Vegetation change impacts on soil organic carbon chemical composition in subtropical forests

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Changes in the chemical composition of soil organic carbon (SOC) might strongly affect the global carbon cycle as it controls the SOC decomposition rate. Vegetation change associated with long-term land use changes is known to strongly impact the chemical composition of SOC; however, data on the impacts of vegetation change following disturbance events of short durations and succession that occur frequently in forest ecosystems via diverse management objectives on SOC chemical composition are negligible. Here we examined the impacts of vegetation changes on the chemical composition of SOC by sampling soils of native broad-leaved forests, planted mixed broad-leaved and coniferous forests, and tea gardens in eastern China. We used nuclear magnetic resonance spectroscopy to quantify SOC chemical composition. We found that among all components of SOC chemical composition, alkyl carbon (C) and aryl C were more liable to change with vegetation than other SOC components. Soil pH was negatively correlated to the relative abundances of alkyl C and N-alkyl C, and Shannon’s index of overstory plant species was positively correlated to the relative abundances of phenolic C and aromaticity. Our results suggest that vegetation changes following short disturbance events and succession may strongly alter SOC chemical composition in forest ecosystems.

Globally, soil organic carbon (SOC) amounts to approximately 2,344 Gt, making it the largest terrestrial carbon (C) pool1–3. Even small changes in SOC pool decomposition might make significant changes in the atmospheric carbon (C) concentration4, and could lead to a strong positive feedback to climate change5. SOC decomposition rates are strongly associated with its chemical composition6, with decomposition rates decreasing from O-alkyl C to alkyl C7,8. Thus, shifts in SOC chemical composition affect SOC stabilization9,10.

The input of C to the soil profile is determined by C allocation, decomposition, and production11. Vegetation change associated with long-term land use changes may result in SOC chemical compositional shifts12–14. For example, when natural forests were converted to hoop pine plantations, O-alkyl C decreased and alkyl C increased13. After native shrub lands were converted to chestnut plantations, both alkyl C and O-alkyl C decreased14. Vegetation change and the corresponding management may alter soil aggregates, which influence SOC chemical composition6. It also leads to lignin variation, which consequently impacts the yield of lignin-derived phenols and carboxyl15. Vegetation change affects lignin degradation through alterations in soil texture16, and increased yields of phenolic CuO oxidation products15. The soil texture consequently affects carbohydrates16, and O-alkyl C (mainly carbohydrate) was found to predominate in the larger fractions, whereas alkyl carbon (primarily polymethylene) dominated the clay fractions17. The influence of vegetation change and the corresponding management toward the initiation of changes in lignin and carbohydrates might be attributed to particular microbes16,18. As a result of the different growth characteristics and nutrient demands of specific microbial groups, the changes in lignin and carbohydrates may favor the growth of certain microbial groups over others19. Variations in SOC chemical composition associated with vegetation change might be attributed to the different patterns of mineralization of plant-derived microbial carbohydrate inputs15,21. However, it remains unclear whether vegetation change following disturbance events of short durations, which frequently occur naturally or via diverse management objectives in forest ecosystems, affects the chemical composition of SOC.

We sampled forest soils from native broadleaf forests, mixed evergreen broadleaved and coniferous forests, and tea gardens in Fengyang Mountain nature reserve, Zhejiang Province, China, to examine the impact of

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vegetation change following disturbance events of short durations and succession on the chemical composition of SOC. The mixed broadleaved and coniferous forests, and tea gardens were converted from native broadleaf forests during 1971–1973, and since 1975, the study forests have been protected from human disturbances. We hypothesized more similar SOC chemical compositions between native broadleaf forests and broadleaf tea gardens than those between native broadleaf forests and mixed broadleaved and coniferous forests, as plants play an important role in SOC dynamics, mainly by means of litter quality, which changes in the quantity and type of fresh organic matter that enters the soil; thus, manipulating the SOC chemical composition accordingly. We alternatively hypothesized that the conversion from native broadleaf forests to tea gardens has greater impacts on the chemical composition of SOC than the conversion to mixed forests, as the conversion to tea garden involved complete clearance, which was a more intensive disturbance than the conversion from native forests to mixed forests via selective deforestation. We also hypothesized that overstory plant diversity, as well as soil physicochemical characteristics, may be attributable to the variation in SOC chemical composition. We addressed: (i) impacts of vegetation change from native broadleaved to mixed forests and tea garden on SOC chemical composition, and (ii) associations between tree species diversity, soil physicochemical characteristics and SOC chemical composition.

