Weathering by tree root-associating fungi diminishes under simulated Cenozoic atmospheric CO₂ decline

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Abstract

Trees dominate terrestrial biotic weathering of silicate minerals by converting solar energy into chemical energy that fuels roots and their ubiquitous nutrient-mobilising fungal symbionts. These biological activities regulate atmospheric CO$_2$ ([CO$_2$]$_a$) over geologic timescales by driving calcium and magnesium fluvial ion export and marine carbonate formation, but the important stabilising feedbacks between [CO$_2$]$_a$ and biotic weathering anticipated by geochemical carbon cycle models remain untested. We report experimental evidence for a negative feedback across a declining Cenozoic [CO$_2$]$_a$ range from 1500 ppm to 200 ppm, whereby low [CO$_2$]$_a$ curtails mineral surface alteration via trenching and etch pitting by arbuscular mycorrhizal (AM) and ectomycorrhizal (EM) fungal partners of tree roots. Optical profile imaging using vertical scanning interferometry reveals changes in nanoscale surface topography consistent with a dual mode of attack involving delamination and trenching by AM and EM fungal hyphae on phyllosilicate mineral flakes. This is consistent with field observations of micropores in feldspar, hornblende and basalt, purportedly caused by EM fungi, but with little confirmatory evidence. Integrating these findings into a process-based biotic weathering model revealed that low [CO$_2$]$_a$ effectively acts as a “carbon starvation” brake, causing a three-fold drop in tree-driven fungal weathering fluxes of calcium and magnesium from silicate rock grains as [CO$_2$]$_a$ falls from 1500 ppm to 200 ppm. The feedback is regulated through the action of low [CO$_2$]$_a$ on host tree productivity and provides empirical evidence for the role of [CO$_2$]$_a$ starvation in diminishing the contribution of trees and mycorrhizal fungi to rates of biological weathering. More broadly, diminished tree-driven weathering under declining [CO$_2$]$_a$ may provide an important contributory mechanism stabilising Earth’s [CO$_2$]$_a$ minimum over the past 24 million years.
1 Introduction

Throughout their evolutionary history, trees have formed ubiquitous symbiotic partnerships with either arbuscular mycorrhizal (AM) or more recently evolved ectomycorrhizal (EM) soil fungi, with both roots and their associated fungal partners recognised as the dominant biotic drivers of terrestrial mineral weathering (Blum et al., 2002; Bonneville et al., 2009; Cochran and Berner, 1996; Landeweert et al., 2001; Quirk et al., 2012; Rosling et al., 2004; Taylor et al., 2009). The activities of both AM and EM fungi are fuelled by delivery of photosynthate from the host plant or tree belowground where extensive networks of root-associating hyphae enhance essential nutrient element mass transfers from solid and liquid phases in soils to plants through greater intimacy of mineral contact and absorptive surface areas than roots (Smith and Read, 2008; Taylor et al., 2009). The mechanistic basis underpinning mycorrhizal hyphal-mineral interaction is being resolved at the nanometre scale with observations demonstrating structural alterations and mineral surface trenching through the effects of organic ligand exudation, such as siderophores and low molecular weight organic compounds, proton extrusion, and cation uptake (Bonneville et al., 2009; Gazzè et al., 2012; Saccone et al., 2011). Such effects are most strongly expressed in EM fungi, which molecular clocks indicate originated 220–135 million years (Ma) ago (Smith and Read, 2008; Taylor et al., 2009), more than 200 Ma after ancestral AM fungi (Brundrett, 2002; Taylor et al., 2009).

Biological weathering of silicate minerals by roots and mycorrhizal fungi is linked conceptually, and through experimental evidence, to primary productivity and photosynthate fluxes belowground from the canopies of the host trees supporting the fungal symbionts (Cochran and Berner, 1996; Rosling et al., 2004; Smits et al., 2012; Taylor et al., 2009). Under atmospheric CO$_2$ ([$\text{CO}_2$]$_a$) enrichment (concentrations of $\sim$600 ppm or higher), delivery of photosynthate to both AM and EM fungal partners can increase, fuelling expansion of fungal networks, increased delivery of labile carbon into soils, and for EM partnerships, greater exudation of low molecular weight organic compounds and chelating agents (Alberton et al., 2005; Drigo et al., 2010, 2013; Fransson, 2012;
Johansson et al., 2009; Pritchard et al., 2008). Intensified exploitation, alteration, and chemical dissolution of mineral resources by mycorrhizal networks of trees under increasing [CO$_2$]$_a$ likely accelerated Earth’s biological weathering engine and global CO$_2$ sequestration to regulate climate during geologic intervals of elevated [CO$_2$]$_a$ throughout the Phanerozoic (past 400 Ma) (Berner et al., 1983; Taylor et al., 2009, 2012; Volk, 1987). However, the effects of the Cenozoic [CO$_2$]$_a$ decline from 1500 ppm to 200 ppm beginning 35 Ma ago (Beerling and Royer, 2011; Pagani et al., 2011) on photosynthate-fuelled weathering processes has not been investigated experimentally.

