- Direct uptake of organic-derived carbon by grass roots and allocation in leaves and phytoliths:
- 2 ¹³C labeling evidence
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Abstract

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In the rhizosphere, the uptake of low molecular weight carbon (C) and nitrogen (N) by plant roots has been well documented. While organic N uptake relatively to total uptake is important, organic C uptake is supposed to be low relatively to the plant's C budget. Recently, radiocarbon analyses demonstrated that a fraction of C from the soil was occluded in amorphous silica micrometric particles that precipitate in plant cells (phytoliths). Here, we investigated whether and in which extent organic-derived C absorbed by grass roots can feed the C occluded in phytoliths. For this purpose we added ¹³C- and ¹⁵N-labeled amino acids (AAs) to the silicon-rich hydroponic solution of the grass Festuca arundinacea. The experiment was designed to prevent C leakage from the labeled nutritive solution to the chamber atmosphere. After 14 days of growth, the ¹³C and ¹⁵N enrichments (¹³C-excess and ¹⁵N-excess) in the roots, stems and leaves, and phytoliths were measured relatively to a control experiment in which no labelled AAs were added. Additionally, the ¹³C-excess was measured at the molecular level, in AAs extracted from roots and stems and leaves. The net uptake of labeled AA derived-¹³C by Festuca arundinacea reached 4.5% of the total AA-¹³C supply. The amount of AA derived-¹³C fixed in the plant was minor but not nil (0.28% and 0.10% of total C in roots and stems/leaves, respectively). Phenylalanine and methionine that were supplied in high amount to the nutritive solution, were more ¹³C-enriched than other AAs in the plant. This strongly suggested that part of AA derived-¹³C was absorbed and translocated into the plant in its original AA form. In phytoliths, AA derived-¹³C was detected. Its concentration was of the same order of magnitude than in bulk stems and leaves (0.15% of the phytolith C). This finding strengthens the body of evidences showing that part of organic compounds occluded in phytoliths can be fed by C entering the plant through the roots. Although this experiment was done in nutrient solution and its relevance for soil C uptake assessment is therefore limited, we discuss plausible forms of AA derived-13C absorbed and translocated in the plant and eventually fixed in phytoliths, and implication of our results for our understanding of the C cycle at the soil-plantatmosphere interface

1. Introduction

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In the rhizosphere, there are numerous known interactions between carbon (C) and nitrogen (N) 38 processes that have yet to be accurately assessed in qualitative and quantitative terms for their 39 consideration in carbon cycle models (Heimann and Reichstein. 2008). Among those interactions the 40 uptake of low molecular weight C and N (e.g. organic acids, sugars and amino acids (AAs)) by plant 41 roots (both mycorrhizal and non-mycorrhizal plants) has been well documented through labeling 42 experiments using hydroponic solutions, artificial substrats or soils (e.g. Bardgett et al., 2003; Kuzyakov 43 and Jones, 2006; Biernath et al., 2008; Jones et al., 2009a; Näsholm et al., 2009; Sauheitl et al., 2009; 44 Rasmussen et al., 2010; Gioseffi et al.. 2012; Moran-Zuloaga et al.. 2015). The aim of most of these 45 studies was to investigate in which extent and under which conditions organic N could be utilized by 46 plants as a direct source of N (i.e. without going through a mineralization step). The answers are still 47 debated (Jones and Darrah, 1992; Jones et al., 2009a; Rasmussen et al., 2010; Moran-Zuloaga et al., 48 2015). Especially, the evidence that plant roots can uptake labeled inorganic C that may bias the results 49 from organic N uptake studies when using bulk measurement of dual-labeling (¹³C and ¹⁵N) has been 50 51 put forward (Biernath et al., 2008; Rasmussen et al., 2010). However, the use of molecular and position-52 specific labeling technics can evidence the uptake and fixation of intact AAs (Sauheitl et al., 2009, Moran-Zuloaga et al., 2015). Organic C uptake was also investigated through the estimation of the net 53 uptake of glucose-C. This uptake has been shown to be low relatively to the plant's C budget, and was 54 often interpreted as the recapture of roots exudates (Jones and Darrah. 1992. 1993. 1996; Kuzyakov and 55 Jones. 2006; Jones et al., 2009a). However, very recently, in the frame of a non-labeling experiment, 56 radiocarbon analyses demonstrated that a fraction of C occluded in amorphous silica micrometric 57 particles that precipitate in plant cells (phytoliths) came from old soil C (Santos et al., 2012; Reverson 58 et al., 2015). Silicon (Si) is the second most abundant element of the earth surface after oxygen. Its 59 uptake by plants is widespread and generates, at the ecosystem scale, important fluxes from the soil to 60 plants (Conley, 2002). For instance, Si absorption represents 2 to 10 times the amount of dissolved Si 61 exported to stream water in tropical ecosystems (Alexandre et al., 2011). If part of the soil C uptake is 62 linked to Si uptake in the rhizosphere, the involved flux may thus also be significant. 63 Here, we aim to investigate whether and in which extent C derived from organic forms such as AAs can 64 be absorbed by grass roots, fixed in the plant and ultimately feed the organic C occluded in phytoliths 65 (phytC). We choose to focus on AAs as they are ubiquitous in soil organic matters of various residence 66 times (Bol et al., 2009). For this purpose we added ¹³C- and ¹⁵N-labeled AAs to the Si-rich nutrient 67 solution of the grass Festuca arundinacea. After two weeks of growth, the ¹³C and ¹⁵N enrichments in 68 the roots, stems/leaves, and phytoliths of the grass (two replicates) were quantified. Enrichments were 69

also measured at the molecular level (four replicates), in AAs of roots and stems/leaves. PhytC could

not be analyzed at the molecular level, due to its very small concentration. The experiment was designed to prevent C leakage from the labeled nutritive solution to the chamber atmosphere.

