

Molecular-level carbon traits of fine roots: unveiling adaptation and decomposition under flooded conditions

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Abstract. Fine roots are vital for plant development and carbon biogeochemical cycling in terrestrial ecosystems. Flooding is known to regulate the physiology and morphology in plant roots; however, its impact on molecular-level characteristics of carbon compounds (carbon traits) in fine roots remains largely unexplored, which limits our understanding of root adaptation and decomposition under changing environments. Here, we used a sequential extraction method, starting from nonpolar to polar solvents, in order to obtain dichloromethane- and methanol-extractable (F_{DcMe}) fractions, base-hydrolyzable (F_{KOHhy}) fractions, and CuOoxidizable (F_{CuOox}) fractions from fine roots of Dysoxylum binectariferum, which is naturally grown in soil and water. Subsequently, we characterized them using targeted gas chromatography-mass spectrometry and nontargeted Fourier transform ion cyclotron resonance mass spectrometry. Also, decomposition experiments were conducted on soil- and water-grown roots under aerobic and anoxic conditions. Results showed a consistent increase in the unsaturation degree and aromaticity of the analytes from F_{DcMe} to F_{CuOox} fractions. Both analyses were sufficiently sensitive to show that, compared to soil-grown roots, the water-grown roots developed more polyphenolics with a high unsaturation degree and aromaticity and had more nonstructural compositions. Furthermore, although flooding provided an anoxic condition that slowed down root decomposition, the adaptive strategy of developing more nonstructural labile components in water-grown roots accelerated root decomposition, thereby counteracting the effects of anoxia. This advances our understanding of biogeochemical processes in response to global environmental change.

1 Introduction

Fine roots are an important part of the plant root system owing to their essential functions in resource absorption and transportation (McCormack et al., 2015). Additionally, with the highest productivity and turnover rate among all underground plant organs, fine roots considerably contribute to organic matter formation and stabilization in soils. Thus, they drive the cycling and distribution of carbon and nutrients in terrestrial ecosystems (Dijkstra et al., 2021). However, the fine-root tissue chemistry is poorly understood (McCormack et al., 2015), significantly hindering our ability to accurately forecast the biogeochemical processes in terrestrial ecosystems.

Carbon compounds are highly diverse in fine roots. We define fine-root carbon traits as the composition and characteristics of carbon compounds in fine roots (Freschet et al., 2021a, b; Wang et al., 2024). The utilization of an extraction method, starting from polar to nonpolar solvents (Otto and Simpson, 2007), revealed a high level of heterogeneity in carbon compounds of fine roots, associated with both

root architecture (Wang et al., 2015) and ancestry (Mueller et al., 2012). Additionally, previous studies have elucidated the prominent influence of carbon compounds (e.g., terpenes, lignin, and bound phenolics) on root decomposition and, thus, on the terrestrial soil carbon cycling and coevolution between plants and their consumers, microbial symbionts, or humans (Sun et al., 2018; Cornelissen et al., 2023; Augusto and Boca, 2022; Jing et al., 2023). Therefore, identifying the variations in the compositional carbon traits of fine roots is important for predicting the function and fate of fine roots in terrestrial ecosystems.

Global climate change has induced more frequent extreme precipitation and flooding events (Mann et al., 2018), which can cause oxygen deprivation in plants. Roots are directly affected by such events that cause hypoxia and anoxia stress, consequently constraining plant growth and development (Herzog et al., 2016). Fortunately, plants can adapt to hypoxia by changing the morphology and anatomical structure of the root system (Pedersen et al., 2021; Yamauchi et al., 2021), such as the outgrowth of adventitious roots, the formation of aerenchyma, a higher cortex-to-stele ratio, and a lower surface-area-to-volume ratio (Colmer et al., 2019; Yamauchi et al., 2019). The alterations in morphology and anatomy are expected to reflect the inevitable alterations in root tissue chemistry. For example, deoxygenated conditions increase the absolute contents of suberin and lignin in the outer part of the roots in rice (Oryza sativa L.; Kotula et al., 2009). However, unlike the well-documented physiological and morphological plasticity, the molecular-level chemical plasticity in plant roots in flooded environments still lacks a holistic understanding.