Geochemical carbon cycle models (Berner et al., 1983, 2006; Volk, 1987) invoke stabilising feedbacks between [CO$_2$]$_a$ and weathering by forests and mycorrhizal fungi, whereby falling [CO$_2$]$_a$ leads to diminished Ca$^{2+}$ and Mg$^{2+}$ ion release from terrestrial silicate rocks into the oceans, which reduces marine carbonate deposition, and lowers the long-term sink for [CO$_2$]$_a$ (Pagani et al., 2009). Critically, these models are not underpinned by the empirical evidence required to resolve the effects of low [CO$_2$]$_a$ on biological weathering processes. This includes understanding of tree-mycorrhiza functioning during episodes of [CO$_2$]$_a$ drawdown to within the narrow range of 180–200 ppm that defines Earth’s Cenozoic [CO$_2$]$_a$ minimum over the past 24 Ma.

We report results from controlled-environment growth chamber experiments testing the hypothesised [CO$_2$]$_a$ regulation of fungal mineral weathering mediated by AM and EM tree species. We grew saplings of a gymnosperm, coast redwood (Sequoia sempervirens D. Don.) and angiosperm, field maple (Acer campestre L.) that form AM partnerships, alongside an angiosperm, silver birch (Betula pendula Roth.) which associates with EM fungi. These species were selected because they belong to families with well characterised mycorrhizal symbioses and molecular clock phylogenetic ages dating from the mid Cenozoic (> 36 Ma ago) or earlier (Crisp and Cook, 2011; Wikström et al., 2001). Saplings were cultivated in individual pots alongside plant free control treatments at three closely regulated (±4 %) [CO$_2$]$_a$ concentrations of 1500 ppm, 500 ppm and 200 ppm, encompassing the upper and lower bounds of the Cenozoic (Beerling and Royer, 2011), and otherwise identical growth conditions.
To isolate the effects of mycorrhizal hyphae on the silicate rocks and minerals, we buried uniform-sized grains of basalt as a source of plant-essential macronutrients (phosphorus, iron, calcium and magnesium) and flakes of potassium-bearing muscovite within each pot in 35 µm pore-size mesh-walled bags that exclude roots but permit colonisation by fungal hyphae. We measured the lengths of hyphae that were in direct contact with the basalt grains and corrected them for background non-mycorrhizal fungi observed in plant free control treatments as an index of AM and EM network size and hyphal interaction with rocks and minerals. We employ vertical scanning interferometry to optically profile the surface micro-topography of the muscovite flakes, enabling us to characterise the physical alteration of the mineral surface following active colonisation by mycorrhizal fungi.

2 Methods

2.1 Experimental design

One-year-old *Sequoia sempervirens*, *Acer campestre* and *Betula pendula* saplings were obtained from a UK nursery and cultivated alongside plant free controls in pots (18 cm diameter; approx. 4.5 L) filled with 1 : 1 (v : v) Chelmsford “52” sand (WBB Minerals, Cheshire, UK) and potting compost (Levington M3, Scotts, Ipswich, UK) mixed with ∼1 % (w : v) mycorrhizal inoculum consisting of soil and root material obtained from beneath mature stands of corresponding species at the National Arboretum, Westonbirt, UK. This ensured that tree root systems were exposed to natural consortia of mycorrhizal fungi and other soil microorganisms. Prior to potting, fresh weights of the trees were taken following removal of adhering compost from the roots. The substrate in plant free control treatments also contained the mycorrhizal inocula. The trees (n = 3 per species per [CO$_2$]$_a$ treatment) were randomly distributed between matched controlled-environment growth chambers (Conviron BDW40, Controlled Environments Ltd. Manitoba, Canada) and grown at 200 ppm, 500 ppm or 1500 ppm [CO$_2$]$_a$ and oth-
otherwise constant growth conditions of 80 % relative humidity, 18/20°C night/day cycle with a 14 h photoperiod. Light intensity at the canopy was 200 µmol m$^{-2}$ s$^{-1}$ for the entirety of the 14 h photoperiod and duration of the six month growth period (Supplement Table S1).