2. Material and methods

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2.1 Hydroponic culture

Festuca arundinacea, commonly referred to as tall fescue, is widely distributed globally as a forage and an invasive grass species (Gibson and Newman, 2001) and can adapt to a wide range of conditions. Festuca arundinacea was grown in hydroponic conditions for 24 days using an experimental procedure adapted from RHIZOtest (Bravin et al., 2010), a plant-based test recently standardized (ISO 16198:2015). Seventy-two plant-receiving pots (i.e. a cylinder closed at the bottom with a polyamide mesh of 30 µm pore size, using an adjustable clamp) were inserted in three perforated platforms covering three 12L tanks containing the nutrient solutions (24 plant pots per tank) (fig. 1). This assembly enabled close contact between seeds or seedling roots and the nutrient solutions. In order to prevent escape of the C- and N-bearing gas from the nutrient solution to the chamber atmosphere, O-rings sealed the plant pots to the perforated platform and the perforated platform to the tank. Additionally, the seeds were covered with agar-agar (polysaccharide agarose). Each tank was hermetically connected to two 20L containers (an input container filled with the nutrient solutions and a waste container). Seeds were first germinated for 10 days in a germination solution and seedlings were then grown for 14 days in a growth solution. The growth solution was entirely renewed once, after 8 days of growth. Otherwise the germination and nutritive solutions were renewed at a rate of 2L/24H using a peristaltic pump (fig. 1). Overall, 10L of germination solution and 64L of growth solution were used per tank. Germination and the growth nutrient solution composition were described in detail in Guigues et al. (2014). The nutrient solutions included 42mg/L of inorganic N (KNO₃) and 18mg/L of inorganic C in Ethylene diamine tetraacetic acid (EDTA), added to chelate metals the plant uses for growth. The solutions were also supplemented with 105 mg/L of SiO₂ (under the form of SiO₂K₂O). The growth chamber parameters were set at (day/night): 25/20°C, 75/70 % relative humidity and 16/8 h with a photon flux density of 450 µmol photons m⁻² s⁻¹ during the day.

At the end of the experiment, all samples were cleaned (to remove agar-agar), rinsed in deionized water and oven-dried at 50°C (to constant mass). When the tanks were filled (1st and 8th days of growth) and emptied (8th and 14th days of growth), the growth solution was sampled and kept frozen for future analyses.

2.2 Isotope labeling

In the two first tanks (two replicates), a mixture containing four ¹³C- and ¹⁵N-labeled AAs was added to the growth solution containing only inorganic C and N (as described above). Those AAs were selected for their following characteristics: L-Alanine (l-ALA) (C/N=2.6) is ubiquist and occurs in high

proportions in soils and plants proteins. The D-enantiomer of Alanine (D-ALA) which is present in natural soils (Hill et al., 2011), was expected to be more resistant to degradation and, if absorbed, less subject to metabolization. Consequently, we speculated that D-ALA may accumulate as a waste product first in cell vacuoles and later in phytoliths. L-Phenylalanine (L-PHE) (C/N=7.7) comprises a phenolic ring resistant to decomposition by microorganisms in soils, solutions or plants. L-Methionine (L-METH) (C/N=4.3) is a sulfur amino acid expected to be recovered at low abundance in plants, but is easily identifiable in Gas Chromatography-Isotope Ratio Mass Spectrometry (GC-IRMS). Commercial 97-99% ¹³C and ¹⁵N molecules (Eurisotop) were diluted with non-labeled amino-acids to reach the following atom abundances: L-PHE (19.51% ¹³C; 19.13% ¹⁵N); L-METH (19.87% ¹³C; 19.49% ¹⁵N); L-ALA (22.05% ¹³C; 16.26% ¹⁵N); D-ALA (7.43% ¹³C; 0.37% ¹⁵N). All AAs were uniformly labeled except i) D-ALA, which was not ¹⁵N-labeled but was ¹³C-labeled on one atom (C-2), and ii) an equivalent fraction of L-ALA because the labeled D-ALA was provided as a racemic mixture (DL-ALA). In each tank, the following amounts of AAs were added to the growth solution: PHE 249.9 mg; MET 125.1 mg; L-ALA 150.3 mg; D-ALA 125.2 mg. The total mixture (two tanks) represented 322 mg of C (26.8 mmol) and 75.7 mg of N (5.4 mmol) with average atom abundances of 18.15% ¹³C and 13.43% ¹⁵N. The maximum AAs concentration in the growth solution on the first or 8th days of the growth period was 0.225mmol/L, equivalent to 6.7mg/L of ¹³C and 1.6mg/L of ¹⁵N.

The third tank was only filled with the growth solution, without labeled AAs. It served as a control experiment to calculate the ¹³C and ¹⁵N enrichments of the plants from the labeled tanks, and verify that ¹³C and ¹⁵N derived from AAs (AA-¹³C and AA-¹⁵N), that may have contaminated the chamber CO₂. were not photosynthesized by the plants.

