Increased Iron-Carbon Interactions Under Long-Term Acid Deposition Enhance Soil Organic Carbon Sequestration in A Tropical Forest in Southern China

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Abstract

Atmospheric acid deposition remains a widespread problem that may influence the protection of carbon (C) in soil by altering organo-mineral interactions. However, the impacts of additional acidity on organo-mineral interactions and soil C sequestration in naturally acidic tropical soils with a high content of reactive iron (Fe) phases have not been well studied. Here we sampled a nearly 10-yr field experiment with a gradient of acidity treatments (0, 9.6, 32, 96 mol H⁺ ha⁻¹ yr⁻¹ as nitric acid + sulfuric acid) to examine how acidification alters organo-mineral interactions and soil organic carbon (SOC) pools in a tropical forest in southern China. As expected, soil acidification significantly enhanced the leaching of base cations (e.g., Ca²⁺), and it also altered the solubility and composition of Fe and Al phases. The acidity treatments converted more crystalline Fe (oxyhydr)oxides to short-range-ordered phases, resulting in a large increase in Fe-bound C vs. a relatively small decrease in Ca-bound C. Overall, the acidity treatments increased the mineral-associated C stock to 32.5–36.4 Mg C ha⁻¹ vs. 28.8 Mg C ha⁻¹ in the control, accounting for 71–83% of the observed increase in total SOC stock. These findings highlight the importance of pH-sensitive geochemical changes and the key roles of Fe in regulating the response of SOC to further inputs of acid deposition even in highly weathered and naturally acidic soils. The magnitude of SOC changes observed here indicates the importance of including pH-sensitive geochemistry in Earth system models to predict ecosystem C budgets under future acid deposition scenarios.

Introduction

Since the industrial revolution, human activities such as fossil-fuel combustion and the use of chemical fertilizer have greatly increased nitrogen (N) and sulfur (S) inputs as acid rain into the terrestrial biosphere (Likens et al. 1996; Galloway et al. 2008). These acid inputs have significant effects on the function and structure of natural ecosystems (Vitousek et al. 1997). In particular, long-term acid deposition can aggravate soil acidification, which has multiple biological and geochemical impacts on soil biogeochemical cycles (Bowman et al. 2008; Meng et al. 2019). However, the net effects of acidification on the multiple biological and geochemical processes that influence soil organic carbon (SOC) cycling are complex (Averill and Waring 2018; Ye et al. 2018), and reconciling the multiple competing impacts of acidification on SOC remains an important knowledge gap with important implications for biogeochemical models.

Recent studies and conceptual frameworks underscored the importance of acid-sensitive organo-mineral interactions in regulating soil carbon (C) dynamics (Averill and Waring 2018). Soil pH plays a central role in understanding the multiple potential impacts of acidification on soil C. For example, in alkaline soils buffered by calcium carbonate (inorganic C), N-induced soil acidification accelerates calcium carbonate dissolution and releases a substantial CO₂ amount from the soil at the global scale (7.48 Tg C yr⁻¹; Zamanian et al. 2018). In Chinese croplands, Raza et al (2020) estimated a 7% (0.15 Pg C; 1.1 Mg C ha⁻¹) loss of carbonate C due to soil acidification from 1980 to 2020. In soils with alkaline to circumneutral
pH, electrostatic bridging between soil organic matter and clay mineral surfaces by divalent base cations (Ca$^{2+}$, Mg$^{2+}$) provides a primary mechanism of physico-chemical protection for SOC (Rasmussen et al. 2018; Yu et al. 2021). In these alkaline to circumneutral soils, inputs of acidity (H$^+$) are mostly buffered by replacement of base cations on ion exchange sites (Chadwick and Chorover 2001; Kirk et al. 2010), possibly releasing SOC bound to clay minerals via divalent cations. However, in acidic soils, exchangeable base cation concentrations are typically much lower, divalent cation bridging is less important for SOC protection (Rasmussen et al. 2018; Yu et al. 2021), and soil acidity is mainly buffered by the dissolution of silicates and aluminum (Al)- or iron (Fe)-bearing minerals (Johnson 2002; Ross et al. 2008; Li and Johnson 2016). The release of hydrolyzing cations (e.g., Fe$^{3+}$ and Al$^{3+}$) at low pH may favor subsequent formation of organo–mineral complexes through ligand exchange between carboxyl groups of organic matter and singly coordinated inorganic hydroxyl groups at mineral surfaces (Kaiser and Guggenberger 2000; Mueller et al. 2012). Consistent with these ideas, a recent study in a northern grassland with circumneutral pH found a large decrease in Ca-bound C vs. a slight increase in Fe-bound C following soil acidification (Ye et al. 2018). However, the impacts of additional soil acidification on organo-mineral interactions in highly weathered soils that are naturally acidic have not yet been addressed (Kuzyakov et al. 2021).