The low [CO$_2$]$_a$ treatment was maintained by routing the chamber air inlet via a filter unit packed with sodalime (Sofnolime, Molecular Products, Essex, UK) fixed to the exterior of the chamber, that scrubbed CO$_2$ from the air. The [CO$_2$]$_a$ level was controlled at set-point using a CO$_2$ sensor (Carbocap GMP242, Vaisala, Finland) linked to a feedback system regulating chamber air inlet between the sodalime scrubbing unit and bursts of ambient air, and was independently verified with an infra-red gas analyser (IRGA) (Li-COR, LI6400, Homburg, Germany). The sodalime was replaced monthly before chamber [CO$_2$]$_a$ began to increase. All trees were of comparable biomass prior to their introduction to experimental conditions (Supplement Table S2). Pots were well watered with reverse osmosis water and their chamber positions rotated weekly.

2.2 Calculated main stem biomass gain

We used digital photographs of each tree taken at the start of the experiment and again after the six month growth period to obtain stem length and diameter at the base and crown (highest point of main stem) with ImageJ 1.43 software (National Institutes of Health, Bethesda, Maryland, USA). Stem volume was calculated using $V = \left(\pi/12\right)(b^2 + bc + c^2)$, where $l$ is stem length, $b$ is stem diameter at the base, and $c$ is stem diameter at the crown. Pre- and post-treatment stem biomass was estimated using measured mean stem density values obtained for each species at the end of the experiment (Sequoia: 0.73 g cm$^{-3}$ ± 0.08; Acer: 0.60 g cm$^{-3}$ ± 0.10; Betula: 0.41 g cm$^{-3}$ ± 0.11).

2.3 Root-excluding mesh bags

Tertiary basalt from Northern Ireland was crushed using a hammer and ball mill, sieved into two size fractions (0.25–0.50 mm and 0.50–1.00 mm), rinsed in deionised wa-
ter (1:100 w:v) until the water was visibly clear (approx. eight rinses) and dried at 50°C. The two grain size fractions were weighed into 2.5 g portions and sealed into 50 × 50 mm right-angled triangle mesh-walled bags (35 µm pore-size woven nylon; Plastok Associates Ltd. Birkenhead, Wirral, UK) using a heat-sealer to melt and fuse the edges. The mesh walls permitted fungal hyphae and soil pore waters to penetrate and interact with the contents, but prevented access by roots. Duplicate basalt-filled mesh bags were inserted into each pot to a depth of five cm and recovered after 185 days of incubation. Previously published X-ray fluorescence data for the basalt used here highlight its importance as a source of macronutrient elements for the trees and mycorrhizal networks in our experiment: P₂O₅ (0.29 % wt), CaO (10.37 % wt), MgO (8.10 % wt), K₂O (0.29 % wt) and Fe₂O₃ (11.93 % wt) (Quirk et al., 2012).

2.4 Hyphal networks colonising rock grains

Basalt sub-samples (2 g) from mesh bags were sonicated for 10 min (Branson B32 sonic bath, Danbury CT, USA) in a 100 mL conical flask with 30 mL of deionised water to release adhering hyphal strands from the grains into solution. Aliquots (5 mL) of the suspension were filtered through 25 mm diameter, 0.45 µm pore size gridded cellulose Whatman® membrane filters. The retained hyphal strands were stained with two mL of Trypan blue lacto-phenol (1.6 mg 100 mL⁻¹ Trypan Blue (C₃₄H₂₄N₆O₁₄S₄Na₄), 200 mg 100 mL⁻¹ phenol (C₆H₅OH), 25 % lactic acid (C₃H₆O₃), 50 % glycerol (C₃H₅(OH)₃) and 25 % deionised water) for 10 min. Excess stain was rinsed through the filter apparatus with deionised water and hyphal strands were measured using a modified line-intersect technique at 200 × magnification (Wallander et al., 2004).

Detailed microscopy of the hyphal strands from Sequoia and Acer treatments confirmed the typical anatomical features of AM fungi, including ready staining under Trypan Blue lacto-phenol, angular projections, terminal spores and the general absence of septa (Smith and Read, 2008). From Betula treatments, higher, predominantly Basidiomycotean fungi that often form EM partnerships with roots were identified through
diagnostic features such as darker pigmentation and melanisation of the hyphal walls and the presence of septa and clamp connections (Smith and Read, 2008). We determined the extent of active hyphal networks by correcting for the background hyphal lengths associated with the basalt in plant free control pots that represent free-living, saprotrophic fungal populations.

### 2.5 Characterisation of mineral surface alteration

Mirau vertical scanning interferometry (VSI) uses white light reflected from a reference mirror that combines with light reflected from a sample surface to produce interference fringes, where best contrast interference occurs at the point of best focus. This “interference” signal is measured at fixed nanoscale intervals as the vertical axis of the instrument moves through focus. As white light has a short coherence length, interference fringes are present only over a shallow depth for each focus point. The vertical position corresponding to peak interference is used to calculate sample surface heights with nanometre vertical resolution (Lüttge et al., 1999). Moreover, using light to characterise surface topography avoids many of the analytical artefacts resulting from the physical mineral surface probing involved in atomic force microscopy, for example (Buss et al., 2007). Each scan produces an \(XY\) data array where each pixel corresponds to surface height \(Z\) relative to the mean surface plane where \(Z\) is equal to zero.