The decomposition of plant litter is a fundamental ecosystem process, the rate of which is known to be regulated by environmental change via reaction kinetics and microbial physiology (Suseela and Tharayil, 2018). Previous studies suggested that flooding caused limitation of oxygen-inhibited aerobic respiration of biological communities (Wen et al., 2006) and slowed the rate of litter decomposition (Jessen et al., 2017), resulting in the accumulation of organic carbon and declination of CO₂ flux (Iqbal et al., 2009). In fact, the oxygen requirement of decomposition varies with different chemical compositions in plant litter (LaRowe and Van Cappellen, 2011). The labile carbon compounds (profitable energy substances such as carbohydrates, free amino acids, and proteins) are preferably decomposed compared with recalcitrant carbon compounds (complex aromatic biopolymers such as lignin and suberin), where the decomposition depends strongly on oxygen (Agethen and Knorr, 2018; Kirschbaum et al., 2021). As mentioned above, the flooded environments are expected to alter the chemical composition of root litter and, thus, indirectly impact decomposition. However, whether this indirect effect strengthens or counteracts the direct effect of anoxia on litter decomposition rate is poorly understood.

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Numerous plants naturally thrive at the waterline within tropical forests, offering an exceptional opportunity to examine and verify the responsiveness of the fine roots of plants to environmental fluctuations. Here, soil-grown fine roots (SGR) and water-grown fine roots (WGR) of Dysoxylum binectariferum Hook.f., an evergreen plant with medicinal value that is common in China, India, and other parts of Asia (Coffin and Ready, 2019; Bharate et al., 2018), were subjected to a series of extraction and chemical degradation steps and then analyzed using targeted (electron ionization gas chromatography-mass spectrometry, GC-MS) and nontargeted (Fourier transform ion cyclotron resonance mass spectrometry, FT-ICR MS) techniques in order to determine their molecular signatures. In addition, the residues after each degradation step were analyzed using an elemental analysis and solid-state ¹³C cross-polarization magic angle spinning nuclear magnetic resonance (NMR) to determine the changes in carbon content and bulk molecular composition during the extraction procedures, respectively. We tested two main hypotheses: (1) compared to SGR, WGR should contain more aromatics, especially in bound compounds, in order to cope with flooding stress; and (2) the WGR would have a lower decomposition rate than the SGR in both aerobic and anoxic conditions. This study establishes a comprehensive tool resolving the molecular-level carbon traits of fine roots that will extend knowledge of the possible mechanism underlying the response of the belowground carbon cycling to global environmental change.

2 Materials and methods

2.1 Study site and sample collection

The study site was in a tropical seasonal rain forest in Xishuangbanna (21°41′ N, 101°25′ E), southwestern China. The forest receives an average annual precipitation of 1539 mm and has an average annual temperature of 21.5 °C. The soils are classified as Oxisols in the soil taxonomy. We collected root samples from three mature trees of Dysoxylum binectariferum Hook.f. that were growing by a slow-flowing stream, with roots growing in both soil and water (Fig. S1 in the Supplement; see Text S1 in the Supplement for more details). At the same time, the soil and sediment near the roots were also collected. All fine roots, soils, and sediments were immediately collected in plastic bags, transported in a cooler to the laboratory, and frozen at -20 °C. Root morphology parameters were acquired by a root image analysis using WinRHIZO software (2007 Pro version; Regent Instruments, Québec, QC, Canada). Before chemical analyses, the fine roots were gently cleaned in cool water, freeze-dried, and ground into fine powder.

2.2 Isolation and targeted analysis by GC-MS of individual compounds

The compounds in fine roots were separated into three fractions via sequential extraction (Fig. 1) based on the methods of Otto and Simpson (2007) and Wang et al. (2015), with modifications. Triplicate extractions of each root sample were conducted. Extraction of the roots with organic solvents followed by alkaline hydrolysis can obtain the free compositions (such as free lipids, steroids, and carbohydrates) and ester-bound compositions (cutin and suberin monomers), respectively. The CuO oxidation yields etherbonded compositions (preferentially lignin phenols). Briefly, solvent extraction by dichloromethane (DCM), methanol (MeOH): DCM (1:1, v/v), and MeOH consecutively for the dichloromethane- and methanol-extractable fraction (F_{DcMe}) was performed with homogenized root samples (R1). The root residues (R2) after solvent extraction were hydrolyzed with $1 \text{ mol } L^{-1}$ methanolic KOH (100 °C, 3 h) for the basehydrolyzable fraction (F_{KOHhy}). Dried root residues (R3) were further subjected to alkaline cupric oxide (CuO) oxidation with CuO, $Fe(NH_4)_2(SO_4)_2 \cdot 6H_2O$, and $1 \text{ mol } L^{-1}$ NaOH for the CuO-oxidizable fraction (F_{CuOox}). The isolated individual compounds were derivatized before GC-MS analysis using an Agilent 7890B gas chromatograph equipped with a 5977B mass spectrometer. Further details are given in Text S2 in the Supplement.