Results
Vegetation change impacts on tree diversity and soil physicochemical characteristics. In this study, when evergreen broadleaved forests were converted to mixed forests and tea gardens, the Shannon index decreased significantly (Table 1). Soil physicochemical characteristics varied among vegetation types, primarily in bulk density and total phosphorus concentration. Bulk density significantly decreased from broadleaved forests to tea gardens, while total phosphorus concentration strongly decreased from broadleaved forests to mixed forests. The impacts of vegetation change on pH, total organic carbon, and total nitrogen concentration were less apparent (Table 1).

Vegetation change impacts on SOC chemical composition. The integration of the major regions of $^{13}$C resonance revealed that alkyl C and O-alkyl C ($C_{0−45}$ ppm and $C_{45−110}$ ppm, respectively) were the dominant C components in all soils (Fig. 1). A multiple-response permutation procedures analysis (MRPP) showed the differences of SOC chemical composition among vegetation types with $P = 0.003$ (Table 2). The relative abundance of alkyl C and aryl C differed significantly among three vegetation types, with the highest relative abundance of alkyl C and lowest aryl C in mixed forests (Fig. 2, Table 3). The A/O-A also differed significantly among vegetation types, with the highest one in mixed forests, while the aromaticity in the mixed forest was lower than the other forest types (Fig. 3a,b, Table 3). However, there was no significant difference in SOC chemical composition.

### Table 1. Species diversity index and physicochemical soil characteristics of sampling stands. Values shown are means and 1 SE (in bracket). Different letters in the same row indicate a significant difference ($P < 0.05$) between vegetation types.

| Characteristic                          | Native forest | Mixed forest | Tea garden |
|-----------------------------------------|---------------|--------------|------------|
| Shannon’s index ($400 \text{ m}^2$)     | 1.61 (0.27)a  | 0.81 (0.28)b | 0.75 (0.17)c|
| Bulk density ($\text{g cm}^{-3}$)       | 1.35 (0.06)a  | 1.35 (0.06)a | 1.27 (0.05)b|
| pH                                      | 5.54 (0.26)a  | 5.24 (0.30)a | 5.18 (0.34)a|
| Total organic carbon ($\text{g kg}^{-1}$)| 34.90 (8.72)a | 30.76 (3.34)a | 28.19 (3.26)a|
| Total nitrogen concentration ($\text{g kg}^{-1}$) | 3.16 (0.84)a | 3.32 (0.76)a | 2.40 (0.70)a|
| Total phosphorus concentration ($\text{g kg}^{-1}$) | 0.49 (0.07)a | 0.40 (0.07)b | 0.55 (0.05)a|

![Figure 1. Solid-state $^{13}$C NMR spectra for soil organic carbon (SOC) in three vegetation types.](image-url)
between broadleaved forests and tea gardens. For the other components of SOC chemical composition, there were no significant differences among three vegetation types (Fig. 2, Table 3).

The NMDS ordination resulted in a final stress of 0.046 (Fig. 4). The distance between broadleaved forest and mixed forest was considerable, and depicted along the NMDS axis 1 and axis 2 with $P = 0.020$ and $P = 0.023$, respectively. However, the distance between tea garden and broadleaved forest was not distinct along either the NMDS axis 1 or axis 2 ($P = 0.640$ and $P = 0.541$, separately). The chemical composition of SOC assemblages was strongly associated with all individual components, particularly along the NMDS axis 1 (Table 4). However, the chemical composition of SOC assemblages had statistically weak association with forest and soil properties. The NMDS axis 1 corresponded mainly to a gradient of increasing Shannon index and pH ($P = 0.059$ and $P = 0.098$, respectively). The NMDS axis 2 corresponded primarily to a gradient of increasing total soil phosphorus (TP) and decreasing total soil nitrogen (TN) ($P = 0.085$ and $P = 0.104$, respectively) (Fig. 5, Table 5).

**Influence of soil physicochemical characteristics and plant species diversity on SOC chemical composition.** Soil pH was negatively correlated with alkyl C and N-alkyl C, with $P = 0.043$ and $P = 0.039$, respectively. The Shannon index was positively correlated with phenolic C, and the aromaticity ($P = 0.018$ and 0.050, respectively). Soil physicochemical characteristics and Shannon index did not significantly correlate to the other components of chemical composition ($P > 0.05$) (Table 6).