Here, approx. four flakes of muscovite (\(\sim 4 \times 4\) mm) from Krantz (Bonn, Germany) were embedded in silicone (commercially available silicone sealant) and mounted on 26 \(\times\) 4 mm glass “VSI slides” (Supplement Fig. S1). Muscovite is an aluminium phyllosilicate mineral ideally suited to such investigations because it yields near atomically smooth cleavage surfaces. Prior to burial, the surface of each muscovite flake was cleaned first using \(\sim 200 \mu\text{L} 1\%\) sodium dodecyl sulphate (SDS) and lens tissue, followed by lens tissue wetted with deionised water (Buss et al., 2007). The VSI slides were mounted on an automated stage platform and randomly selected localities (500 \(\times\) mag., 126 \(\times\) 94 \(\mu\)m scanned area, 640 \(\times\) 480 pixel array) on the surface of each mineral flake were scanned using a Wyko NT9100 VSI instrument (Bruker AXS,
Madison WI, USA) positioned on a vibration-minimising air table. We obtained the root mean square (RMS) roughness (Buss et al., 2007) of each surface location \((n = 2\) surface locations on each piece of muscovite) using the instrument’s analytical software (Vision 4.10, Bruker AXS, Madison WI, USA) following Eq. (1):

\[
\text{RMS roughness} = \sqrt{\frac{1}{MN} \sum_{j=1}^{M} \sum_{i=1}^{N} Z^2 (x_i, y_j)}
\]  

(1)

where \(M\) and \(N\) are the number of data points in the \(X\) and \(Y\) directions of the data array (640 x 480 pixels), and \(Z\) is the surface height relative to the reference mean plane. As the height values are squared, the development of low points, such as trenches, on the mineral surface over the experiment can be resolved more easily (Buss et al., 2007). Each scan location has known stage coordinates that, in combination with the automated platform and slide mount, allow precise relocation and re-characterisation of surface topography at each scan location after burial in the pots.

The VSI slides were buried along with 0.5 g of crushed basalt grains (0.50–1.00 mm grain size) in parallel with the root-excluding mesh bags used to study hyphal colonisation of basalt grains. After 185 days the slides were recovered, cleaned with lens tissue wetted with 1 % SDS and deionised water as described above and their RMS roughness re-measured. We used the ratio of the post-experiment and pre-experiment RMS roughness to calculate a metric of muscovite mineral surface alteration associated with each pot in the experiment, including the plant free control pots \((n = 3\) per species, per \([\text{CO}_2]_a\)). We corrected the roughness ratios obtained for each tree by subtracting the muscovite roughness ratios obtained from samples in plant free control pots, which were independent of \([\text{CO}_2]_a\) (200 ppm: 2.0 ± 0.5; 500 ppm: 2.1 ± 0.8; 1500 ppm: 2.1 ± 0.2; \(F_{2,6} = 0.03; P = 0.967\)) and demonstrates a degree of surface roughness increase under plant free conditions associated with soil pore waters and free-living populations of microorganisms.
2.6 Fungal trenching of muscovite

Once the post-experiment RMS roughness measurement was made, each piece of muscovite was analysed for evidence of hyphal trenches, identified as linear features with characteristics consistent with fungal hyphal morphology, such as diameter, branching patterns, angular projections and terminal spore formation. Where hyphal trenching had occurred, measurement of the dimensions (width and depth) of surface trenches was undertaken using Vision 4.10 software. Two-dimensional transects were analysed at approx. 20 µm-spaced intervals at right angles to the linear trenches allowing assessment of the width at the top of the trench and maximum depth relative to the surrounding planar surface. Trench profiles exhibited a V-shaped morphology in cross-section, so we used trench width and depth measurements to estimate trench cross-sectional areas at each intersection based on the area of a triangle (area = 0.5 × width × depth). We obtained mean trench dimensions for each pot that demonstrated evidence of mineral surface trenching.