2.3 FT-ICR MS analysis and molecular formula assignment

Aliquots of the extracts (F_{DcMe} , F_{KOHhy} , and F_{CuOox}) were dissolved in MeOH at a target carbon concentration of approximately 50 µg mL⁻¹. The solution was infused into a 9.4 Tesla Bruker Apex-Ultra FT-ICR MS instrument (Bruker, Germany) using the ESI mode for ionization. Signals were accumulated with 128 scans, a 2 M word size, and a mass range of m/z 200–800.

Mass peaks were internally calibrated with a target mass measurement accuracy of < 1.0 ppm. The signal-to-noise threshold and absolute intensity threshold for peak selection were set as 7 and 100, respectively. Following the compound identification algorithm (Kujawinski and Behn, 2006), elemental formulae were assigned using the Formularity software (Tolić et al., 2017), with the additional requirements of ${}^{12}C_{1-60}$, ${}^{1}H_{1-120}$, ${}^{14}N_{0-3}$, ${}^{16}O_{0-30}$, and ${}^{32}S_{0-1}$. The double bond equivalent (DBE), modified aromaticity index (AImod), and nominal oxidation state of carbon (NOSC) were calculated to reveal the degrees of unsaturated states (Koch and Dittmar, 2006, 2016), aromaticity (Koch and Dittmar, 2006, 2016), and relative nominal oxidation state of organic matter (Kellerman et al., 2015), respectively. Furthermore, the unique formulae in SGR and WGR were identified and analyzed based on Kendrick mass defect (KMD) plots in order to display the reaction trends of (1) hydrogenation/dehydrogenation (H₂ series), (2) methylation/demethylation (CH₂ series), (3) ring-opening reactions (O₂ series), and (4) carboxylation/decarboxylation (COO series; Waggoner and Hatcher, 2017; Khatami et al., 2019).

2.4 Solid-state ¹³C-NMR analysis

Approximately 100 mg of root sample (R1) or residue after each extraction step (R2, R3, and R4) was packed into a 4 mm zirconium rotor with a Kel-F cap. The solid-state ¹³C NMR spectra were acquired using a Bruker Avance III 600 spectrometer at a resonance frequency of 150.9 MHz, with a 4 mm magic angle spinning probe, which was spun at 12 kHz. A contact time of 4 ms and a recycle delay of 2 s were used for a ramp cross-polarization magic angle spinning NMR measurement. The spectra were divided into seven central resonance regions according to the chemical shift as follows (Nelson and Baldock, 2005): 0-45 ppm for alkyl carbon, 45–60 ppm for N-alkyl/methoxy carbon, 60–95 ppm for O-alkyl carbon, 95–110 ppm for di-O-alkyl carbon, 110– 145 ppm for aromatic carbon, 145–165 ppm for O-aromatic carbon, and 165–210 ppm for carbonyl and carboxyl carbon. The relative abundance of these seven carbon regions was then calculated.

2.5 Fine-root decomposition

Both SGR and WGR were incubated in a mixed matrix of A horizon soil near SGR and surface sediment near WGR under anoxic or aerobic conditions. Before incubation, any visible plant residues and stones with a diameter of > 2 mmwere removed from the soil and sediment. Then the soil and sediment were homogenized based on a 1:1 fresh mass ratio. For anoxic conditions, a glass tube (4.5 cm diameter \times 10 cm depth) was filled with 100 g of the freshly mixed matrix (65 g in dry weight) and 80 mL of deionized water (5 cm of overlying water). For aerobic conditions, the fresh mixture matrix was air-dried to 60 % water holding capacity (WHC) and then filled into another glass tube (65 g in dry weight). Anoxic and aerobic incubation experiments of the SGR and WGR were set up with three replicates, which resulted in 12 microcosms in total. Then, 1 g of freeze-dried root sample was sealed in a clean polyamide litterbag $(3.5 \text{ cm} \times 7 \text{ cm})$ 40 µm mesh). All litterbags were completely buried in microcosms and in close contact with the mixed soil and sediment matrixes. The experimental microcosms were kept in a dark incubator at 25 °C. The mixed matrixes in an aerobic condition were held at 60 % WHC, while those in an anoxic condition were held with 5 cm of overlying water via weekly additions of deionized water. All microcosms were harvested after 180 d. After that, decomposed fine roots were collected, cleaned, and freeze-dried, after which they were weighed and ground for carbon content analysis.



Figure 1. Schematic chart of the sequential extraction of organic compounds from fine roots to obtain different operational fractions and residues. GC-MS: gas chromatography–mass spectrometry. FT-ICR MS: Fourier transform ion cyclotron resonance mass spectrometry. ¹³C-NMR: ramp solid-state ¹³C cross-polarization magic angle spinning nuclear magnetic resonance. F_{DcMe} : dichloromethane- and methanol-extractable fraction. F_{KOHhy} : base-hydrolyzable fraction. F_{CuOox} : CuO-oxidizable fraction.