In China, about 40% of the total territory is affected by acid rain due to the continuing increase in anthropogenic activities (coal combustion, industrial emissions, and automobile exhaust) in recent decades. Particularly in southern China, acid deposition was currently estimated as 34.4 kg N ha$^{-1}$ yr$^{-1}$ and 32.6 kg S ha$^{-1}$ yr$^{-1}$ (Jiang et al. 2018), comparable with the highest levels of acid deposition in Europe and North America (Duan et al. 2016). High rates of acid deposition have increased public concern about its widespread impacts on the natural ecosystems in southern China (Liu et al. 2013). Forests in southern China account for approximately 34 % of the forest area in China (Tang et al. 2018), and play an important role in global C balance (Yu et al. 2014). Soils in these forest ecosystems are acidic (often with pH < 4.5, and some even with pH < 4) and are highly weathered with a high content of reactive Fe phases (Zhang et al. 2010; Hu et al. 2015). Thus, Fe-bound C is likely to comprise an important portion of the total SOC stock in these forests (Coward et al. 2017). However, whether the mineral phases and organo-mineral associations in these highly acidic soils still respond significantly to further inputs of acid deposition has not been tested, which may have important implications for forest soil C sequestration in southern China and in other global regions with naturally acidic soils.

We carried out a long-term (nearly 10 yrs) field experiment with a gradient of acidity treatments (0, 9.6, 32, 96 mol H$^+$ ha$^{-1}$ yr$^{-1}$ as nitric acid + sulfuric acid) to examine acidification impacts on organo-mineral interactions and SOC pools (divided light and heavy fractions) in a tropical forest in southern China. Earlier work at this study site demonstrated that the acidity treatments significantly increased SOC stock and altered its biochemical composition (Wu et al. 2016). In this study we hypothesized that the acidity treatments enhanced SOC sequestration mainly arising from increased C in heavy (mineral-associated) fraction. We also expected that pH-sensitive organo-mineral interactions, especially changes in Fe phases, could explain the responses of SOC pools. Specifically, under the acidity treatments we quantified
1) the shift of mineral-associated C pools (Fe-bound C vs. Ca-bound C) and its absolute contribution to the increase in total SOC stock, and 2) the extent of cation losses and changes in Fe phases.

**Materials And Methods**

**Site description**

This study was conducted in Dinghushan National Nature Reserve (1133 ha), located in the middle part of Guangdong Province in southern China (112°30′ – 112°33′E, 23°09′ – 23°11′N). This area is characterized by a typical south subtropical monsoon climate. The mean annual temperature is 21°C, with the maximum and minimum average monthly temperature of 28.0°C in July and 12.6°C in January, respectively. Annual average relative humidity is 82%, and the mean annual rainfall is 1927 mm. Acid rain is a threat in this area with high deposition rates of 34.4 kg N ha⁻¹ yr⁻¹ and 32.6 kg S ha⁻¹ yr⁻¹ in recent decades (Liu et al. 2007), which has significantly lowered the soil pH of some forests to < 4 (Jiang et al. 2018).

In the reserve, there are three types of forests of different successional stages: the broadleaf forest, mixed pine and broadleaf forest, and pine forest. In this study, the experimental site was set up in the broadleaf forest, the most mature forest with age > 400 year. The forest is located 250–300 m above sea level and occupies approximately 600 ha. The dominant species were *Castanopsis chinensis*, *Cryptocarya concinna*, *Cryptocarya chinensis*, *Machilus chinensis*, and *Schima superba* (Yan et al. 2006). The bedrock is sandstone and shale belonging to the Devonian Period. Soil type is lateritic red earth (Yan et al. 2015) and is classified as an Oxisol according to the Keys to Soil Taxonomy (Soil Survey Staff 2014).

**Experimental treatments and design**

The acidity experiment was initiated in June 2009. Twelve plots were established and divided into four acidity treatments with three replicates each. Each treatment plot was measured 10 × 10 m² and was surrounded by a 3 m wide buffer strip. All plots and treatments were randomly arranged. The acidity treatments were irrigated with water of different pH values: CK (control, pH 4.5), T1 (pH 4.0), T2 (pH 3.5), and T3 (pH 3.0). To reflect the real mole ratio of S:N in the region, acidic solutions were prepared by adding a mixture of H₂SO₄ and HNO₃ in a 1:1 mole ratio to the local lake water based on the local acid rain records (Du et al. 2015). The simulated acid rain was applied to each plot below the canopy using a gasoline engine powered sprayer and sprayed twice a month. The amount applied to each plot was 40 L per application, equal to 4 mm rainfall. The H⁺ added in each plot was estimated as 0, 9.6, 32, 96 mol H⁺ ha⁻¹ yr⁻¹ in the T1, T2, T3 and T4 treatments, respectively, which was equal to about 0, 0.6, 2.0 and 6.0 times of that in the through-fall of the forest. More details on the experimental design and methods can be found in Wu et al (2016).

**Sample collection and soil properties measurements**
In September 2018, four cores (diameter = 5 cm) were randomly collected from the topsoil (0–10 cm) and subsoil (10–20 cm) in each plot, and combined to yield one composite sample per depth and plot. The soil sample was passed through a 2 mm sieve to remove rocks and plant roots, and then divided into two parts (field-moist soil sample and air-dried soil sample) for subsequent analysis. Fine roots (diameter ≤ 2mm) were picked and thoroughly rinsed in deionized water and dried at 60°C to constant biomass.