**Discussion**
Vegetation controls the magnitude of SOC stocks as well as the composition of SOC in soils$^{24}$, and is thus regarded as one of critical factors in SOC composition$^{25}$. We found that the differences in the chemical composition of SOC are primarily represented by alkyl C and aryl C, which is similar to previous findings$^{24,26-28}$. This indicated that alkyl C and aryl C were more sensitive to vegetation changes.

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**Table 2.** Results of multiple-response permutation procedures (MRPP) testing the null hypothesis of no significant differences in the organic carbon chemical composition of the soil among three vegetation types. Observed delta = 6.954, expected delta = 8.363, chance-corrected within-group agreement, $A = 0.169$, $P = 0.003$.

| Vegetation   | Delta | n |
|--------------|-------|---|
| Native forests | 7.858 | 9 |
| Mixed forests  | 4.772 | 6 |
| Tea gardens    | 8.193 | 4 |

**Table 3.** Analysis of variance results for the soil organic carbon chemical composition among three vegetation types.

| Composition        | df | $F$   | $P$   |
|--------------------|----|-------|-------|
| Alkyl C            | 2, 16 | 16.460 | <0.001|
| N-alkyl C          | 2, 16 | 1.496  | 0.254 |
| Carbohydrate C     | 2, 16 | 0.500  | 0.616 |
| Di-O-alkyl C       | 2, 16 | 0.192  | 0.827 |
| Aryl C             | 2, 16 | 2.808  | 0.090 |
| Phenolic C         | 2, 16 | 1.822  | 0.194 |
| Carboxyl C         | 2, 16 | 2.057  | 0.160 |
| Alkyl C: O-alkyl C | 2, 16 | 8.824  | 0.003 |
| Aromaticity        | 2, 16 | 2.654  | 0.101 |

**Figure 2.** The chemical composition of soil organic carbon in three vegetation types. For each chemical shift range, different letters refer to a significant difference ($p < 0.05$).
As we anticipated, the conversion of subtropical native broadleaved evergreen forest to other vegetation types impacted the chemical composition of SOC, where the impact differed between vegetation types. We hypothesized that the conversion of native broadleaved evergreen forests to tea garden may result in stronger alterations.

**Figure 3. Ratio of alkyl C to O-alkyl C and the aromaticity in three vegetation types.** For each index, different letters refer to a significant difference ($p < 0.05$).

**Figure 4. Non-metric multi-dimensional scaling ordination of three vegetation types.** Site scores represent soil organic carbon chemical composition assemblages. The large symbols represent the centroids of all samples from three vegetation types, and lines indicate the SE along each NMDS axis. Circles represent native broad-leaved forests; triangles represent mixed forests, and squares represent tea gardens.

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in SOC chemical composition. It was considered that the disturbance of soil aggregates was much more potent during the conversion to tea gardens than the conversion to mixed forests, which might significantly impact the chemical composition of SOC. Contrary to our hypothesis, the difference of SOC chemical composition in the conversion to mixed forests was more distinct.

This may be partially attributable to the different biochemical content (e.g., lignin or carbohydrates) in the litter of pine needles in the support of its preferred microbial community, as trees can alter soil properties through root-microbe interactions. This resulted in a distinct difference of the chemical composition of SOC in mixed

| Composition       | NMDS 1  | P     | NMDS 2 | P     |
|-------------------|---------|-------|--------|-------|
| Alkyl C           | −0.618  | <0.001| −0.786 | <0.001|
| N-alkyl C         | −0.997  | <0.001| 0.073  | 0.844 |
| Carbohydrate C    | −0.478  | <0.001| 0.878  | <0.001|
| Di-O-alkyl C      | −0.412  | <0.001| 0.911  | <0.001|
| Aryl C            | 0.993   | <0.001| 0.119  | 0.294 |
| Phenolic C        | 0.975   | <0.001| −0.222 | 0.379 |
| Carboxyl C        | 0.840   | <0.001| −0.543 | 0.092 |
| Alkyl C:O-alkyl C | −0.190  | <0.001| −0.982 | <0.001|
| Aromaticity       | 0.997   | <0.001| −0.072 | 0.330 |

Table 4. Correlation matrix of associations between NMDS axes scores and organic carbon chemical composition for three vegetation types. Values indicate Pearson coefficients.