2.6 Statistical analysis

Significant differences in the chemical characteristics between SGR and WGR were assessed using a two-way independent-samples *t* test. Furthermore, significant differences among the three fractions were analyzed using a twoway analysis of variance (ANOVA) followed by Tukey's post hoc test for pairwise comparisons. All statistical analyses were conducted using IBM SPSS Statistics 23. Statistical difference was considered when p < 0.05.

3 Results

3.1 Biomass and carbon fractions in sequential extraction steps

Each sequential extraction step could extract considerable proportions of biomass and carbon for all root samples. F_{KOHhy} and F_{CuOox} had similar (p > 0.05) and the highest biomass fractions, accounting for 34 % and 32 % of the total biomass, respectively, while F_{DcMe} had the lowest biomass fraction with 10 % of the total biomass (Fig. 2a; Table S1 in the Supplement). There was approximately 24 % of biomass residue after the three-step extraction. Regarding the chemical element composition, nearly 90 % of carbon was extracted via the three-step chemical extraction, and more than 70 % of carbon was evenly distributed in F_{KOHhy} and F_{CuOox} (Fig. 2b; Table S1). Although WGR had near twice the number of biomass and carbon fractions in F_{DcMe} of SGR (p <0.05), the two classes of roots were similar in both biomass and carbon content in the other three fractions (p > 0.05; Table S1). The proportion of ash content in SGR was approximately twice that in WGR (p < 0.05; Fig. 2c; Table S1).

Solid-state ¹³C NMR spectroscopy showed the variations in carbon structure in different extracted fine root residues (Fig. 3; Fig. S2 in the Supplement). After solvent extraction, the intensity of the alkyl carbon region significantly decreased (p < 0.05) in all fine-root samples. Base hydrolysis significantly reduced the signal intensity in the carbonyl and carboxyl carbon region in SGR and the *O*-aromatic carbon region in WGR (p < 0.05), which indicated that these structures were extracted by base hydrolysis. The carbonyl and carboxyl and aromatic carbon regions in SGR decreased after CuO oxidation, as did the carbonyl and carboxyl and *N*-alkyl/methoxy C regions in WGR (p < 0.05).

3.2 Composition based on GC-MS analyses

The dichloromethane- and methanol-extractable fractions (F_{DcMe}) contained more than 70 compounds detected by GC-MS and identified by comparing mass spectra with mass spectra from the NIST library (Table S2 in the Supplement). Based on their functions, chemical characteristics, and specificities, according to Macabuhay et al. (2022), these compounds in F_{DcMe} were subdivided into six principal categories: phenolics, glycerolics, prenolics, carbohydrates, fatty acyls, and steroids (Fig. 4a). The total concentration of these compounds in F_{DcMe} reached approximately 100 mg g⁻¹ OC in all fine roots. Compared with SGR, WGR had significantly more phenolics (p = 0.007), prenolics (p < 0.001), steroids (p < 0.001), and total lipids (p = 0.001) but fewer glycerolics (p = 0.028).



Figure 2. Average biomass fractionation (a) and fractional carbon contents (b) based on sequential extraction and ash content (c) of watergrown (WGR, blue bar) or soil-grown roots (SGR, red bar). Four shades, from light to dark, indicate the dichloromethane- and methanolextractable fraction (F_{DcMe}), base-hydrolyzable fraction (F_{KOHhy}), CuO-oxidizable fraction (F_{CuOox}), and residue after CuO oxidation. Bars represent means \pm standard errors (n = 3).



Figure 3. Solid-state ¹³C cross-polarization magic angle spinning nuclear magnetic resonance spectra of fine-root samples (R1) and the residues after dichloromethane and methanol extraction (R2), base hydrolysis (R3), and CuO oxidation (R4). The shadowed areas represent the range of means \pm standard errors (n = 3).

The base-hydrolyzable fractions (F_{KOHhy}) yielded 23 aliphatic compounds and 15 phenolic compounds (Table S2), with a total concentration of 20–30 mg g⁻¹ OC in all fine roots (Fig. 4b). The aliphatic compounds predominantly comprised even-numbered, mid- to long-chain homologs (C₁₀–C₂₆) of *x*- and ω -hydroxyalkanoic acids (*x*-/ ω -OH

acids), *n*-alkanols, saturated normal fatty acids (FAs), saturated normal fatty diacids (DAs), and unsaturated normal fatty acids (UAs). Compared with SGR, WGR were significantly enriched in more phenolics (p < 0.001) but in fewer aliphatics (p = 0.022).