2.3. Analyses

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- For each tank (one control tank and two labeled tanks) stems/leaves were separated from the roots into two samples. The six resulting samples were ground finer than 200 µm. After alkaline fusion, they were analyzed in SiO₂ using Inductively Coupled Plasma Optical Emission Spectrometry (ICP-OES).
- For each sample, total C, ¹³C/¹²C, total N and ¹⁵N/¹⁴N were determined after dry combustion by IRMS using a Carlo Erba NA 1500 elemental analyzer (EA) coupled to a Thermo-Finnigan Delta-plus mass-spectrometer. Solutions were also analyzed for total C and ¹³C/¹²C by IRMS, after evaporation by dry combustion in tin capsules.
- Proteic carbon was analyzed as the sum of 19 individual AAs representing ca. 95% of all AAs. Quantification and ¹³C/¹²C determination of individual AAs were performed using a GC-IRMS (Thermo Fisher Scientific). The extraction and purification procedure was a slightly modified version (Rubino et al., 2014) of the protocol developed by Amelung et al. (2006). Briefly, dry plant samples were hydrolyzed in 6 M HCl (20 h. 100 °C). AAs were purified on Dowex 50 W X8 cation exchange resin

(100-200 mesh size, Arcos Organics, Thermo Fisher Scientific), dried by rotary evaporation, redissolved in 0.1 M HCl and dried again by speed-vacuum evaporation. AAs were separated and quantified as tert-butyl dimethyl silyl derivatives (TBDMS-aa): AAs were dissolved in N-Methyl-N-(tbutyldimethylsilyl) trifluoroacetamide (MTBSTFA) mixed with 1 % trimethylchlorosilasane (TMCS) (Sigma-Aldrich Co. Ltd.) and acetonitrile and heated at 120 °C for 1 hour. One µl of TBDMS-AA solution was injected into the GC through a GCCIII combustion interface (Thermo Fisher Scientific). TBDMS-AA were separated on a DB5 column (30 m. 0.25 mm i.d., 0.25 µm film thickness) with helium as a carrier gas. AA identification and quantification were performed using commercial mixtures of 20 proteinogenic AAs (Sigma Aldrich). Norvaline (Sigma-Aldrich Co. Ltd.) was added to plant samples before hydrolysis as an internal standard for quantification. Due to the addition of non-labeled carbon by TBDMS, AA ¹³C enrichment was subsequently calibrated for each AA (Shinebarger et al., 2002). Briefly, this calibration was based on the independent measurement of ¹³C of the TBDMS-derivatives of the commercial AA and an additional set of four ¹³C-labelled AA (Rubino et al., 2014). The calibration equation involves the number of carbon atoms added as TBDM and the isotopic composition of the latter. This isotopic composition term disappears in the calculation of the isotope excess (cf Eq. (1) below). When multiple peaks were encountered for a single AA, the main or the best individualized peak was chosen for both quantification and isotope ratio determination. The isotope ratios were calculated using ammonium sulfate IAEA-N1 (δ^{15} N = 0.43 ± 0.07 ‰), IAEA- N2 (δ^{15} N = 20.41 ± 0.07 %) and polyethylene IAEA CH7 (δ^{13} C = -32.15 ± 0.05%) as secondary standards. The sucrose standard IAEA CH6 used as a control yielded a mean value of 10.43 ‰.

Phytoliths were extracted from plants using a high purity protocol based on acid digestion and alkali immersion steps previously described in detail by Corbineau et al. (2013). Phytolith samples were observed in natural light microscopy to determine their morphological assemblage and check for the absence of residual organic matter particles. An additional purity check was done via Scanning Electron Microscopy (SEM) (Corbineau et al., 2013; Reyerson et al., 2015). Then, phytolith samples were analyzed for their C and N contents, as well as their ¹³C and ¹⁵N abundances by EA (Fisons NA 1500NC) coupled to a continuous flow IRMS (Finnigan Delta-Plus). About 6-10 mg of phytolith concentrates were weighed using a pre-calibrated microbalance (Sartorius AG, Göttingen, Germany) into tin capsules (5x9 mm capsules, Costech Analytical Technologies Inc., Valencia, CA, USA) and pre-baked at 100°C for 2 hours to remove extraneous contaminants. To assure accurate integration and linearization of the raw analytical data obtained from the lower C and N peaks, we decreased the helium carrier flow rate and measured several aliquots of in-house collagen L-Cystine 99% (δ^{13} C = -28.74 ± 0.13 % and δ^{15} N = -6.14 ± 0.07 %; from Sigma Aldrich Co. Ltd) and Atropina (δ^{13} C = -21.30 ± 0.06 % and δ^{15} N = -2.90 ± 0.10 %; from Costech 031042) as well as the internationally certified reference materials (e.g. Graphite USGS24 δ^{13} C = -16.05 ± 0.07 % and Amonium sulfate - IAEA-N1 δ^{15} N = +0.43 ± 0.07‰). Aliquots

of baked-clean silicon dioxide (SiO₂; mesh# -325, Sigma Aldrich, St. Louis, MO, USA) and fossil

phytoliths (MSG70; Crespin et al., 2008) were also analyzed to provide independent blank data (Santos

et al., 2010) and to check accuracy. To serve as quality assurance, note that the reproducibility obtained

on the phytolith laboratory standard MSG70 was s.d. $\pm 0.2\%$ for $\delta^{13}C$ and $\delta^{15}N$, and s.d. $\pm 0.01\%$ for C

178 and N.

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2.4. Calculations

- For the control tank (no labeling), results are reported as δ values in % relative to the Vienna Pee Dee
- Belemnite (V-PDB) for δ^{13} C, and in % relative of the atmospheric N₂ for δ^{15} N. For the labeled tanks.
- AA derived-¹³C and AA derived-¹⁵N plant concentration, recovery and net uptake were calculated on
- the basis of ¹³C-excess and ¹⁵N-excess of a sample relatively to the control tank samples according to
- 184 Eq. (1):

$$^{13}\text{C-excess}_{\text{sample}}(\%) = ^{13}\text{C atom}_{\text{sample}}(\%) - ^{13}\text{C atom}_{\text{control}}(\%)$$
 (1)

- where ¹³C atom_{sample} is the ¹³C atom abundance of a sample (stems/leaves, roots or phytoliths) from a
- labeled tank and ¹³C atom_{control} is the ¹³C atom abundance of the same sample from the control tank.
- The concentration of AA derived-¹³C in a sample, expressed in % of total C in the sample, was calculated
- 189 using Eq. (2):