| Characteristic     | NMDS 1  | P     | NMDS 2 | P     |
|--------------------|---------|-------|--------|-------|
| Shannon index      | 0.973   | 0.059 | 0.230  | 0.778 |
| Bulk density (g·cm\(^{-3}\)) | −0.977 | 0.137 | 0.214  | 0.840 |
| pH                 | 0.702   | 0.098 | 0.713  | 0.302 |
| Total organic carbon (g·kg\(^{-1}\)) | −0.293 | 0.605 | 0.956  | 0.319 |
| Total nitrogen concentration (g·kg\(^{-1}\)) | −0.196 | 0.548 | −0.981 | 0.085 |
| Total phosphorus concentration (g·kg\(^{-1}\)) | 0.103  | 0.769 | 0.995  | 0.104 |

Table 5. Correlation matrix of associations between axes scores, tree diversity, and soil physicochemical characteristics for three vegetation types. Values indicate Pearson coefficients.

Figure 5. Non-metric multi-dimensional scaling ordination of soil organic carbon chemical composition in three vegetation types. Black lines with long arrows indicate soil organic carbon chemical composition. Grey lines with short arrows indicate tree diversity and soil physicochemical characteristics. Shannon = Shannon index.
Carboxyl C
Carbohydrate C
Di-O-alkyl C
Alkyl C
Aromaticity
Aromaticity

| Composition | Bulk density | pH | Total carbon concentration | Total nitrogen concentration | Total phosphorus concentration | Shannon's index |
|-------------|--------------|----|---------------------------|-----------------------------|-------------------------------|----------------|
|             | Cor.         | P  | Cor.                      | P                           | Cor.                          | P               |
| Alkyl C     | 0.251        | 0.299 | –0.469    | 0.043                       | –0.025                       | 0.918           | –0.326            | 0.173            | 0.430            | 0.065 |
| N-alkyl C   | 0.336        | 0.16 | –0.476    | 0.039                       | –0.023                       | 0.925           | –0.209            | 0.389            | –0.271            | 0.262  | –0.137 | 0.577 |
| Carbohydrate C | 0.233 | 0.337 | –0.078    | 0.751                       | 0.21                         | 0.389           | 0.1               | 0.683            | 0.313            | 0.192  | –0.309 | 0.201 |
| Di-O-alkyl C | 0.258 | 0.286 | 0.055     | 0.824                       | 0.3                          | 0.213           | 0.101             | 0.682            | 0.175            | 0.474  | –0.219 | 0.375 |
| Aryl C      | –0.355       | 0.136 | 0.405     | 0.085                       | –0.202                       | 0.407           | –0.209            | 0.39            | 0.014            | 0.955  | 0.362  | 0.127 |
| Phenolic C  | –0.173       | 0.48 | 0.297     | 0.216                       | –0.084                       | 0.734           | –0.156            | 0.522            | –0.249            | 0.304  | 0.536  | 0.018 |
| Carboxyl C  | –0.291       | 0.227 | 0.232     | 0.34                        | –0.013                       | 0.958           | –0.069            | 0.78            | 0.219            | 0.368  | 0.418  | 0.077 |
| Alkyl C; O-alkyl C | 0.081 | 0.741 | –0.388    | 0.101                       | –0.112                       | 0.648           | 0.034             | 0.89            | –0.396           | 0.093  | –0.210 | 0.382 |
| Aromaticity | –0.324       | 0.176 | 0.37      | 0.119                       | –0.157                       | 0.521           | –0.202            | 0.407            | –0.018            | 0.942  | 0.456  | 0.05  |

Table 6. Correlations between SOC chemical composition, soil physiochemical characteristics and Shannon’s index. *Correlation is significant at $P < 0.05$.

Our study highlights that vegetation change is a critical factor that impacts the SOC chemical composition in forests ecosystems, where the impacts were found to be variable from native broadleaved forests to other vegetation types following disturbances of short durations and natural succession. The impact of conversion from broadleaved forests to mixed forests on the chemical composition of SOC was stronger than the conversion to tea gardens. The difference of the chemical composition of SOC were primarily represented by alkyl C and aryl C, indicating that alkyl C and aryl C were more liable to change following vegetation changes. The relative abundance of alkyl C was significantly higher, and aryl C was considerably lower in mixed forests than the other vegetation types. The physicochemical characteristics of the soil, as well as tree species diversity were correlated to SOC chemical composition, indicating that they contributed to SOC chemical compositional shifts associated with the vegetation changes. Our results suggest that the conversion of native broadleaved evergreen forests to the other vegetation types following disturbances of short durations and succession may shift the equilibrium states of SOC composition to a different degree, which may potentially drive alterations in SOC and other nutrient cycling in these ecosystems.