Figure 4. Molecular-level compositions of dichloromethane- and methanol-extractable fractions (F_{DcMe}) (**a**), base-hydrolyzable fractions (F_{KOHhy}) (**b**), and CuO-oxidizable fractions (F_{CuOox}) (**c**) of soil-grown roots (SGR, in red) and water-grown roots (WGR, in blue). The *y* axes represent organic carbon (OC)-normalized concentrations. Bars represent mean \pm standard error (n = 3). The coefficient above each pair of bars is the *p* value of two-way *t* test analyses of the significant differences between SGR and WGR. *x-/ω*-OH acids: *x-* or *ω*-hydroxyalkanoic acids. FAs: saturated normal fatty acids. DAs: saturated normal fatty diacids. UAs: unsaturated normal fatty acids. PAL: *p*-hydroxybenzaldehyde. PON: *p*-hydroxyacetophenone. PAD: *p*-hydroxybenzoic acid. VAL: vanillin. VON: acetovanillone. VAD: vanillic acid. SAL: syringaldehyde. SON: acetosyringone. SAD: syringic acid. CAD: *p*-coumaric acid. FAD: ferulic acid. Detailed concentration data of individual compounds are shown in Table S2.

A total of 10 phenols (total concentration of $27 \pm 5 \text{ mg g}^{-1} \text{ OC}$) were the predominant products in F_{CuOox} after CuO oxidation, with CAD not detected (Fig. 4c; Table S3 in the Supplement). The eight lignin phenols (except PAL, PON, and PAD) amounted to $24 \pm 5 \text{ mg g}^{-1} \text{ OC}$ of the root samples. Of these phenols, vanillyl phenols (VAL, VON, and VAD) were the main components, accounting for more than 60% of all phenols, followed by syringyl phenols (SAL, SON, and SAD) that accounted for over 20%

of the total concentration. The roots from different growing media differed in the composition of monomeric phenols. Compared with SGR, WGR had more *p*-hydroxyl phenols (p = 0.047) and vanillyl phenols (p = 0.072) and exhibited higher values of acid-to-aldehyde ratios of vanillyl and syringyl phenols (i.e., (Ad : Al)_V and (Ad : Al)_S); p = 0.056 and p = 0.020, respectively) but lower syringyl-to-vanillyl ratio (S : V) values (p = 0.002; Table S3).

3.3 Molecular composition based on FT-ICR MS analyses

The FT-ICR MS spectra of all samples provided 8631 assigned formulae (Fig. S3; Table S4 in the Supplement). From F_{DcMe} to F_{CuOox} , the average H/C ratio (weighed by peak intensity) continuously decreased, and the average O/C ratio, DBE, AI_{mod}, and NOSC continuously increased (p < 0.05) (Table S5 in the Supplement), demonstrating the increasing unsaturation degree and aromaticity. In the three fractions, the formulae containing only carbon, hydrogen, and oxygen (CHO) accounted for more than half of the total number and nearly 90% of the total abundance of all assigned formulae (Table S5). In addition, there was a considerable quantity of nitrogen-containing and/or sulfur-containing compounds (CHON, CHOS, and CHONS), but these only accounted for approximately 1/10 of the total formulae abundance. Compared with SGR, WGR had a higher average O/C ratio, DBE, AI_{mod}, and NOSC and a lower average H/C ratio (p < 0.05; Table S5), which represented a higher unsaturation degree and aromaticity in WGR. However, there were lower proportions of sulfur-containing compounds (CHOS and CHONS) in WGR than in SGR.

The common formulae in all three replicates covered an average of 76 % and 90 % of the total number and signals in a replicate, respectively (Fig. S3; Table S4), and, thus, had a strong molecular characteristic representativeness. A total of 4368 common formulae (2500 in F_{DcMe}, 2070 in F_{KOHhy}, and 4214 in F_{CuOox}) in the three SGR replicates and 3461 (2594 in F_{DcMe} , 1962 in F_{KOHhy} , and 1746 in F_{CuOox}) in the three WGR replicates were identified (Fig. S4 in the Supplement). The number of coexisting formulae in the three fractions (F_{DcMe}, F_{KOHhy}, and F_{CuOox}) accounted for approximately 30% of the total formulae in both SGR and WGR. More aromatic formulae with low average H/C ratios and high average O/C ratios were present in the F_{CuOox} of SGR and F_{KOHhy} and F_{CuOox} of WGR than that in the F_{DcMe} (Fig. 5; S4 in the Supplement). Compared with SGR, WGR had more formulae with high O/C ratios and low H/C ratios in specific and common formulae in the three fractions (Fig. 5).

