL microscopy in fossil phytoliths from soils and sediments were burnt organic remains indicative of past fire occurrence (Kealhofer and Penny, 1998; Elbaum et al., 2003; Parr, 2006; Piperno, 2006). However, when observed by NL microscopy, the empty dissolution depressions evidenced by SEM on monocellular phytoliths from soils (Fig. 9a) also appeared as opaque areas, especially when they were oriented downwards (Fig. 9c). This is probably due to trapped air in the dissolution depressions that caused an optical artifact at the place where the air met the mounting medium. This feature implies that opaque areas in fossil phytoliths should not be considered as unequivocal evidence of burnt organic compounds. Similarly, internal cavities may also appear as opaque spots due to the occurrence of trapped air, independent of the presence of organic compounds.

5 Conclusions

3-D X-ray microscopy reconstructions of GSC phytoliths from harvested grasses, and SEM observations of their cross sections, showed that the silica structure contains micrometric internal cavities. These cavities were sometimes observed isolated from the outside. Their opening may be an original feature or may result from the silica dissolution during the chemical extraction procedure, mimicking the beginning of dissolution process that may happen in natural environments. The phytC that may originally occupy those cavities is thus susceptible to rapid oxidation. It was not detected by the NanoSIMS technique. On the contrary, another pool of phytC, continuously distributed in and protected by the silica structure was evidenced by NanoSIMS. Its N/C ratio (0.27) is in agreement with the presence of amino acids. These findings constitute a basis to further characterize the origin, occlusion process, nature and accessibility of phytC, necessary for assessing its significance in the global C cycle.

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