2.7 Canopy transpiration

Atmospheric CO$_2$ regulation of canopy transpiration has implications for soil moisture content and mineral weathering rates and this effect is built into the process-based weathering model framework. After five months’ acclimation, the saplings were watered to field capacity and the soil was covered in polyvinyl chloride (PVC) sheeting (Caterwrap, Wrap Film Systems, Telford, UK) to prevent evaporation from the soil surface. Over the following week, all pots were weighed twice daily to calculate canopy transpiration. The vapour pressure deficit (VPD) of each growth chamber was calculated using the mean photoperiod temperature and relative humidity (Yoder et al., 2005). Canopy conductance (G$_w$) was calculated as transpiration (mol H$_2$O tree$^{-1}$ s$^{-1}$) divided by daytime VPD (0.47 kPa) for each tree.
2.8 Process-based modelling of mycorrhizal mineral weathering

We modified a leading global weathering model (Taylor et al., 2011, 2012) to calculate weathering fluxes from the basalt-filled mesh bags for each treatment in response to \([\text{CO}_2]_a\). The model is driven by the nutrient ion uptake required to support tree net primary productivity (NPP), along with diffusion of soil CO\(_2\), as influenced by \([\text{CO}_2]_a\) and respiration, and, for EM networks only, organic acid exudation (van Hees et al., 2006). NPP was estimated from stem biomass gain, following a stoichiometric assumption that half the mass was carbon (Zhang et al., 2009), with all parts of the pot contributing equally to the nutrition of the organisms. For temperate and boreal tree species, respiration is \(\sim 6 \times \text{NPP}\) (Malhi et al., 1999), so that gross primary productivity (GPP) is \(7 \times \text{NPP}\). We assumed that organic acid exudation by EM hyphae was \(\sim 0.2\%\) of GPP (van Hees et al., 2006), and that the negative charge of each organic anion (treated as oxalate) was balanced by protons (Casarin et al., 2003).

We also defined a separate hyphosphere within the mesh bags for basalt under the influence of hyphae, which is determined by the measured hyphal lengths colonising basalt grains and a hyphal radius of 1.4 microns (van Hees et al., 2006). Like the rhizosphere, which extends up to several millimetres from fine roots of sub millimetre radius (Nye, 1981), we consider the hyphosphere to extend \(5 \times 1.4\) micrometres from the hyphal surface. This hyphosphere is acidified by the uptake of nutrient cations and, for EM hyphae only, by the protons exuded with organic anions. Because the model calculates weathering based on the pH and oxalate concentration of the water interacting with mineral surfaces following well established reaction mechanisms (Brantley, 2008), it requires moisture content and mineral surface area. We interpolated moistures content at four equally-spaced 21 h time-steps using starting and ending values for one watering cycle calculated using the canopy transpiration rates of our saplings. A geometric surface area of 72 cm\(^2\) g\(^{-1}\) derived from the particle size distribution of our basalt grains, scaled with the roughness ratio data for muscovite normalised to the 500 ppm
treatment for each species, allowed conversion of calculated weathering rates (per unit surface area) to hyphosphere weathering fluxes for each treatment.

2.9 Statistics

We verified that no differences in initial biomass between \([\text{CO}_2]_a\) treatments existed prior to the introduction of saplings into their respective growth \([\text{CO}_2]_a\) regimes using one-way ANOVA testing for \([\text{CO}_2]_a\) within each species following a general linear model factorial design in Minitab 12.21 (Supplement Table S2). AM and EM plant free-corrected hyphal colonisation of basalt grains, the plant free-corrected roughness ratio of the muscovite flakes, calculated stem mass gain, root dry weight, and total plant dry weight at the end of the experiment were all subjected to two-way ANOVA testing for effects of species type and \([\text{CO}_2]_a\). The absence of \([\text{CO}_2]_a\) effects on the muscovite surface roughness ratio in plant free control pots was verified with a one-way ANOVA testing for effects of \([\text{CO}_2]_a\). The dimensions (width, depth and cross-sectional area) of the hyphal trench features on the surface of the muscovite flakes (based on multiple transects per trench feature per flake) were averaged \((n = 3\) per species, per \([\text{CO}_2]_a\)) and subjected to one-way ANOVA testing for effects of species type.

3 Results and discussion

3.1 Hyphal colonisation of basalt rock grains

After six months under the different \([\text{CO}_2]_a\) treatments, AM and EM hyphal colonisation of basalt grains was significantly lower for all three tree species experiencing the late Cenozoic \([\text{CO}_2]_a\) minimum of 200 ppm compared to higher \([\text{CO}_2]_a\) treatments \((F_{2,18} = 6.36; P = 0.008)\) (Fig. 1a). AM fungal networks supported by *Sequoia* and *Acer* saplings decreased by 20–40% at 200 ppm \([\text{CO}_2]_a\), and EM fungal networks with *Betula* saplings decreased by 25%, relative to 500 and 1500 ppm \([\text{CO}_2]_a\). The decline in
Low CO$_2$ diminishes weathering by trees and fungi

J. Quirk et al.

3.2 Mineral surface alteration

Based on our VSI analyses of muscovite, decreasing [CO$_2$]$_a$ was significantly and non-linearly linked to lower mineral surface roughness ratios across all tree species ($F_{2,18} = 4.55; P = 0.025$). This response is particularly marked for AM Acer and EM Betula angiosperm trees (Fig. 1b), but absent from plant free controls across the same three [CO$_2$]$_a$ treatments ($F_{2,6} = 0.03; P = 0.991$). For all tree species grown at 200 ppm [CO$_2$]$_a$ mineral surface roughness ratios were zero after correcting for the values of plant free controls (Fig. 1b). By extension, these findings indicate that in soils beneath trees and forests experiencing low [CO$_2$]$_a$, physical mineral alteration by fungal hyphae would essentially be reduced to levels found in treeless and unvegetated soils. In such situations biological weathering would be limited to interactions between natural populations of free-living microorganisms and their associated soil pore waters.