Kendrick mass defect (KMD) plots showed a unimodal clustering of KMD plotted against m/z stretching from the molecular formulae with a high KMD for aliphatics as compared to those with a low KMD for aromatics (Fig. 6). A considerable number of new molecules existed in WGR compared with SGR, which exhibited a definite shift towards lower m/z values for the KMD CH₂ and KMD H₂ series (Fig. 6a–f) and higher m/z values for the KMD O₂ and KMD COO series (Fig. 6g–l). The results suggested the oxidation of aliphatic chains and aromatic rings as well as the ring opening of aromatic rings and carboxylation of new molecules in WGR.

3.4 Decomposability of fine roots

To investigate the influencing factors regulating fine-root decomposition, we detected the morphological traits and nitrogen content in origin roots. The results showed that SGR have a higher specific root length and nitrogen content, with a lower average diameter and carbon-to-nitrogen ratio than WGR (p < 0.05; Table S6 in the Supplement). After 180 d of decomposition, the two-way t test analyses revealed that the redox condition and root source significantly affected the percentage of root carbon remaining (Table S7 in the Supplement). Both kinds of fine roots had more proportion of carbon remaining in the anoxic condition than in the aerobic condition (Fig. 7). WGR had 14 % and 13 % less carbon remaining in the aerobic condition (p = 0.003) and anoxic condition (p < 0.001) than SGR, respectively. Therefore, the decomposability of organic carbon was consistently higher in WGR than in SGR.

4 Discussion

4.1 Extended information captured by nontargeted analyses of root fractions

Fine roots contain abundant carbon compounds with heterogeneous and macromolecular structures. In this study, the biomass and carbon directly extracted using organic solvents only accounted for 11 % and 16 % of the total biomass and total carbon in roots, respectively (Fig. 2). This indicated that most biomass and carbon were distributed in the nonfree biopolymers of roots. Thus, the carbon compounds obtained via simple solvent extraction could hardly represent the full characteristics of fine roots. Alkaline hydrolysis is an established method to obtain many ester-bound compounds from organic heteropolymers (Otto and Simpson, 2006), after which the oxidation with CuO preferentially yields mostly ether-bound monomers (Kaiser and Benner, 2012). A combination of the three approaches selectively broke down and extracted approximately 76 % of the total biomass and 85 % of the total carbon in roots (Fig. 2). The residue after the threestep sequential extraction contained mostly O-alkyl and di-O-alkyl carbon (Fig. 3), which was widely attributed to the contribution of cellulose and hemicellulose (Foston, 2014). Our results suggest that the chemical depolymerization is a practical approach to obtaining carbon fractions from root biopolymers for further characterization.

Upon combining the GC-MS and FT-ICR MS, our results showed that the compositional carbon traits of fine roots after sequential chemical extraction were further supported by the ¹³C-NMR data. For example, the results of targeted and nontargeted approaches consistently indicated that the unsaturation degree and aromaticity of components increased from F_{DcMe} to F_{CuOox} (Fig. 4; Table S5), which corresponded to a decrease in the alkyl carbon region from R1 to R2 and



Figure 5. Comparison of van Krevelen diagrams of soil-grown roots (SGR, in red) and water-grown roots (WGR, in blue) for specific formulae detected by Fourier transform ion cyclotron resonance mass spectrometry (FT-ICR MS) from (a) dichloromethane- and methanol-extractable fractions (F_{DcMe}), (d) base-hydrolyzable fractions (F_{KOHhy}), and (g) CuO-oxidizable fractions (F_{CuOox}). Shown are van Krevelen diagrams of formulae that are common but have a higher relative abundance in SGR than in WGR (b, e, h) or a higher relative abundance in WGR than in SGR (c, f, i).

a further decrease in the carbonyl and carboxyl carbon, *O*aromatic carbon, and aromatic carbon regions of residues from R2 to R4 (Figs. 3 and S2). GC-MS results showed that, based on the qualitative and quantitative detection of specific compounds in the extracted fractions, 78, 38, and 10 compounds were identified and quantified in F_{DcMe} , F_{KOHhy} , and F_{CuOox} , respectively (Fig. 4; Table S2). As a complementary tool, FT-ICR MS is a nontargeted technique that has sufficient mass-resolving power to separate and accurately assign elemental compositions to individual molecules (Qi et al., 2022; He et al., 2022). FT-ICR MS spectra provided a total of 8631 assigned formulae in all samples (Fig. S3; Table S4), which was approximately 100 times more than the total number of components identified by GC-MS. In addition, a considerable number of nitrogen- and sulfur-containing compounds (CHON, CHOS, and CHONS) were also identified (Table S5), which were not well captured by GC-MS. Therefore, concordant with our first hypothesis, the nontargeted and targeted analyses were consistent and complementary in showing analyte polarity and an unsaturation degree after sequential chemical extraction. These two techniques, combined with sequential chemical extraction, significantly extended the analytical windows of the compositional carbon traits of fine roots.