190 [AA derived-
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C_{sample}] (%) = 13 C-excess _{sample} (%) / (13 C atom_{solution}(%) - 13 C atom_{control}(%)) x 100 (2)

- 191 Recovery of AA derived-¹³C in a sample, expressed in μg/g of the dry matter weight (d.wt), was
- calculated using Eq. (3):

193 Recovery AA derived-
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C (μ g/g) = [AA derived- 13 C _{sample}] x [C]_{sample}(μ g/g) (3)

- where [C]_{sample} is the concentration of total C in the sample.
- Net uptake of AA derived-¹³C, expressed in % of AA¹³C initially supplied to the solution, was calculated
- 196 using Eq. (4):

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- 197 AA derived- 13 C uptake (%) = recovery AA derived- 13 C (mg/g) x d.wt_{sample} (g)/ AA 13 C_{max supplied}(mg) (4)
- where d.wt_{sample} is the dry weight of sample and AA¹³C_{max supplied} is the maximum AA¹³C supplied to the
- 199 solution (322mg of AA-¹³C).
- 200 The same calculations were applied to ¹⁵N.

3. Results

3.1. Total C, N and AAs concentrations in the plants

203 After 14 days in the growth solution, the above-ground part of the plants of Festuca arundinacea were

30cm high but had not reached maturity or flower development (fig. 1). The C/N ratios of the stems and

leaves were similar in the labeled and control plants (8.4 and 9.0), whereas the C/N ratios of roots were higher in labeled (11.0) than in control (4.2) plants (Table 1; fig. 2). Recovered AAs accounted for 52 mgC/g of the dry matter in both roots and leaves. This is a high AAs content, also attested by the low C/N ratio, in agreement with the high N level requirement of young plants sufficiently fertilized to support rapid protein synthesis (Mattson, 1980). However AA derived-¹³C and AA derived-¹⁵N only accounted for ca. 13-14% of total C, and 40% of total N. This N contribution was lower than what might be expected. Indeed, at any stage of growth, AAs (in the form of protein or free molecules) should account for more than 50% of grass N. This discrepancy can be attributed to an underestimation of AAs by the extraction-purification process (i.e. incomplete hydrolysis recovery due to recombination into strong acids, and incomplete silylation). An underestimation of the AAs is consistent with the fact that amino-acid TBDMS-derivatives, such as tryptophane or cystine (the cystein-dimer), could not be recovered. However such an underestimation should not bias the measured relative proportion of methionine, phenylalanine and alanine.

3.2. Excess, uptake and recovery of AA derived-¹³C and AA derived-¹⁵N in the plants

- The δ values of the roots and aerial parts of the control plants were respectively -31.0 and -31.7% for
- δ^{13} C and 13.8 and 14.5% for δ^{15} N (Table 1). These values were in the range of the ones measured for
- 221 C₃ grasses in natural conditions (δ¹³C from -22 to -34 ‰; e.g. O'Leary, 1988), ensuring that CO₂
- 222 potentially produced by decomposition of the ¹³C-labeled molecules inside the tanks did not contaminate
- 223 the growing chamber atmosphere. The amount of labeled C recovered in plants and phytoliths was thus
- 224 considered as exclusively resulting from root uptake.

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- 225 The low number of replicates (2 labeled tanks and 2 molecular extracts per sample) was a compromise
- basically constrained by the large amount of matter required to isolate phytolith-occluded C and the
- experimental/analysis/cost effort. Table 1 shows that standard deviations calculated on the two replicates
- 228 were always one order of magnitude lower than the mean values.
- Relative to plants from the control tank, plants from the labeled tanks were enriched by 0.05% in ¹³C
- and 0.5% in 15 N in the roots and by 0.02% in 13 C and 0.3% in 15 N in the stems and leaves (Table 1).
- Overall, the net uptake of AA derived-¹³C and AA derived-¹⁵N by the plant represented respectively
- 4.5% and 46.9% of the AA¹³C and AA¹⁵N added to the solution (Table 1). Whereas 35% of absorbed
- 233 AA derived-¹³C and 19.2% of absorbed AA derived-¹⁵N were stored in the roots, 64.4% and 81.1% of
- 234 the absorbed AA derived-¹³C and AA derived-¹⁵N, respectively, were allocated to the stems and leaves
- 235 (after Table 1). The associated AA derived-¹³C/AA derived-¹⁵N ratios were 0.8 in the roots and 0.3 in
- 236 the stems and leaves (Table 1).
- 237 Concentrations of AA derived-¹³C and AA derived-¹⁵N represented only 0.13% of total C and 2.8% of
- 238 total N of the plant, respectively (Table 1; fig. 2). This contribution was higher in roots (0.28% of C and

- 4.1% of N) than in stems and leaves (0.10% of C and 2.6% of N). In roots, AA derived-¹³C was more
- concentrated in AAs than in total plant matter (0.70 vs 0.28% of C) (Table 1). When translocated to
- leaves, AA derived-¹³C concentration in AAs decreased to reach that of the bulk leaf matter (0.10% of
- 242 C) (Table 1).