Soil moisture was determined by drying the field-moist soil samples at 105°C for 24 h. Soil pH values were measured using a glass electrode after shaking the samples for approximately 30 min in deionized water. The soil to water ratio was 1:2.5. Microbial biomass carbon (MBC) was measured using the chloroform fumigation extraction technique (Jenkinson 1987). The soil extractable dissolved organic carbon (DOC) was measured on the same samples used for the analysis of MBC and calculated as the K$_2$SO$_4$ – extractable C concentration. Total SOC was measured with the air-dried soil samples by using the K$_2$Cr$_2$O$_7$ + H$_2$SO$_4$ oxidation method (Schollenberger 1927). Total soil N was determined by semimicro + Kjeldahl (Menefee and Overman 1940; Bremner 1960). Total Ca, Al and Fe in soil were extracted with HNO$_3$-HF-HClO$_4$ and measured using atomic absorption spectrophotometry with a graphite furnace. Soil exchangeable cations (Ca$^{2+}$, Al$^{3+}$, and Fe$^{3+}$) were extracted with 0.1 M BaCl$_2$ (soil:solution ratio of 1:50) and measured on an Agilent 5100 ICP-OES (Galí Navarro et al. 2011).

**Density fractionations**

Each field-moist soil sample (30 g dry mass equivalent) was separated into two operationally defined soil fractions, a light fraction (LF) and a heavy fraction (HF), following a modified density fractionation technique (Ye et al. 2018). Given that the soils used in this study were rich in clay, the LF was separated by flotation after immersing soils in NaI solution at a density of 1.85 g cm$^{-3}$ combined with a ultrasonic treatment (a total energy input of 200 J mL$^{-1}$) in order to disrupt soil aggregates. The residual soil consisted of the remaining mineral-associated organic matter (HF). The separated soil fractions were then dried in an oven at 60°C and ground to a homogenized fine powder for SOC analysis (interpreted as LF-C and HF-C).

**Determination of Ca-bound C and Fe-bound C**

The HF soil samples were sequentially extracted with 0.5 M Na$_2$SO$_4$ and the C released in the solutions was interpreted as organic C associated with Ca bridges (Xu and Yuan 1993). The difference between HF-C and the C concentrations of the residual soils after Na$_2$SO$_4$ extraction was calculated as Ca-bound C. Next, Fe-bound organic C was measured following a procedure adapted from Lalonde et al (2012). The residual soil after Na$_2$SO$_4$ extraction was added to a solution containing sodium bicarbonate and trisodium citrate in a 50 mL polycarbonate centrifuge tube and heated to 80°C in a water bath, and then sodium dithionite was added to the tube and maintained at 80°C for 15 min. After centrifugation at 3000 g for 10 min, the supernatant was separated from the solid fraction. The procedure was repeated three times and the residual soil was rinsed three times with deionized water and then oven dried at 80°C for
organic C analysis, respectively. The concentrations of organic C in soil after dithionite-citrate-bicarbonate (DCB) extraction were subtracted from the Na$_2$SO$_4$-extracted samples to obtain Fe-associated organic C.

**Determination of Fe phases**

During the determination of Fe-bound C, the extracted solution was also analyzed by ICP-OES to measure the total free Fe (oxyhydr)oxides (Fe$_d$). Dry HF subsamples were separately extracted with acid ammonium oxalate or sodium pyrophosphate to measure poorly crystalline (i.e., short-range-ordered, SRO) Fe (Fe$_o$) and organically complexed and colloidal Fe (Fe$_p$), respectively. Briefly, soils were extracted in the dark using 0.175 M ammonium oxalate and 0.1 M oxalic acid at pH = 3 with a soil:solution ratio of 1:60 and shaken for 2 h. Soils were extracted by 0.1 M sodium pyrophosphate (soil:solution ratio of 1:20) at pH = 10 and the supernatant was syringe-filtered through a 0.2 µm nylon membrane filter after centrifugation at 2000 g for 15 min. The Fe concentrations were determined by ICP-OES.

Additionally, the field-moist soil samples were extracted in 0.5 M hydrochloric acid (HCl) to measure Fe(II) and Fe(III) in HCl, denoted Fe(II)$_{\text{HCl}}$ and Fe(III)$_{\text{HCl}}$ (Huang and Hall 2017). Briefly, field-moist soil samples (3 g dry mass equivalent) were immersed in a 1:10 ratio with HCl in the field, and then vortexed, shaken for 1 hr, and filtered to 0.22 µm. Concentrations of Fe(II)$_{\text{HCl}}$ were measured using a colorimetric ferrozine assay and corrected for Fe(III)$_{\text{HCl}}$ interference (Viollier et al. 2000). Field-moist soil samples were also extracted in 0.2 M sodium citrate/0.05 M ascorbic acid solutions to measure the easily reducible Fe oxides (Fe$_{\text{ca}}$) in short-range-order minerals and organic fractions using microplate-based ferrozine methods (Huang and Hall 2017). Briefly, field-moist soil samples (1.5 g dry mass equivalent) were added to freshly prepared citrate/ascorbic solution in a 1:30 ratio, vortexed, shaken in the dark for 16 h, and centrifuged for 10 min at 10,000 g. 40 µL of extract was added to 40 µL of 10% hydroxylamine hydrochloride and 200 µL of color reagent (1 g L$^{-1}$ Ferrozine in 600 mM HEPES buffer at pH 8), the absorbance at 562 nm was then recorded at 1h. The supernatant solution was decanted to a clean HDPE bottle for dark storage at 4°C for analysis within two weeks of collection. The difference between Fe$_d$ and Fe$_{\text{ca}}$ (Fe$_d$-$\text{ca}$) was interpreted as crystalline Fe (oxyhydr)oxides.