4.2 Different compositional carbon traits between SGR and WGR

Environmental stress elicits adaptive responses from plants via the preferential biosynthesis and production of specific



Figure 6. Kendrick mass defect (KMD) plots of soil-grown roots (SGR, in red) and water-grown roots (WGR, in blue) for specific formulae detected by Fourier transform ion cyclotron resonance mass spectrometry (FT-ICR MS) from (**a**, **d**, **g**, **j**) dichloromethane- and methanol-extractable fractions (F_{DcMe}), (**b**, **e**, **h**, **k**) base-hydrolyzable fractions (F_{KOHhy}), and (**c**, **f**, **i**, **l**) CuO-oxidizable fractions (F_{CuOox}). The four rows display the KMD for H₂, CH₂, O₂, and COO series.

compounds from assimilated carbon, which provide efficient protection against specific stressors (Suseela and Tharayil, 2018). This study demonstrated that, compared with SGR, WGR contained a high proportion of free compounds and free carbon (Fig. 2a and b), specifically more free phenols, terpenoids, and steroids (Fig. 4a). It is consistent with a previous finding that plants exposed to environmental stress (including waterlogging) tended to produce more extractable labile compounds to counter the environmental imbalance than those growing in optimal conditions (Suseela and Tharayil, 2018). However, WGR contained less ash (Fig. 2c) and fewer free organic acids than SGR (Fig. 4a). In general, one of the functions of endogenous organic acids in roots is to modulate the imbalance of cellular electroneutrality and osmotic pressure caused by a large number of cations absorbed from the medium (Crawford and Tyler, 1969). The relatively lower concentrations of organic acids in WGR than in SGR tend to match the low solute concentration and low osmotic pressure in an aquatic medium.

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Figure 7. Percentage of carbon remaining in fine roots collected from soil (in red) and water (in blue) after 180 d of decomposition in aerobic and anoxic conditions. Bars represent mean \pm standard error (n = 3). The coefficient above each pair of bars is the p value of two-way t test analyses of the significant differences between root sources.

In addition, the whole plant structural matrix is notably altered under environmental stress (Le Gall et al., 2015) because many recalcitrant heteropolymers, including lignin and suberin phenolics, are biosynthesized via the phenylpropanoid pathway and can be directly regulated by environmental stressors (Dong and Lin, 2021) including flooding (Yamauchi et al., 2019), drought, and salt stresses (de Silva et al., 2021). Our results showed that, compared with $F_{\rm KOHhv}$ of SGR, WGR had higher average values of DBE, AI_{mod}, and NOSC (p < 0.05; Table S5) and more formulae (both specific and common formulae but had over 20 % more relative intensity) with high O/C ratios and low H/C ratios (Fig. 5d-f), especially approximately twice more phenols ion $F_{\rm KOHhv}$ (Fig. 4b; Table S2), indicating more biosynthesis and deposition of suberin phenolics (for example, ferulic acids) in WGR (Ranathunge et al., 2011). Although there were no significant differences in the concentrations of eight lignin phenols in F_{CuOox} between two classes of roots (Fig. 4c; Table S3), F_{CuOox} of WGR had higher average values of DBE, AI_{mod}, and NOSC (p < 0.05; Table S5) and more formulae with high O/C ratios and low H/C ratios (Fig. 5g-i) than that of SGR. These were consistent with the previous study based on histochemistry and microscopy, which suggested that apoplastic barriers with suberin lamellae and secondary lignification in the cortex of fine roots could promote adaptation of Metasequoia glyptostroboides to aquatic environments (Yang et al., 2019). Moreover, higher oxidation and carboxylation were observed in specific formulae of WGR (Fig. 6), suggesting increased oxidative stress and further regulation of the shikimate pathway under flooding conditions (Karlova et al., 2021; Dong and Lin, 2021).

Notably, the nontargeted analyses also showed that there were over two-thirds less quantity and abundance of sulfur-

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containing compounds (CHOS and CHONS) in WGR than in SGR (p < 0.05; Table S5). This may be because the lower sulfur content in water rather than in soil resulted in reduced sulfur absorption and assimilation in plant roots (Hawkesford and De Kok, 2006). Thus, it is necessary to monitor possible sulfur deficiencies in trees resulting from flooding-prone regions under future global environmental change.