VSI optical profiling allows us to directly link hyphal colonisation with mineral surface alteration features on muscovite flakes in response to varying [CO$_2$]$_a$ supply to the host trees. Under AM Sequoia trees, VSI profiling of muscovite flakes revealed branched linear trenches, $\sim$ 0.2–0.3 $\mu$m deep, that contained etch pits extending to a maximum depth of $\sim$ 0.5 $\mu$m below the adjacent unaltered surface (Fig. 2a–d, f). These trenches share close morphological similarity with Glomeromycotean (AM) fungal hyphae, including diagnostic angular projections (Fig. 2a) (Smith and Read, 2008), and...
correspond to features seen in minerals colonised by AM trees in the field (Quirk et al., 2012). Muscovite buried beneath AM *Sequoia* and *Acer* trees at 1500 ppm [CO$_2$]$_a$ also developed ~ 50 µm diameter semi-spherical depressions in the silicate crystal layers, with morphology and dimensions matching AM fungal terminal spores (Smith and Read, 2008), linked by linear channels with widths also matching those of AM fungal hyphae (2–5 µm width) (Fig. 2c, d). Spore features were independently confirmed through direct observation with conventional light microscopy (Fig. 2c inset). No such features were found in muscovite buried under the EM *Betula* saplings, consistent with higher EM-forming Basidiomycotean fungal species that produce sporulating fruit bodies rather than terminal hyphal spores. This VSI evidence challenges the recent assumption that AM fungi are passive components of Earth’s biological weathering engine (Lambers et al., 2009), and further implicates the involvement of ancestral mycorrhizal fungi in biotic weathering processes over the past 400 Ma.

In addition to alteration of the exposed upper surface of the silicate sheets, some of the trench features are consistent with physical and chemical disruption by hyphal penetration between the repeating layers of the muscovite flakes. This is confirmed using VSI imaging demonstrating fungal penetration of step features in the mineral (Supplement Fig. S2), challenging recent claims that linear trench features in minerals widely reported from field studies are of totally abiotic origin (Sverdrup, 2009). Flakes recovered from beneath EM *Betula* saplings (Fig. 2e, f) also show extensive linear trenching punctuated with deeper pits, but with wider trenches (3.7 ± 0.3 µm) than those from AM *Sequoia* (2.7 ± 0.2 µm) and *Acer* (3.2 ± 0.2 µm) ($F_{2,18} = 2.77; P = 0.072$) trees.

Trenching and pitting of minerals by fungal hyphae indicates dissolution and mass loss from the mineral, as shown for biotite and chlorite weathered by EM fungi in axenic culture with host tree seedlings (Bonneville et al., 2009; Gazzè et al., 2012; Saccone et al., 2011). Collectively, the evidence indicates that fungal attack of the mineral occurs on exposed outer surfaces and also between the sheets of muscovite. This is consistent with field observations of 3–10 µm diameter tunnels in non-phylllosilicate minerals, like feldspar and hornblende (Hoffland et al., 2003; Jongmans et al., 1997), and in silicate
rocks like basalt (Cochran and Berner, 1996), that are often ascribed to the actions of EM fungi mobilising nutrients from minerals in soils beneath forested ecosystems (Blum et al., 2002; Finlay et al., 2009; Hoffland et al., 2003) but with little confirmatory evidence. Our experiments now establish this second mode of attack by AM and EM fungi on minerals, in this case involving active delamination and internal trenching of phyllosilicates by both groups of mycorrhizal fungi (Fig. 2 and Supplement Fig. S2).