**Statistical analyses**

Data analyses were conducted using SPSS 20.0 for Windows (IBM Corporation, Armonk, New York, USA) and R version 4.0.2. Analysis of Variance (ANOVA) was used to determine the statistical significance ($\alpha = 0.05$) of the acidity treatment, soil layer and their interactive effects on bulk SOC and its fractions/pools as well as mineral elements (Ca, Al and Fe including Fe oxides). Tukey's multiple comparison test (HSD) was conducted if significant effects of acid addition or soil layer were found. Pairwise relationships between soil pH and biogeochemical variables, as well as biogeochemical variables and SOC fractions/pools were assessed using Pearson correlation coefficients.

A structural equation modelling (SEM) approach was also used to test a conceptual model for the acid deposition impacts on LF-C and HF-C. The SEM analysis was performed with the IBM SPSS Amos 22.0.
using the maximum likelihood estimation method. Several tests were used to assess model fit: the Chi-square ($\chi^2$)-test, comparative fit index (CFI) and root square mean error of approximation (RMSE).

**Results**

**Soil general characteristics**

Soil pH values decreased significantly under the acidity treatments compared to the control ($p < 0.05$, Table 1), and pH was significantly lower in topsoil than in subsoil ($p < 0.05$, Table 1). There was no significant difference in soil moisture or bulk density among treatments ($p > 0.05$ for both, Table 1). The MBC in both topsoil and subsoil decreased significantly with the acidity treatments compared to the control ($p < 0.05$, Table 1), except for the subsoil in the T1 treatment ($p > 0.05$, Table 1). DOC in topsoil was significantly higher under the T1 and T3 treatment compared to control ($p < 0.05$, Table 1), while in subsoil, DOC was significantly lower under the T3 than that under the T1 and T2 treatments ($p < 0.05$, Table 1). The fine root biomass significantly decreased under the acidity treatments compared to the control ($p < 0.05$, Table 1), except for the T2 treatment ($p > 0.05$, Table 1).
Table 1
Changes of soil properties, microbial biomass C, dissolved organic C and fine root biomass under the acidity treatments. Values are means (SE).

| Property                        | Depth (cm) | CK         | T1          | T2          | T3          |
|---------------------------------|------------|------------|-------------|-------------|-------------|
| Soil pH                         | 0–10       | 3.95 (0.04)a | 3.76 (0.04)b | 3.78 (0.02)b | 3.76 (0.04)b |
|                                 | 10–20      | 4.04 (0.02)a | 3.94 (0.04)b | 3.86 (0.02)bc| 3.82 (0.03)c |
| Soil moisture (% weight)         | 0–10       | 0.34 (0.01) | 0.36 (0.01) | 0.35 (0.01) | 0.33 (0.01) |
|                                 | 10–20      | 0.35 (0.01) | 0.32 (0.01) | 0.32 (0.02) | 0.34 (0.01) |
| Soil bulk density (g cm⁻³)      | 0–10       | 1.24 (0.08) | 1.25 (0.06) | 1.28 (0.11) | 1.23 (0.09) |
|                                 | 10–20      | 1.16 (0.06) | 1.13 (0.08) | 1.16 (0.05) | 1.15 (0.09) |
| MBC (g kg⁻¹)                    | 0–10       | 0.55 (0.02)a | 0.48 (0.01)b | 0.37 (0.02)c | 0.36 (0.02)c |
|                                 | 10–20      | 0.42 (0.02)a | 0.40 (0.03)ab| 0.35 (0.03)bc| 0.32 (0.02)c |
| DOC (g kg⁻¹)                    | 0–10       | 0.30 (0.01)b | 0.36 (0.00)a | 0.32 (0.02)b | 0.38 (0.01)a |
|                                 | 10–20      | 0.25 (0.01)ab| 0.26 (0.03)a | 0.23 (0.01)b | 0.23 (0.02)b |
| Fine root biomass (kg m⁻²)      | 0–20       | 0.28 (0.01)a | 0.24 (0.01)b | 0.25 (0.02)ab| 0.17 (0.01)c |

The acidity treatments are: CK = control (0 mol H⁺ ha⁻¹ yr⁻¹), T1 = 9.6 mol H⁺ ha⁻¹ yr⁻¹, T2 = 32 mol H⁺ ha⁻¹ yr⁻¹, and T3 = 96 mol H⁺ ha⁻¹ yr⁻¹. Different letters in each soil layer denote significant difference (p < 0.05) among the treatments. MBC, microbial biomass carbon; DOC, dissolved organic carbon.