In summary, our results partially supported the second hypothesis: waterlogging altered the distribution and accumulation of free and bound components. The waterlogging tends to increase free carbon fractions (especially free phenols, terpenoids, and steroids) and bound carbon fractions (e.g., bond phenols and the $V : \Lambda 8$ ratio) but decrease the contents of free organic acids, bond aliphatics, and the $S : \Lambda 8$ ratio in lignin-derived phenols, supporting the fact that plants remarkably regulate biomass and carbon partitioning under changing environments (Karlova et al., 2021).

4.3 Impact of molecular-level carbon traits of fine roots on decomposition

Plant organic matter is the most important source of soil organic carbon pool. The environmental condition was a key factor affecting litter decomposition (Table S7). In this study, fine roots had slower decomposition rates of carbon in anoxic conditions than in aerobic conditions (Fig. 7). Similar to our findings, many studies also observed a general negative effect of O₂ limitation on aerobic respiration (LaRowe and Van Cappellen, 2011) and organic matter decomposition in soil (Kirschbaum et al., 2021; Sjøgaard et al., 2017), wetlands (LaCroix et al., 2019), and marine sediment (Jessen et al., 2017). Moreover, environmental stressors can alter the structure and metabolic activity of the decomposer community (Chuckran et al., 2020). For example, flooding of soils induced the function increase in CH₄ production, S cycle, and Fe cycle (e.g., Bathyarchaeota, Bacteroidetes, and Geobacter) and the function decrease in environmental remediation and biological control (e.g., Blastococcus and Roseiflexus) (Luo et al., 2023), which consequently regulated organic carbon sequestration and biogeochemical cycling.

Our observation that decomposition was also strongly affected by root type (Table S7) supports the view that litter properties were one of the overriding factors in decomposition (Huys et al., 2022; Rummel et al., 2020). Although conventional wisdom held that high specific root length and a lower carbon nitrogen ratio commonly correlated with fast decomposition rates (Guo et al., 2021), our decomposition experiment exhibited that the organic carbon decomposability was higher in WGR with lower specific root length and a higher carbon nitrogen ratio than in SGR (Fig. 7; Table S6). Therefore, the commonly used parameters, like morphological traits and stoichiometric indicators, cannot well predict the decomposition characteristics of fine roots under flooded conditions. In addition, contrary to the third hypothesis, WGR with more phenolics and a higher unsaturation

degree and aromaticity experienced nearly 15% more carbon loss than SGR after 180d of incubation (Fig. 7). That is, the higher bound phenolics in WGR than in SGR could not explain the faster organic carbon decomposition in WGR than in SGR. Instead, the high carbon content in the free fraction in WGR compared to that in SGR (F_{DcMe} ; Fig. 2) was likely the major reason. It conformed to the previous conceptual model, which suggested that free organic carbon and nonstructural compounds controlled the amount of C lost from root litter in the early and middle decomposition phase (Soong et al., 2015; Shipley and Tardif, 2021). Moreover, WGR exhibited consistently higher decomposability of organic carbon than SGR in anoxic conditions, approaching the carbon decomposition potential of SGR in aerobic conditions (Fig. 7). The result suggested that, although flooding provided an anoxic condition that slowed down root decomposition, the adaptive strategy of developing more nonstructural labile tissue components in WGR in regard to flooding would accelerate root decomposition in order to counteract the effect of an anoxic condition.

5 Conclusions

This study developed and applied the complementary targeted and nontargeted analyses of sequentially extracted fractions of fine roots to determine the molecular-level fineroot carbon traits. The sequential chemical extraction allowed the analyses of both nonstructural and structural (e.g., biopolymers) root carbon components, and the combination of GC-MS and FT-ICR MS analyses successfully provided complementary quantitative and qualitative compositional information on organic compounds. This combined method was applied to further reveal the distinct differences in molecular-level carbon traits between SGR and WGR. The targeted and nontargeted analyses were sufficiently sensitive to show that WGR were enriched in more phenolics with a higher unsaturation degree and aromaticity and a more nonstructural composition to help plants cope with flooding stress than SGR. Also, we highlight the fact that, although flooding provided an anoxic condition that slowed down root decomposition in the short-term, the adaptive strategy of developing more nonstructural labile tissue components in WGR in regard to flooding would accelerate short-term root decomposition in order to counteract the effect of anoxic conditions. Nevertheless, the long-term effects of chemical adaptation under flooded environments on root degradation and carbon biogeochemical processes still require further evaluation. Together, we incorporated the molecular-level carbon traits of fine roots into trait-based ecology, which could greatly expand the root trait pools and thereby improve our understanding of plant functioning and biogeochemical progress in response to global environmental change.

Data availability. The data that support the findings of this study are available on Figshare (https://doi.org/10.6084/m9.figshare. 23726082, Wang, 2024). Correspondence and requests for materials should be addressed to Junjian Wang.

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