Methods

Study area. This study was conducted at the Fengyang Mountain nature reserve, Zhejiang Province, China (119°06′ E to 119°15′E, 27°46′ N to 27°58′ N, 600 m to 1929 m a.s.l.), which comprises an area of 15,171 ha. The nature reserve is characterized as a humid subtropical climate with ~2,400 mm of annual rainfall, and an average annual temperature of 12.3 °C. Prior to 1970, this area was dominated by native evergreen broadleaved forests (composed mainly of Camellia japonica Linn., Cyclobalanopsis multiervis W. C. Cheng et T. Hong, Schima superba Gardn. et Champ., and bamboo plantations at different elevation following complete clearance. Subsequent to the establishment of the nature reserve in 1975, the entire study area, including the tea gardens has been protected from anthropogenic disturbances. There has since been no fire or insect infestation disturbances recorded as yet. An overview of the main vegetation composition of this study area is presented in Supplementary Information Table S1.

Sampling. In June 2013, we randomly sampled nine native evergreen broadleaved forest stands, six mixed-forest stands, and four tea garden stands. All sample stands were located on well-drained and mesic sites with slopes of less than 5% to minimize the site effects in soil characteristics. In each stand, we established a sample plot of 20 × 20 m², in which all trees >2 cm in diameter at breast height (DBH, 1.3 m above root collar) were identified and counted. In each tea garden stand, the tea bush clusters were counted as the quantity of tea bush “individuals”. Smith, et al.30 deemed that the effects of vegetation or land use changes on C content of soil is tree species-dependent. That the conversion of native broadleaved forests to tea gardens resulted in no significant difference in the chemical composition of SOC may be attributed to the less differences in the organic carbon chemical composition in leaves or roots between broadleaved trees35. Collectively, these findings indicate that the vegetation type is a critical factor that influences SOC chemical composition.

The physicochemical characteristics of the soil may favor certain microbial groups over others, thereby leading to shifts in the composition of microbial communities26. In this study, alkyl C was found to be significantly correlated with soil pH, and this may be attributed to the microbial community composition of different pH optima32, which controlled SOC formation processes accordingly31. As we anticipated, tree species diversity contributed to SOC chemical composition. The impact of plant diversity on SOC deposition is chiefly through influence physiochemical and biological pathways5. Briefly, plant diversity may influence the quantity and quality of organic carbon inputs from litterfall and root decomposition, as well as microclimates, to subsequently alter physicochemical characteristics of the soil16. Collectively, the belowground soil microbial biomass and microbial activities are stimulated, leading to differences in SOC composition.
Four sampling points were randomly determined within each plot, where sampling points were at least 8 m apart. At each sampling point, we collected soil samples for nuclear magnetic resonance (NMR) spectroscopy, pH, total organic carbon concentration, total nitrogen concentration, and total phosphorus concentration analysis. Using a sharp knife and a trowel at the depth from 0–10, 10–20, 20–30 cm after removing litter. For bulk density soil analysis, we carefully extracted soil samples with a metal corer at four randomly allocated sampling points (diameter: 5.5 cm, height: 5 cm). Samples for nuclear magnetic resonance (NMR) spectroscopy analysis were stored at −80°C in laboratory until the analysis was conducted.

**Soil physicochemical properties analysis.** Soil physicochemical properties of each plot were determined by the mean value of three layers (0–10, 10–20, 20–30 cm) of the four sampling points. Soil bulk density was determined by drying the samples in an oven at 105°C until a constant weight was achieved, which was then corrected for root and stone volume. Soil samples for other physicochemical analyses were air-dried in the laboratory and then sieved (2 mm mesh) and stored in air-tight plastic bags. Soil pH was measured using a PB-10 pH meter (Sartorius GmbH, Göttingen, Germany) at a 1:5 ratio of soil to water. The total organic carbon concentration was determined using the sulfuric acid-potassium external heating method. Total nitrogen and phosphorus concentrations were determined using a Bran+Luebbe Autoanalyzer 3 Continuous Flow Analyzer (Bran+Luebbe GmbH, Norderstedt, Germany) according to the manufacturer’s instructions.