Soil geochemical characteristics

Total Ca and Fe contents in both topsoil and subsoil and total Al content only in subsoil decreased significantly under the acidity treatments compared to the control (p < 0.05 for all, Table 2). The exchangeable Ca²⁺ in topsoil was significantly lower in T1 than in T3 treatment (p < 0.05, Table 2), but neither significantly differed from the control (p > 0.05, Table 2). In subsoil, exchangeable Ca²⁺ in the control was significantly higher than in the T1 and T2 treatments (p < 0.05, Table 2), but not in the T3 treatment (p > 0.05, Table 2). Exchangeable Al³⁺ and Fe³⁺ in both topsoil and subsoil increased significantly with the acidity treatments compared to the control (p < 0.05 for both, Table 2).
Table 2
Changes of the total and extractable Ca, Al, Fe under the acidity treatments. Values are means (SE).

| Property         | Depth (cm) | CK      | T1     | T2     | T3     |
|------------------|------------|---------|--------|--------|--------|
| Total Ca (g kg⁻¹) | 0–10       | 0.58 (0.10)a | 0.36 (0.06)b | 0.25 (0.04)b | 0.22 (0.04)b |
|                  | 10–20      | 0.56 (0.08)a | 0.23 (0.06)b | 0.20 (0.02)b | 0.09 (0.02)b |
| Total Al (g kg⁻¹) | 0–10       | 77.54 (1.64)ab | 79.86 (1.45)a | 73.44 (1.97)b | 78.84 (2.87)ab |
|                  | 10–20      | 89.60 (2.28)a | 83.13 (2.26)b | 77.64 (1.22)b | 77.73 (2.24)b |
| Total Fe (g kg⁻¹) | 0–10       | 42.23 (0.49)a | 40.74 (0.54)b | 37.94 (0.33)c | 39.90 (0.39)bc |
|                  | 10–20      | 43.38 (0.70)a | 41.03 (0.44)b | 40.68 (0.36)b | 39.32 (0.66)b |
| Exchangeable Ca²⁺ (g kg⁻¹) | 0–10 | 0.11 (0.01)ab | 0.08 (0.01)b | 0.10 (0.01)ab | 0.12 (0.01)a |
|                  | 10–20      | 0.07 (0.01)a | 0.02 (0.00)c | 0.06 (0.00)b | 0.09 (0.01)a |
| Exchangeable Al³⁺ (g kg⁻¹) | 0–10 | 0.67 (0.02)b | 0.93 (0.05)a | 0.88 (0.02)a | 0.88 (0.03)a |
|                  | 10–20      | 0.60 (0.02)c | 0.71 (0.05)b | 0.73 (0.03)ab | 0.82 (0.03)a |
| Exchangeable Fe³⁺ (mg kg⁻¹) | 0–10 | 16.91(0.66)c | 36.87(1.00)b | 34.02(3.80)b | 42.50(2.73)a |
|                  | 10–20      | 14.81(1.22)c | 21.53(1.41)b | 27.39(2.61)a | 20.60(2.35)b |

The acidity treatments are: CK = control (0 mol H⁺ ha⁻¹ yr⁻¹), T1 = 9.6 mol H⁺ ha⁻¹ yr⁻¹, T2 = 32 mol H⁺ ha⁻¹ yr⁻¹, and T3 = 96 mol H⁺ ha⁻¹ yr⁻¹. Different letters in each soil layer denote significant differences (p < 0.05) among the treatments.

Soil iron phases

The concentrations of Fe(II)HCl, Fe(III)HCl, and Fe_ca in both topsoil and subsoil increased significantly under the acidity treatments compared to the control (p < 0.05 for all, Table 3). Soil Fe_d concentrations did not vary consistently among treatments and between soil layers (Table 3). However, the concentrations of Fe_d-ca in both topsoil and subsoil decreased significantly under the acidity treatments compared to the control (p < 0.05 for both, Table 3). Soil Fe_p concentrations in both topsoil and subsoil also decreased significantly with the acidity treatments (p < 0.05, Table 3), while Fe_o concentrations were not significantly influenced by the acidity treatments (p > 0.05, Table 3).
Table 3
Changes of Fe phases under the acidity treatments. Values are means (SE).

| Fe pool              | Depth (cm) | CK           | T1           | T2           | T3           |
|----------------------|------------|--------------|--------------|--------------|--------------|
| Fe(II)$_{\text{HCl}}$ (g kg$^{-1}$) | 0–10       | 0.19 (0.01)c | 0.29 (0.01)a | 0.22 (0.02)b | 0.24 (0.01)ab |
|                      | 10–20      | 0.18 (0.01)b | 0.20 (0.01)ab | 0.19 (0.01)ab | 0.21 (0.01)a |
| Fe(III)$_{\text{HCl}}$ (g kg$^{-1}$) | 0–10       | 1.52 (0.02)c | 2.32 (0.08)a | 2.05 (0.18)b | 2.05 (0.12)b |
|                      | 10–20      | 1.67 (0.05)b | 1.86 (0.06)a | 1.69 (0.09)ab | 1.90 (0.04)a |
| Fe$_{\text{ca}}$ (g kg$^{-1}$)   | 0–10       | 3.93 (0.17)c | 4.62 (0.13)b | 4.46 (0.04)b | 4.91 (0.07)a |
|                      | 10–20      | 3.49 (0.14)c | 4.16 (0.05)b | 4.99 (0.10)a | 4.21 (0.04)b |
| Fe$_{\text{d}}$ (g kg$^{-1}$)    | 0–10       | 13.38 (0.18)b | 14.05 (0.27)a | 13.61 (0.53)ab | 14.16 (0.26)a |
|                      | 10–20      | 14.54 (0.29)a | 13.86 (0.52)ab | 14.78 (0.24)a | 13.69 (0.30)b |
| Fe$_{\text{d-ca}}$ (g kg$^{-1}$)  | 0–10       | 9.46 (0.12)a | 9.41 (0.06)ab | 9.14 (0.53)ab | 9.24 (0.03)b |
|                      | 10–20      | 11.05 (0.05)a | 9.39 (0.12)c | 9.79 (0.11)b | 9.48 (0.34)c |
| Fe$_{\text{p}}$ (g kg$^{-1}$)     | 0–10       | 9.84 (0.11)a | 8.89 (0.20)b | 8.49 (0.20)b | 8.15 (0.35)b |
|                      | 10–20      | 10.24 (0.22)a | 8.25 (0.32)b | 9.01 (0.20)b | 8.49 (0.31)b |
| Fe$_{\text{o}}$ (g kg$^{-1}$)     | 0–10       | 3.45 (0.11) | 3.63 (0.13) | 3.58 (0.10) | 3.53 (0.24) |
|                      | 10–20      | 3.94 (0.08) | 3.99 (0.04) | 4.00 (0.15) | 3.88 (0.09) |