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- Among the measured AAs in plant, alanine and phenylalanine were more abundant by more than a factor
- 244 10 relative to methionine (Table 1). However, alanine was not more enriched in ¹³C than most of the
- 245 AAs (fig. 3). Instead, phenylalanine and methionine were significantly more enriched in ¹³C than other
- 246 AAs in roots, stems and leaves (fig.4)

3.3. Concentrations of phytoliths, phytC, phytN, AA derived-¹³C and AA derived-¹⁵N in phytC and phytN

- Silica content measured by ICP-AES accounted for 0.08% and 0.26% of the dry weight (d.w.) of roots
- 250 and stems/leaves respectively, which is lower than the >1% d.w. concentration previously measured for
- 251 this species harvested 8 weeks after sowing (Hartley et al., 2015). This is possibly due to the fact that
- 252 the plants did not reach maturity or/and that the volume of the roots in contact with the Si-enriched
- solution was small, limited by the RHYZOtest configuration. Most of the stems and leaves silica was in
- 254 the form of phytoliths (0.19% d.w) (Table 2) that constituted a morphological assemblage characteristic
- of the Festucoideae grass subfamily which Festuca arundinacea belongs to (Honaine et al., 2006) (fig.
- 4). As expected, root phytoliths were not abundant enough to be quantified. PhytC represented 0.51%
- d.w. of phytoliths (Table 2), which is in the range of values previously measured for phytC (Santos et
- al., 2010; Alexandre et al., 2015; Reyerson et al., 2015). Occluded N (phytN) accounted for 0.10% d.w.
- of phytoliths.
- 260 Phytoliths were slightly more enriched in 13 C (13 C-excess of $0.026 \pm 0.002\%$) than the leaves (13 C-excess
- of $0.017 \pm 0.001\%$) (p-value<0.05) (Table 1). The AA derived- 13 C concentration in phytoliths
- represented $0.15 \pm 0.01\%$ of phytC which is low but in the same order of magnitude than the
- 263 concentrations in the bulk matter and AAs of stems and leaves (0.10 ± 0.003) (Table 1). The AA derived-
- ¹³C/AA derived-¹⁵N ratio in phytoliths was low (0.8) but higher than in bulk stems and leaves (0.3)
- 265 (Table 1; fig. 2).

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4. Discussion

4.1. Plausible forms of AA derived-¹³C and AA derived-¹⁵N absorbed and translocated

Festuca arundinacea may have absorbed AA derived-¹³C and AA derived-¹⁵N from the labeled solution

in multiple organic and inorganic forms, as detailed below. The AA derived-¹³C/¹⁵N ratios of roots and

leaves (0.8 and 0.3, respectively) that were much lower than C/N ratios of the supplied AAs (from 2.6

to 7.7) suggested that the grass absorbed most of ¹⁵N from already mineralized N in the tank. Indeed,

272 under non-sterile conditions, microbial activity around roots can biodegrade the AAs in a range of hours

(Jones et al., 2005; Kielland et al., 2007; Jones et al., 2009) and produce derived metabolites plus mineralized N and CO₂ (Biernath et al., 2008; Rasmussen et al., 2010). Regarding C, both organic and inorganic forms may enter into the plant (Biernath et al., 2008; Rasmussen et al., 2010). Inorganic C can be transported through the plant passively, in link with transpiration (Vuorinen et al., 1989) and contribute to the carbon budget of the leaves through decarboxylation of the dissolved CO2 and photosynthetic refixation of released CO₂ (anaplerotic fixation; Viktor and Cramer, 2005). Both organic and inorganic compounds can thus be used in the build-up of new molecules or as energetic resources (e.g. Näsholm et al., 2009), or be lost through respiration (Gioseffi et al., 2012) or exudates (e.g. Jones and Darrah, 1993). Organic C can also enter the plant as intact molecules, such as AAs (Sauheitl et al., 2009: Whiteside et al., 2009), and be either translocated by AA transporters or subject to deamination. At least, rhizospheric and endophytic microorganisms, that acquired their labeled signature from the labeled solution, may also account for the AA derived-13C and AA derived-15N recovery in the plant. The present labeling experiment does not allow to precisely trace the form under which the AA derived-¹³C and AA derived-¹⁵N were absorbed and fixed in roots, stems and leaves, as recently done using a position-specific C and N labeling technique (Moran-Zuloaga et al., 2015). All the processes described above may have occurred jointly. The significant decrease of ¹³C-excess (or AA derived-¹³C concentration) in AAs from roots to leaves (p-value<0.05) (fig. 2 and 3), suggested either an uptake and fixation of organic ¹³C, or an anaplerotic fixation of inorganic ¹³C in the roots themselves. However, the fact that ¹³C was more concentrated in the extracted AAs than in bulk roots and stems/leaves (Table 1; fig. 2), and further, that ¹³C-excess values of methionine and phenylalanine were significantly higher than ¹³C-excess values of other AAs in the roots and stems/leaves (fig. 3), supported that a small amount of AA¹³C entered the plant and was subsequently translocated and fixed in roots and stems/leaves in its original molecular form.

4.2. AA-¹³C fixation in phytoliths

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In agreement with the radiocarbon evidence for soil C occlusion in phytoliths (Reyerson et al., 2015), AA derived-¹³C accounted for a measurable part of phytC. The phytolith C/N value (5.0) was close to a value previously measured in cultivated wheat phytoliths (3.7; Alexandre et al., 2015) and in the range of C/N values characteristic of AAs (4-5, Jones et al., 2009). However, the AA derived-¹³C/¹⁵N ratio (0.8) was far from this range. Thus, although our experiment allowed to trace for the first time that C absorbed by grass roots can feed the C ultimately fixed in organic compounds subject to occlusion in stem and leaf phytoliths, the forms under which AA derived-¹³C entered the plant, was translocated and ultimately occluded in phytoliths still remain unknown.