**Nuclear magnetic resonance (NMR) analysis.** The 13C NMR spectroscopy was conducted on composite samples, which were obtained by mixing three layers and four sampling points from each sampling plot for analysis. Similar to physicochemical analysis, the mixed samples were air-dried in the laboratory and then sieved (2 mm mesh) and stored in air-tight plastic bags in a refrigerator at 4°C for further investigation. To enhance the signal to noise ratio of the instrument, the hydrofluoric acid (HF) pretreatment was performed to remove Fe3+ and Mn2+ from the soil. Samples were processed according to previously published methods. Briefly, 5 g of air-dried soil samples were weighed and transferred into a 100 ml plastic centrifuge tube for the HF pretreatment, followed by the addition of 50 ml HF solution (10% v/v) into the tube. Following vibratory agitation for 1 h and centrifugation for 10 min, the supernatant liquid was removed from the tube, and the residue was continuously treated with a HF solution. These steps were repeated 8 times, but varied in the vibration time (4 × 1 h, 3 × 12 h, 1 × 24 h). Soil samples were rinsed 4 times with double distilled water to remove any residual HF that remained. Subsequently, the residues were dried in oven at 40°C. Following all of the above processes, the residues were ground to the extent that they could pass through a 60-mesh screen, and then loaded into a bag for NMR measurement. The HF-treated soil samples were subjected to 13C NMR analysis with a Bruker (Spectrospin, Rheinstetten, Germany) Avance 600 MHz NMR spectrometer. The experiments were carried out using a 7 mm CPMAS probe, at a carbon frequency of 150 MHz, MAS spinning frequency at 6 kHz, contact time of 2 ms, and a recycle delay time of 5 s.

Similar to previous NMR studies, the NMR spectra were divided into the following regions: alkyl C region (0–45 ppm), N-alkyl C region (45–60 ppm), carbohydrate C region (60–90 ppm), di-O-alkyl C region (90–110 ppm), aryl C region (110–145 ppm), phenolic region (145–165 ppm), carboxyl C region (165–215 ppm). The relative intensities of the different SOC fractions were obtained through the measurement and integration of the area under the curve for each region. Additionally, two indices of organic carbon stability were calculated: the ratio of alkyl C to O-alkyl C (A/O-A) and aromaticity, which has been used as an indicator of organic carbon decomposition potential and aromaticity.

**Data analysis.** We quantified plant species diversity by employing the Shannon’s index ($H'$):

$$H' = -\sum_{i=1}^{n} p_i \times \ln(p_i)$$

where $p_i$ is the proportion of the individuals of $i$th overstory species in the plot based on stem counts, and $n$ is the number of overstory species in the plot, and the index was calculated for each sampling plot. To examine the impacts of vegetation changes on the Shannon index of overstory plants, soil physicochemical characteristics and each SOC chemical composition, one-way analysis of variance (ANOVA) was conducted, followed by a LSD test ($P < 0.05$).

Multiple-response permutation procedures analysis (MRPP) was performed to test the difference of SOC chemical composition of among three vegetation types. The relationship between SOC chemical composition, tree diversity, and soil physicochemical characteristics was analyzed by non-multidimensional scaling (NMDS). Pearson correlation tests were also applied to explore the impacts of the chemical composition of SOC, tree diversity, and soil physicochemical characteristics variables on SOC chemical composition assemblages of NMDS axes. This was achieved by calculating Pearson correlation coefficients between all of the variables and NMDS axes scores. To examine the chemical compositions of SOC, tree diversity, and soil physicochemical characteristics variables, which influenced the chemical composition of SOC assemblages, linear regressions test were performed. To further discover the potential impact of tree diversity and soil physicochemical characteristics on SOC chemical composition, Pearson correlations analysis was also conducted between all components of SOC chemical composition, tree species diversity, and soil physicochemical characteristics. All analyses were performed in R (vegan package).
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**Author Contributions**
J.Z. and X.G. designed the experiments. X.G. and M.M. conducted the experiments. All authors wrote the early version of manuscript draft. H.Y.H.C., M.M. and X.G. wrote the revised version. H.Y.H.C. and X.G. conducted all the data analysis. X.G. and M.M. prepared all figures. All authors reviewed the manuscript.

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