The acidity treatments are: CK = control (0 mol H$^+$ ha$^{-1}$ yr$^{-1}$), T1 = 9.6 mol H$^+$ ha$^{-1}$ yr$^{-1}$, T2 = 32 mol H$^+$ ha$^{-1}$ yr$^{-1}$, and T3 = 96 mol H$^+$ ha$^{-1}$ yr$^{-1}$. Different letters in each soil layer denote significant differences ($p < 0.05$) among the treatments. Fe(II)$_{\text{HCl}}$, reduced Fe in 0.5 M HCl; Fe(III)$_{\text{HCl}}$, oxidized Fe in 0.5 M HCl; Fe$_{\text{ca}}$, citrate-ascorbate extractable Fe; Fe$_{\text{d}}$, Dithionite–bicarbonate–citrate extractable Fe; Fe$_{\text{p}}$, sodium pyrophosphate extractable Fe; Fe$_{\text{o}}$, ammonium oxalate extractable Fe.

Soil organic carbon fractions

The total SOC content increased significantly under the acidity treatments compared to the control, except for the T1 treatment in both topsoil and subsoil ($p < 0.05$, Fig. 1a). Both LF-C and HF-C concentration and stock also increased significantly under the acidity treatments compared to the control, except for the HF-C in the T1 treatment in subsoil ($p < 0.05$, Fig. 1b, c). Since the bulk density did not change among treatments ($p > 0.05$, Table 1), the stock of SOC and its fractions varied similarly with their content (Fig. 1). Overall, the acidity treatments increased LF-C to 3.1–4.1 Mg C ha$^{-1}$ vs. 1.6 Mg C ha$^{-1}$ in the control and increased HF-C to 32.5–36.4 Mg C ha$^{-1}$ vs. 28.8 Mg C ha$^{-1}$ in the control. The increase in LF-C and HF-C accounted for 22–32% and 78–83%, respectively, in the increase of the total SOC stock.
The content of Ca-bound C in the topsoil showed no significant change among treatments \((p > 0.05, \text{Fig.~} 1d)\), while in the subsoil it decreased significantly under acidity treatments compared to the control \((p < 0.05, \text{Fig.~} 1d)\). Fe-bound C in both topsoil and subsoil increased significantly with the acidity treatments compared to the control \((p < 0.05, \text{Fig.~} 1e)\).

Relationships among the SOC fractions and geochemical characteristics

Across all treatments, soil pH was negatively correlated with SOC, LF-C, HF-C and Fe-bound C, but not with Ca-bound C (Fig. 2a). SOC was positively correlated with both LF-C and HF-C (Fig. 2b,c). HF-C was positively correlated with Fe-bound C only, but not with Ca-bound C (Fig. 2d,e).

Soil pH was also negatively correlated with exchangeable Al\(^{3+}\), exchangeable Fe\(^{3+}\), Fe(II)\(_{\text{HCl}}\), Fe(III)\(_{\text{HCl}}\), and Fe\(_{\text{ca}}\), while total Ca, total Fe, total Al, and Fe\(_p\) showed opposite relationships (Fig. 3a). The Ca-bound C was positively correlated with the loss of total Ca only, but not with exchangeable Ca\(^{2+}\) (Fig. 3b). However, the Fe-bound C was positively correlated with exchangeable Fe\(^{3+}\) and Al\(^{3+}\) (Fig. 3c,d). The Fe-bound C was also positively correlated with the concentrations of Fe(II)\(_{\text{HCl}}\), Fe(III)\(_{\text{HCl}}\), and Fe\(_{\text{ca}}\) (Fig. 3e,f,g).

Controls on SOC and fraction changes under the acidity treatments

The SEM model implied by our data showed that the decreased soil pH under the acidity treatments was the optimal predictor and directly explained 53%, 63% and 83% of the variance in fine root biomass, Fe-bound C, and LF-C, respectively (Fig. 4). There was no significant relationship of soil pH with MBC or Ca-bound C. Fine root biomass had a direct negative relationship with MBC \((R^2 = 0.46)\), whereas neither fine root biomass nor MBC was significantly related to LF-C. Change in LF-C directly explained 50% of the variance in DOC, which in turn together with Fe-bound C directly explained 81% of the variance in HF-C. There was no significant relationship of HF-C with MBC and Ca-bound C, respectively.