Previous Nano Secondary Ion Mass Spectrometry (NanoSIMS) investigation of phytC indicated that at least part of phytC is continuously distributed in the silica structure, at the sub-micrometric scale (Alexandre et al., 2015). This has been further supported by Raman spectroscopy mapping (Gallagher

et al., 2015). The process of silica precipitation has been investigated by environmental scanning electron microscope (ESEM) and TEM-EDX analyses that highlighted that silica first precipitates in the inner cell wall, probably triggered by the presence of callose or lignin, then infills the cell lumen in a centripetal way, until most of the cell becomes silicified (e.g. Perry et al., 1987; Motomura, 2004; Laue et al., 2007; Law and Exley, 2011; Zhang et al., 2013). During this process, an organic template probably participates in the silica formation (Harrison. 1996; Laue et al., 2007). When the cell silicification is complete, residual organic compounds that were not already occluded probably gather in any remaining spaces within the cell and delimitate micrometric central cavities characteristic of most phytoliths (Alexandre et al., 2015). In the present case, the concentration of AA derived-¹³C (relatively to total C) in phytoliths, which is in the same order of magnitude than in leaves, supports a random fixation of AA derived-¹³C in these residual organic compounds subject to occlusion in the silica structure. There are two plausible hypotheses for this fixation. The first hypothesis is that AA derived-¹³C may be associated with Si when absorbed by the roots, translocated in the plant and introduced into the cells. However, ²⁹Si NMR spectroscopy of ²⁹Si-labeled exudate of wheat xylem previously indicates only the occurrence of the dissolved forms of Si (Casey et al., 2004). Although this does not preclude the subsequent formation of organo-silicate complexes it weakens the hypothesis of Si and C being associated since their uptake by the roots. Additionally, in our experiment, the roots that contain the lowest amount of Si also contain the highest amount of AA derived-¹³C which is not in agreement with AAderived-¹³C and Si being associated when absorbed by the roots. The second hypothesis is that AA derived-¹³C may be isolated as an unwanted substance in cell vacuoles and subsequently trapped in the silica structure. In order to check this hypothesis we used D-ALA, expected to be less metabolized than L-AAs, although recent investigations suggest that plants are able to utilize D-AAs at rate comparable to those of other N forms (Hill et al., 2011). D-ALA was not specifically taken-up or retained as an intact molecule (fig. 3) and cannot account for the AA derived-13C measured in phytC. D-ALA may thus not be appropriate for tracing unwanted substances. Further investigations, including the use of spectroscopies relevant for characterizing phytC at the molecular level, are necessary to support or refute the above hypotheses.

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4.3. Implication for our understanding of the C cycle at the plant-soil interface

In the experiments presented here, the net uptake of AA derived-¹³C by Festuca arundinacea represented 4.5% of AA derived-¹³C supplied to the nutrient solution, part of it being absorbed as intact AA molecules (here methionine and phenylalanine). AA derived-¹³C fixed in the plant represented only 0.13% of total C, the root absorption of AA derived-¹³C being clearly marginal compared to photosynthesis. The present experiment was done in nutrient solution and its relevance for soil C uptake assessment is therefore limited. It may underestimate the extent of the process under natural and field conditions. Indeed, AAs uptake was shown to be inhibited by the high concentrations of mineral N preferred by the plant (Paungfoo-Lonhienne et al., 2008. Sauheitl et al., 2009. Gioseffi et al., 2012). In

the present case, mineral N may come from KNO₃ initially presents in the original nutrient solution and from the supplied AAs dissociated by microbes, mycorrhizas or root exsudates (Paungfoo-Lonhienne et al., 2008; Keiluweit et al., 2015). The maximum AA content in the growth solution (0.2 mmol/L) was higher than AAs concentrations that have been measured in soil solutions (from 0 to 0.1 mmol/L; Vinolas et al., 2001; Jämtgård et al., 2010), thus potentially inhibiting AAs uptake. Additionally, due the RHIZOtest configuration, roots were confined to a small volume, and their contact with the renewed labeled solutions and AA derived-¹³C available for uptake was limited. Given the above considerations, the use of the AA derived-¹³C concentrations, experimentally measured, as a proxy of soil derived-C concentrations should be considered with caution.

However, to gain a rough idea on the order of magnitude of the C flux that may occur from soil to plant, at the ecosystem scale we used the 0.13% AA derived-¹³C concentration in plants obtained in the present experiment and extrapolated that value to the grassland ecosystem scale. Grasslands cover a global surface of 2.4 10⁹ ha (Scurlock and Hall. 1998) and are characterized by a Net Primary Production (NPP) ranging from 7 to 20 10⁹tC/ha/yr (Scurlock et al., 2002). The global grassland productivity thus ranges from 16.8 to 48 109tC/vr. The obtained flux of AA-derived C absorbed by grasses then would range from 21.8 to 62.4 10^6 tC/yr. This is nonsignificant when compared to the 2.6×10^9 tC yr⁻¹ estimate for the land C sink (IPCC Staff. 2007), or to the 0.4×10^9 tC yr⁻¹ estimate for the global long term soil C accumulation rate (Schlesinger, 1990). It is however higher than a possible CO2 phytolith biosequestration flux (e.g. Parr & Sullivan, 2005; Parr et al., 2010; Song et al., 2014). The CO₂ phytolith biosequestration concept is based on the assumptions that phytC is exclusively derived from atmospheric CO₂, and has a long residence time in soils. A recent re-examination of the CO₂ biosequestration flux by phytoliths, in the light of a lower and more realistic estimates of phytolith residence time in soils, vielded a value of 4.1×10^4 tC/yr for the world grasslands (Reyerson et al., 2015). The present study further minimizes the significance of CO₂ biosequestration by phytoliths showing that it could be counteracted by the flux of C potentially mobilized from soils by grass root uptake.

Recent experiments have demonstrated that root exudates promote a net loss of soil C previously assumed to be stable at the millennial scale thanks to its protection by mineral constituents (e.g. clays or amorphous minerals). Root exudates would stimulate microbial and fungi digestion (priming effect) (Fontaine et al., 2003; Fontaine et al., 2011) and promote dissolution of the mineral phase through oxalic acid production (Keiluweit et al., 2015). From the present experiment, we suggest that direct uptake of soil derived-C by roots, in conjunction with the N uptake, should be accounted for when investigating the role of roots in soil C mobilization.