4 Stabilising low [CO₂] a feedback on biotic weathering

Critically, our results establish that AM and EM trees growing at low [CO₂] a are associated with reductions in the extent of hyphal colonisation of minerals, which is further compounded by the formation of trench features on minerals with smaller cross-sectional areas that highlight less aggressive fungal-mineral interactions (Fig. 3a). The cross sectional area of fungal trenches on the muscovite were significantly smaller at 200 ppm [CO₂] a (0.23 ± 0.03 µm²) than those at 1500 ppm [CO₂] a (0.52 ± 0.11 µm²) \( (F_{2, 18} = 5.30; P = 0.015) \) (Fig. 3a, c). This may reflect reduced demand for mineral nutrients by the trees and fungal partners under conditions of [CO₂] a starvation. Foraging mycorrhizal hyphae likely target and colonise basalt for acquisition of phosphorus and other nutrient elements from olivine, apatite or other silicate mineral inclusions in the rock. These processes are associated with the parallel dissolution of excess Ca and Mg that is not taken up by the fungi, as evidenced by the formation of calcium oxalate crystals on hyphae (Arocena et al., 2001; Tuason and Arocena, 2009). Decreased fungal colonisation of mineral grains at 200 ppm [CO₂] a is associated with diminished muscovite surface roughness ratios (Fig. 3b). The intensity of the fungal-mineral interactions indicated by the cross-sectional areas of trenches in muscovite is proportional to the surface roughness. Across all tree-mycorrhiza partnerships this relationship is linked to declining [CO₂] a with convergence to near zero surface roughness ratios at 200 ppm CO₂ (Fig. 3c).
Our findings support the recently hypothesised “top-down” control by host trees on mineral weathering through the capture and delivery of photosynthate carbon from the top of the canopy down to the roots and mycorrhizal hyphal tips in soils (Brantley et al., 2011). In all three tree species investigated, calculated host tree stem productivity ($F_{2,18} = 27.7; \ P<0.0001; \ Fig. \ 3d$), root biomass ($F_{2,18} = 5.08; \ P = 0.019$) and total tree biomass ($F_{2,18} = 6.41; \ P = 0.008$) (Fig. 4) showed significant effects of declining $[\text{CO}_2]_a$, particularly from 500 ppm to 200 ppm. Most importantly, as tree productivity declines as $[\text{CO}_2]_a$ approaches 200 ppm, AM and EM hyphal network sizes also diminish (Fig. 3d). Both hyphal network size and weathering intensity are linked along a continuum driven by the $[\text{CO}_2]_a$ treatments of the host trees, with low $[\text{CO}_2]_a$ down-regulating mineral weathering by AM and EM hyphal networks (Fig. 3a–d).

4.1 Process-based modelling of $\text{CO}_2$-driven biological weathering

Simulated total basaltic $\text{Ca}^{2+}$, $\text{Mg}^{2+}$ and $\text{Si}^{4+}$ dissolution fluxes from the hyphospheres of AM and EM trees over the course of the study decline markedly (3-fold) when $[\text{CO}_2]_a$ drops towards 200 ppm (Fig. 5a). We focus on basalt dissolution as the most important source of continental silicate-bound $\text{Ca}^{2+}$ and $\text{Mg}^{2+}$ regulating long-term $[\text{CO}_2]_a$ (Dessert et al., 2003) and because field evidence indicates AM and EM fungi of temperate trees preferentially colonise basalt grains and significantly enhance their weathering (Quirk et al., 2012). The order of magnitude greater simulated dissolution fluxes from the hyphosphere of EM Betula compared to the AM trees (Fig. 5a) is driven primarily by the greater size of EM hyphal soil networks across all $[\text{CO}_2]_a$ treatments in our study (Fig. 1a) and the exudation of oxalate from active EM hyphae. Reduced cation fluxes from the basalt at low $[\text{CO}_2]_a$ are driven by the feedback of low $[\text{CO}_2]_a$ on host tree productivity, which diminishes mycorrhizal hyphal network size and activity (Fig. 3). The simulated rates of basalt dissolution in the immediate vicinity of the mycorrhizal hyphae, and in the bulk soil, are well-validated by comparison with rates derived from field observations of catchment-scale weathering and laboratory studies (Navarre-Sitchler and Brantley, 2007) (Fig. 5b).
In general, acidification of the microenvironment surrounding the hyphae accelerates weathering with a pH drop of 2–3 units (Fig. 5b). However, smaller hyphal networks release fewer protons into pore waters and moisture films surrounding the basalt grains, which are themselves influenced by the effect of low [CO$_2$]$_a$-driven increases in canopy transpiration rates (Fig. 6). Our results indicate that the combination of smaller mycorrhizal soil networks and lower proton extrusion rates into a drier hyphosphere and soil environment under low [CO$_2$]$_a$ would create less acidic weathering conditions and lower dissolution rates of Ca$^{2+}$, Mg$^{2+}$ and other base cations. Smaller mycorrhizal soil networks under low [CO$_2$]$_a$ would effectively reduce the surface area of active hyphae interacting with mineral grains, thereby diminishing the importance of these soil microorganisms in regulating mineral weathering reactions during geologic intervals of low [CO$_2$]$_a$.