Discussion

We found that the acidity treatments significantly increased total SOC content and stock including both light and heavy fractions (Fig. 1a, b, c). The increased SOC was unlikely to have come from greater plant C input, as soil acidification was previously shown to limit plant productivity and led to decreased fine root biomass as reported in this study (Liang et al. 2013; Table 1). The acidity treatments also decreased MBC (Table 1) and could inhibit soil respiration (Wu et al. 2016), which might also have contributed to the observed SOC accumulation. However, decreased MBC might be expected to cause higher LF-C but lower HF-C, because microbial residues derived from catabolism of LF-C can potentially be important constituents contributing to mineral-associated C (Cotrufo et al. 2013; Liang et al. 2017). In contrast to this reasoning, we found that the HF-C increased much more in absolute amount than the LF-C under the acidity treatments. Consistent with our hypothesis, the acidity treatments increased the HF-C stock, which accounted for most of the total increase in SOC stock (Table 4).
With a SEM model implied by our data, we further found that pH-sensitive organo-mineral interactions particularly changes in Fe phases, to a large extent, explained the increase of HF-C under the acidity treatments. The increase in Fe-bound C under the acidity treatments was much larger than the decrease in Ca-bound C (Fig. 1d, e). Consistent with pH buffering by dissolution of Al and Fe phases in acidic soils, we found that the acidity treatment generally decreased exchangeable Ca$^{2+}$, while soil extractable Al$^{3+}$ and Fe$^{3+}$ increased significantly (Table 2). In an alpine grassland of Europe with soil pH < 3.5, acid deposition similarly led to buffering by Al$^{3+}$ and Fe$^{3+}$ (Bowman et al. 2008). Increases of extractable Al$^{3+}$ and Fe$^{3+}$ under the acidity treatments reflect the enhanced solubility of Al and Fe mineral phases (Gu et al. 1994; Mueller et al. 2012), causing losses of these metals via leaching as indicated by decreases in total Al and Fe in the subsoil (Table 2). However, increased Al and Fe solubility likely also promoted formation of new organic−mineral complexes between DOC and Fe/Al phases, perhaps through ligand exchange between carboxyl groups of organic matter and singly coordinated inorganic hydroxyl groups at metal/mineral surfaces (Masiello et al. 2004; Chen et al. 2014). Negative relationships between DOC leaching and extractable Al$^{3+}$ or Fe$^{3+}$ with increasing soil acidity have frequently been observed in acid soils (Kalbitz et al. 2000; Filep and Rékási 2011; Lu et al. 2013; Xiao et al. 2013). Previous work at our study site also found that the acidity treatments significantly decreased the DOC leaching flux (Qiu et al. 2013, 2015), and here we found positive relationships between soil exchangeable Al$^{3+}$ and Fe$^{3+}$ with mineral-associated C, consistent with DOC sorption or co-precipitation (Fig. 3). In addition, soil acidification could decrease the solubility of organic matter due to the protonation of organic acid functional groups at pH values approaching zero net charge (Chorover and Sposito 1995), as well as by increasing molecular aggregation (Louzao et al. 1990).

Changes in Fe phase compositions under the acidity treatments were also likely related to the large increase in Fe-bound C pool in this study (Fig. 4b). We found that the acidity treatments significantly decreased Fe$_{d-ca}$ while Fe(II)$_{HCl}$, Fe(III)$_{HCl}$, and Fe$_{ca}$ increased (Table 3). These contrasting changes in Fe pools likely reflect the progressive transformation of crystalline Fe oxides into increasingly short-range-ordered Fe phases. Decreased Fe$_{d-ca}$ indicates that soil acidification increased the solubility of crystalline Fe oxides, as observed in other acid forest soils (Guo et al. 2007). Intriguingly, we also found that Fe$_p$ decreased under the acidity treatments, possibly indicating competition between H$^+$ and Fe$^{3+}$ for binding sites in soil organic matter under these highly acidic conditions, leading to release of Fe$^{3+}$ from monomeric organo-Fe complexes and its subsequent precipitation in Fe phases that were not extracted by Fe$_p$. The formation of new short-range-ordered Fe phases was consistent with the observed increase in the Fe$_{ca}$ and Fe(III)$_{HCl}$ pools, which include highly reactive portions of soil Fe (Hall and Silver 2015). The fact that Fe(II)$_{HCl}$ was a small portion of the 0.5 M HCl-extractable Fe pool indicated that acid-mediated dissolution, rather than reductive dissolution, was likely the dominant process controlling the observed changes in soil Fe (Table 3). Finally, the new Fe phases in the acidity treatments likely associated with DOC inputs via adsorption and co-precipitation (Wagai and Mayer 2007; Chen et al. 2014; Mikutta et al. 2014; Coward et al. 2017), as evidenced by the positive relationships of soil Fe(III) and Fe$_{ca}$ with Fe-bound C under the acidity treatments (Fig. 3d,e,f).
Our results have also important implications for understanding the impacts on SOC sequestration of ongoing acidification from atmospheric N deposition in tropical forests. At the global scale, soil pH decreases linearly with N addition (Tian and Niu 2015), and at our study site, a parallel experiment with N addition alone also revealed significant soil acidification (Lu et al. 2014). N-induced increases in SOC have been detected from temperate to tropical forests, but the underlying mechanisms in tropical forests remain poorly understood (Lu et al. 2021). Because many tropical forests are more strongly limited by phosphorus (P) than N, increased plant productivity and C input of plant residue cannot likely explain increases in SOC (Lu et al. 2021). N-induced soil acidification is widely known to suppress certain aspects of microbial decomposition activity, such as production of oxidative enzymes (Chen et al. 2018), and these changes have previously been invoked to explain the increased SOC stock and the change of biochemical composition under N addition (Cusack et al. 2011). Rather, a recent study by Lu et al (2021) showed that in tropical forests N addition increased SOC sequestration, mainly arising from increased C in heavy (mineral-associated) fraction accompanied by acidification-induced the decreases in both soil CO₂ efflux and DOC leaching. Here our data suggest that pH-sensitive organo-mineral interactions, especially formation of new Fe-C associations that provide protection from SOC decomposition and DOC leaching, play a key role in regulating the response of SOC pools in tropical forests and explain the observed increases in mineral-associated C.