5. Conclusion

In agreement with previous studies, the present labeling experiment supports that C absorbed by grass roots and allocated to stems and leaves preserve in a small extent its original organic molecular form (here methionine and phenylalanine). Moreover, the experiment shows for the first time that AA derived-C absorbed by grass roots and allocated to stems and leaves can partly feed the C ultimately fixed in organic compounds subject to occlusion in stem and leaf silica. Further analyses are required to identify the form in which AA derived-C and more generally phytC is occluded. Our findings complements previous radiocarbon evidence of soil C contribution to phytC (Santos et al., 2012; Reyerson et al., 2015) and raise questions about the mechanisms that drive soil C mobilization by plant roots, for a better

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understanding of soil/plant interactions involved in the terrestrial C cycle.

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497 **Captions**

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- Figure 1: The labeling experiment in the growth chamber. (A) The two labeled tanks are connected to
- 499 the solution containers. A perilstatic pump facilitates the solution renewals. (B) A platform is sealed to
- each tank. (C) Plant-receiving pots are cylinders closed at the bottom with a polyamide mesh. (D)
- Twenty-four plant-receiving pots are inserted into each platform. (E) In each pot. seeds are covered with
- agar-agar to limit gas exchanges.
- Figure 2. Concentration of AA derived-¹³C and AA derived-¹⁵N in bulk matter, phenylalanine (PHE)
- and methionine (MET) of roots, stems and leaves and phytoliths of Festuca arundinacea grown in labeled
- tanks (in % of bulk C, N, PHE and MET respectively).
- Figure 3. Concentration (A) and ¹³C-excess (B) of amino acids (AAs) measured by GC-IRMS in roots,
 - stems and leaves. Bars stand for one standard deviation of 4 replicates (2 tanks x 2 AAs extractions).
- Figure 4. Natural light microscopy image of the phytolith assemblage produced by the stems and
 - leaves of Festuca arundinacea dominated by (e) the elongate type and (gsc) the grass short cell
- trapeziform type (Madella et al., 2005).

Table 1: Allocation of AA derived-¹³C and AA derived-¹⁵N in Festuca arundinacea grown in labeled and control solutions. Mean values and standard deviations (numbers in brackets) are given. The numbers in bold refer to the percentage of the individual AA applied. Un. is for unanalyzed.

	Dry weight Total elements			Isotopic composition			Label concentration		Recovery of label from solution			Label net uptake			
		[C]	[N]		$\delta^{13}C$	¹³ C-excess	$\delta^{15}N$	15N-excess	[AA- ¹³ C _{sample}]	$[AA^{-15}N_{sample}]$	Recovery AA- µgC/g d.wt	Recovery AA-15N	AA-13C/15N	AA-13C	AA-15N uptake
	g	mg/g o	1.wt	C/N	‰	Atom%	‰	Atom%	% plant C	% plant N	μgC/g d.wt	$\mu g N/g \; d.wt$		% supplied C	%supplied N
Control tank															
Arial Part	16.2	389.4	43.4	9.0	31.7		13.8								
Phytolith	0.03	0.0	0.0	5.2	28.8		3.2								
Roots	4.4	270.3	63.9	4.2	31.0		14.5								
Labeled tank-Aerial															
Total	23.9	395.0	46.8	4.9		0.017		0.338	0.10	2.6	385.0	1201.7	0.3	2.86	37.94
	(0.5)	(15.0)	(4.0)			(0.001)		(0.047)	(0.003)	(0.4)	(1.7)	(66.4)		(0.05)	(1.23)
Phytolith	0.040	5.1	1.05	4.9		0.026		0.118	0.15	0.9	0.02	0.0002	0.8	2 10-7	1 10-8
	(0.001)	(0.9)	(0.07)	(1.2)		(0.002)		(0.038)	(0.01)	(0.3)	(0.003)	(0.00006)		(4 10-8)	(3 10-9)
Sum AAs		52.5	13.2	4.0		0.020			0.10	N.D.	50.7	N.D.		0.38	Un.
		(6.4)	(1.6)			(0.009)			(0.000)		(0.0)			(0.000)	
Phenylalanine (PHE)		5.4	0.7	7.7		0.038			0.20	N.D.	12.1	N.D.		0.18	Un.
		(0.5)	(0.1)			(0.010)			(0.000)		(3.2)			(0.047)	
Methionine (MET)		0.3	0.1	4.3		0.187			1.10	N.D.	3.2	N.D.		0.15	Un.
		(0.2)	(0.0)			(0.027)			(0.200)		(0.5)			(0.022)	
Alanine (ALA)		4.7	1.8	2.6		0.012			0.07	N.D.	3.3	N.D.		0.07	Un.
		(1.1)	(0.4)			(0.001)			(0.000)		(0.3)			(0.006)	
Labeled tank- Roots															
Total	4.9	376.0	34.3	11.0		0.048		0.534	0.28	4.1	1065.0	1387.2	0.8	1.61	8.98
	(0.5)	(13.0)	(3.6)			(0.000)		(0.100)	(0.002)	0.8	(43.6)	(117.5)		(0.24)	(1.74)
Sum AAs		52.0	14.6	3.6		0.112			0.70	N.D.	347.0	N.D.		0.53	Un.
		(5.3)	(1.5)			(0.013)			(0.100)		(49.6)			(0.075)	
Phenylananine (PHE)		2.3	0.3	7.8		0.499			2.90	N.D.	66.9	N.D.		0.20	Un.
		(0.2)	(0.0)			(0.155)			(0.900)		(20.8)			(0.063)	
Methionine (MET)		0.3	0.1	4.3		0.546			3.20	N.D.	9.8	N.D.		0.10	Un.
		(0.1)	(0.0)			(0.049)			(0.300)		(0.9)			(0.009)	
Alanine (ALA)		3.8	1.5	2.6		0.101			0.60	N.D.	22.6	N.D.		0.10	Un.
		(0.2)	(0.1)			(0.005)			(0.000)		(1.2)			(0.005)	
Labeled tank-total															
Total plants	28.8	391.5	44.7	8.8		0.022		0.371	0.13	2.8	500.7	1232.3	0.4	4.5	46.9
		(14.1)	(4.1)			(0.000)		(0.060)	(0.001)	(0.4)	(22.0)	(79.0)		(0.197)	(3.0)

Table 2. Concentration of phytoliths. phytolith occluded C ([PhytC]) and phytolith occluded N ([PhytN]) in Festuca arundinacea grown in labeled and control solutions. Numbers in italics refer to the standard deviation associated with the averaged values (one value per labeled tanks).