5 Conclusions

Our experimental evidence and modelling results support the anticipated negative feedbacks between trees and weathering which underpin current models of global geochemical carbon cycling over Earth’s recent geologic history (Berner, 2006; Volk, 1987). We report mechanistic details of biotic weathering processes that respond non-linearly to falling [CO$_2$]$_a$. In particular, we have shown that as the global [CO$_2$]$_a$ environment approaches a Cenozoic Earth system minimum of $\sim$200 ppm, it appears to act as a “carbon starvation” brake (Beerling et al., 2012) curtailing weathering by diminishing forest tree productivity and the associated intensity of fungal-mineral interactions. Our experiments have yet to account for a broad suite of soil types or characterise the role of soil nutrient status in limiting tree responses to high [CO$_2$] (Norby et al., 2010; Reich et al., 2006). Nevertheless, in general terms and under our experimental conditions, we have obtained evidence indicating weathering processes and rates by forest trees and symbiotic fungi are sensitive to declining [CO$_2$]$_a$. This implies terrestrial biological processes may be important in buffering substantial [CO$_2$]$_a$ and climate change.
when episodes of tectonic activity increase the long-term sink for CO$_2$ over millions of years. Such stabilising biotic feedbacks would operate in combination with the recently confirmed abiotic CO$_2$-climate weathering feedback (Zeebe and Caldeira, 2008).

Supplementary material related to this article is available online at http://www.biogeosciences-discuss.net/10/15779/2013/bgd-10-15779-2013-supplement.pdf.

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Fig. 1. $[\text{CO}_2]_a$-regulated hyphal colonisation and mineral surface alteration. (a) Mycorrhizal hyphal lengths colonising basalt grains and (b) muscovite roughness ratios, decrease with falling $[\text{CO}_2]_a$. All values are mean ±s.e.m.
Fig. 2. Alteration of silicate mineral surfaces by symbiotic fungal hyphae. (a) Linear trench features on muscovite in an AM *Sequoia* treatment at 500 ppm [CO$_2$]$_a$ exhibiting the angular hyphal projections (ap) diagnostic of Glomeromycotean (AM-forming) fungi. The area within the rectangular box in (a) is magnified and projected in 3-D in (b) to emphasise the morphology of the fungal trenches. The profile of the dashed/black transect arrow is plotted in (f). (c) Physical disruption of muscovite beneath AM *Sequoia* at 1500 ppm [CO$_2$]$_a$ associated with a terminal spore (ts), as confirmed by light microscopy (inset; scale bar is 50 µm). (d) Surface disruption, associated with a terminal spore produced in muscovite beneath an AM *Acer* sapling at 1500 ppm [CO$_2$]$_a$. (e) Surface alteration by trenching and pitting of muscovite beneath EM *Betula* saplings; the area within the rectangular box is magnified in (f) to reveal morphological details of etch pits (ep) and the solid/red transect arrow is plotted. All units are µm.
Fig. 3. \([\text{CO}_2]_a\) -regulated hyphal trenching and mineral surface alteration. Effect of decreasing \([\text{CO}_2]_a\) on the relationship between hyphal length per unit mass of basalt and (a) cross-sectional area of trenches in muscovite and (b) surface roughness ratio of the mineral. Circles are *Sequoia* (AM), triangles are *Acer* (AM) and squares are *Betula* (EM). All values are mean ±s.e.m. (c) Cross-sectional areas of hyphal trenches on muscovite flakes and surface roughness of the mineral both increase with \([\text{CO}_2]_a\). Values are means ±s.e.m. (d) Positive relationship between stem mass gain at different \([\text{CO}_2]_a\) and hyphal lengths supported by each tree species and associated with basalt grain weathering.
Fig. 4. Root biomass (a), calculated stem mass gain (b) (a proxy for net primary production) and measured total plant biomass (c) at the end of the experiment. Values are mean ± s.e.m ($n = 3$). Bars sharing the same letter are not statistically different at $P > 0.05$ (two-way ANOVA testing for effects of species and [CO$_2$]$_a$ with Tukey multiple comparisons.)
Fig. 5. Numerical simulations of [CO$_2$]$_a$-regulated mycorrhizal fungal weathering. (a) Simulated [CO$_2$]$_a$-regulated total Ca and Mg weathering flux from basalt by AM and EM fungal networks beneath each tree over the 185 day experiment (note the axis break). (b) Simulated release of Si from basalt by the hyphosphere and bulk solution interactions for trees versus calculated pH, compared with Si release rates and pH values from field and laboratory studies of basalt weathering (Navarre-Sitchler and Brantley, 2007).
Fig. 6. $[\text{CO}_2]_a$-regulated canopy conductance to water vapour. Whole canopy transpiration rates ($G_w$) increase with declining $[\text{CO}_2]_a$. Values are means ±s.e.m. ($n = 3$).