Conclusions

Using a long-term acidity manipulation field experiment, we found that increased inputs of acid deposition in a highly weathered and naturally acidic tropical soil could enhance SOC sequestration, which mainly arising from increased C in mineral-associated fraction. Accordingly, soil organo-mineral interactions responded significantly to further inputs of acid deposition, with increased the leaching of base cations (e.g. Ca²⁺) and the solubility of Al and Fe phases. These acidity treatments converted more crystalline Fe (oxyhydr)oxides to short-range-ordered phases, resulting in a large increase in Fe-bound C vs. a relatively small decrease in Ca-bound C. With a SEM model implied by our data, our results further highlight the importance of pH-sensitive geochemical changes and the key roles of Fe in regulating the response of SOC to acid deposition in highly weathered soils. Therefore, accounting for pH-sensitive geochemistry is critical to predict ecosystem C budgets under future acid deposition scenarios, and that these principles could be usefully incorporated in mechanistic biogeochemical models.

Declarations

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**Author contributions**

QD and DZ planned the research; JC, YH, JL and LS conducted the field research; JC also conducted the laboratory research, analyzed results and wrote the original draft; SJH, QD and DH provided constructive comments and suggestions in the manuscript writing; and all authors edited the manuscript.

**Conflict of interest**

The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

**Data availability**

Data and materials are available upon request from the corresponding author.

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Figures
Figure 1

Responses of soil organic carbon fractions to the acidity treatments. The acidity treatments are: CK = control (0 mol H+ ha−1 yr−1), T1 = 9.6 mol H+ ha−1 yr−1, T2 = 32 mol H+ ha−1 yr−1, and T3 = 96 mol H+ ha−1 yr−1. SOC, soil organic carbon; Fe-bound C, iron-bound organic carbon; Ca-bound C, calcium-bound organic carbon; LF-C and HF-C indicate the organic carbon in light fraction and heavy fraction, respectively. Error bars represent ± standard errors. The columns and scatters represent soil organic carbon content and stock, respectively. Different lowercase letters within the columns in each soil layer
denote significant difference (p<0.05) in the contents of soil organic carbon fractions among the treatments. Different capital letters above the scatters denote significant difference (p<0.05) in the stocks of soil organic carbon fractions among the treatments. As none of the interactions between treatment and depth on soil organic carbon fractions were significant, only main effects of treatment and soil depth are presented. *, ** and *** indicates that factors are significant at p<0.05, p<0.01 and p<0.001, respectively.

Figure 2

Pearson’s correlation coefficients between soil pH and soil organic carbon fractions. SOC, soil organic carbon; Fe-bound C, iron-bound organic carbon; Ca-bound C, calcium-bound organic carbon; LF-C and HF-C indicate the organic carbon in light fraction and heavy fraction, respectively. The red and blue squares indicate positive or negative correlations between parameters at a significance level of $\alpha=0.05$ respectively.
Figure 3

Pearson’s correlation coefficients between soil pH, Ca-bound C, Fe-bound C and geochemical characteristics. Fe-bound C, iron-bound organic carbon; Ca-bound C, calcium-bound organic carbon; Feca, short-range-order Fe phases; Fed, Dithionite–bicarbonate–citrate extractable Fe phases; Fep, sodium pyrophosphate extractable Fe phases; Feo, ammonium oxalate extractable Fe phases. The red and blue squares indicate positive or negative correlations between parameters at a significance level of $\alpha = 0.05$ respectively.
Figure 4

Structural equation model (SEM) analysis for the effects of the acidity treatment on soil organic carbon fractions via the pathways of soil acidification. Results of the model fitting: Chi-square (X2) = 27.381, p = 0.055, degree of freedom (df) = 18, comparative fit index (CFI) = 0.942, root square mean error of approximation (RMSEA) = 0.159. Numbers at arrows are standardized path coefficients. Arrow thickness represents the strength of the relationships. Grey arrows indicate insignificant relationships (p>0.05), while black and red arrows indicate significant positive and negative relationships, respectively (p<0.05). In addition, R2 values associated with response variables indicate the proportion of variation explained by relationships with other variables. MBC, microbial biomass carbon; Fe-bound C, iron-bound organic carbon; Ca-bound C, calcium-bound organic carbon; LF-C and HF-C indicate the organic carbon in light fraction and heavy fraction, respectively.