	Phytolith	[PhytC]	[PhytN]			
	%d.wt plant	% d.wt phytolith				
Control tank						
Aerial part	0.102	0.88	0.17			
Roots	0.075					
Labeled tank						
Aerial parts	0.19	0.51	0.1			
	(0.002)	(0.08)	(0.007)			
Roots	0.014					
	(0.01)					

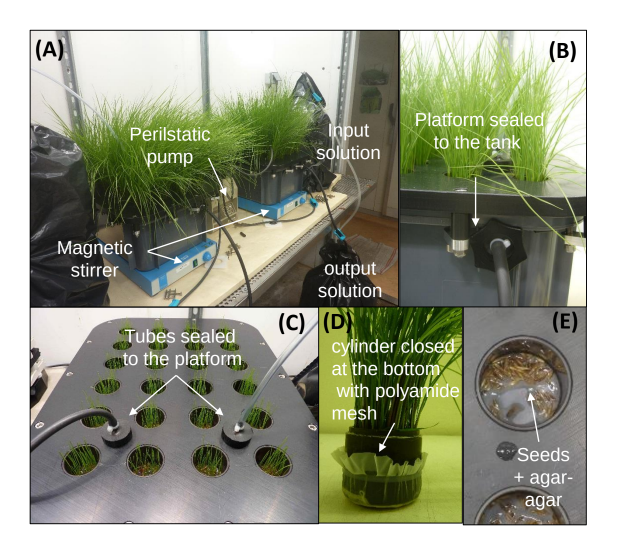


Figure 1

Phytoliths



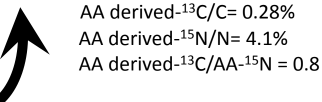
AA derived- 13 C/C= 0.15% AA derived- 15 N/N= 0.9% AA derived- 13 C/AA- 15 N = 0.8

Stems Leaves



AA derived- 13 C/C= 0.10% AA derived- 15 N/N= 2.6% AA derived- 13 C/AA- 15 N = 0.3 AA derived- 13 C/C_{PHE}: 0.2% AA derived- 13 C/C_{MET}: 1.1%

Roots



AA derived- 13 C/C_{PHE}: 2.9% AA derived- 13 C/C_{MET}: 3.2%

Concentration

AA derived- 13 C/C= 0.13% AA derived- 15 N/N = 2.8% AA derived- 13 C/AA- 15 N = 0.4

Solution

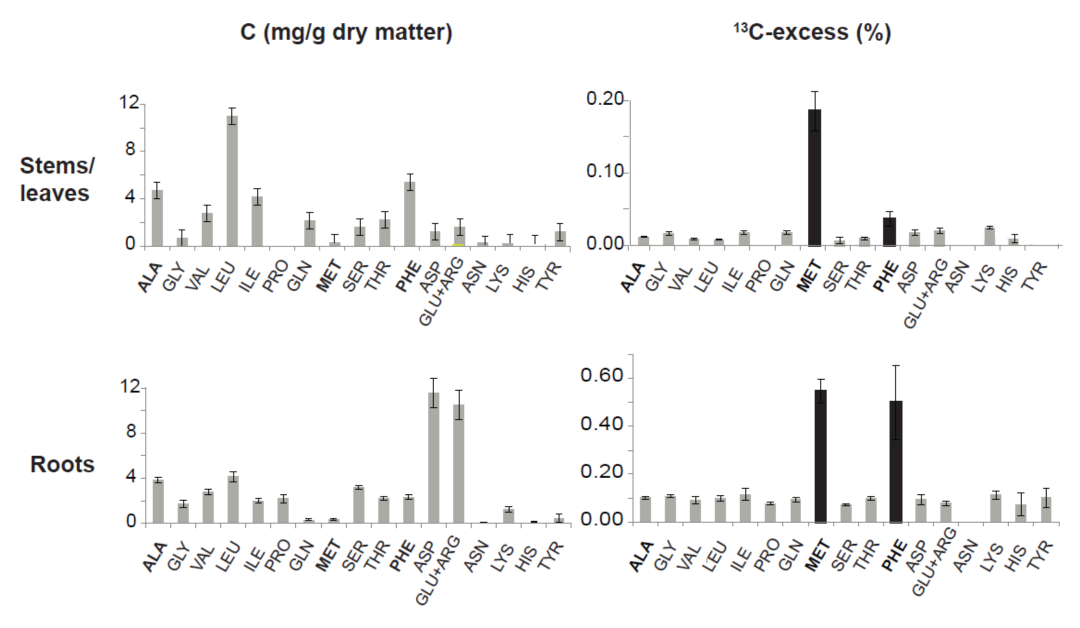


Figure 3

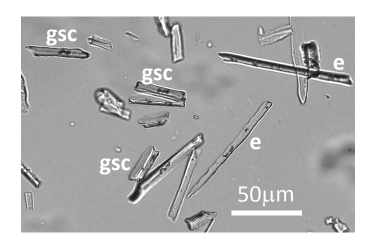


Figure 4