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A pulse of simulated root exudation alters the composition and temporal dynamics of microbial metabolites in its immediate vicinity

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ABSTRACT

Root exudation increases the concentration of readily available carbon (C) compounds in its immediate environment. This creates 'hotspots' of microbial activity characterized by accelerated soil organic matter turnover with direct implications for nutrient availability for plants. However, our knowledge of the microbial metabolic processes occurring in the immediate vicinity of roots during and after a root exudation event is still limited.

Using reverse microdialysis, we simulated root exudation by releasing a¹³C-labelled mix of low-molecularweight organic C compounds at mm-sized locations in undisturbed soil. Combined with stable isotope tracing, we investigated the fine-scale temporal and spatial response of microbial metabolism, soil chemistry, and traced microbial respiration and uptake of exuded compounds.

Our results show that a 9-h simulated root exudation pulse leads to i) a large local respiration event and ii) alteration of the temporal dynamics of soil metabolites over the following 12 day at the exudation spot. Notably, we observed a threefold increase in ammonium concentrations at 12 h and increased nitrate concentrations five days after the pulse. Moreover, various short-chain fatty acids (acetate, propionate, formate) increased over the following days, indicating altered microbial metabolic pathways and activity. Phospholipid and neutral lipid fatty acids (PLFAs, NLFAs) of all major microbial groups were significantly ¹³C-enriched within a 5 mm radius around the microdialysis probes, but not beyond. The highest relative ¹³C enrichment was observed in fungal NLFAs, indicating that a significant proportion of the exuded compounds had been incorporated into fungal storage compounds.

Our findings indicate that the punctual release of low-molecular-weight organic C compounds into intact soil significantly changes microbial metabolism and activity in its immediate surroundings, enhancing mineralization of native organic nitrogen. This highlights the versatility of microbial metabolic pathways in response to rapidly changing C availability and their effectiveness in increasing nutrient availability near plant roots.

1. Introduction

Roots release a variety of C-rich compounds into their immediate environment (Jones et al., 2009; Canarini et al., 2019). These compounds consist of high-molecular weight compounds such as mucilage and proteins, and low-molecular-weight compounds such as amino acids, organic acids, and sugars (Badri and Vivanco, 2009; Vives-Peris et al., 2020). The latter are predominantly released at the

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root tips, either through passive diffusion or active exudation (Badri and Vivanco, 2009; Jones et al., 2009; Canarini et al., 2019). Consequently, due to their constant movement through the soil, tips of growing roots create distinct 'exudation hotspots' of transiently high concentration of low-molecular-weight C compounds (Vogel et al., 2014).

Such pulses of easily available C are thought to stimulate the activity of previously partly starving and thus inactive microbes in the surrounding of the pulse (De Nobili et al., 2001; Mondini et al., 2006; Kuzyakov et al., 2015; Vetterlein et al., 2020). This process is marked by an immediate respiration of the assimilated C (Butler et al., 2004; Kaštovská and Šantrůčková, 2007), followed by an enhanced respiratory activity driven by increased microbial biomass turnover (Blagodatskaya and Kuzyakov, 2008), which is eventually followed by accelerated SOM decomposition - a phenomenon called the rhizosphere priming effect (Kuzyakov, 2002, 2010; Dijkstra et al., 2013; Haichar et al., 2014; Huo et al., 2017). While priming effects are observed in the majority of investigated cases, instances of no or 'negative' priming (i.e. retardation of SOM decomposition) have been reported (Kuzyakov, 2002; Dijkstra et al., 2013; Luo et al., 2016). These findings underscore the complexity of the rhizosphere C-cycling and variability of microbial interactions.

As a result of this accelerated microbial activity, a variety of microbial metabolites is released into the soil solution. Pulses of root exudation and their consequences for microbial activity are referred to as 'hot moments' in soil, implying that these changes in the process rates and metabolite levels last only for a short duration (Kuzyakov et al., 2015). The amount and composition of microbial metabolites released likely reflect the microbial processes occurring in response to the root exudation event but have not been previously measured at the hotspot. Moreover, the actual timeframe of accelerated or altered process rates at hotspots is uncertain and could range from minutes to days (Kuzyakov et al., 2015; Vetterlein et al., 2020). By studying the dynamics of soil metabolites that emerge in response to root exudation, valuable insights can be gained into the metabolic pathways involved in the degradation and transformation of exudate compounds, as well as the timeframe and persistence of the rhizosphere priming effect.

A substantial fraction of the C exuded by roots is incorporated into the rhizosphere microbial biomass (Butler et al., 2004; Denef et al., 2007). It was observed that exudates of young beech trees where preferentially utilized by gram-negative bacteria and fungi (Esperschütz et al., 2009), while gram-positive bacteria appeared to have little reliance on rhizodeposits but perhaps benefitted from later priming effects (Butler et al., 2003; Esperschütz et al., 2009; Balasooriya et al., 2013). Conversely, Cesarz et al. (2013) reported that beech roots did not affect bacterial biomass, but rather led to an increase in the fungi-to-bacteria ratio. This highlights how root exudates may be preferentially utilized by certain microbial groups and shape the microbial community composition (Zhalnina et al., 2018; Seitz et al., 2022), though the specific outcomes remain uncertain.

Researchers have employed various approaches, e.g. rhizoboxes (Wenzel et al., 2001; Chaignon et al., 2002; zu Schweinsberg-Mickan et al., 2010; Spohn et al., 2013) or planar optodes (Santner et al., 2015; Li et al., 2019), to study a range of root and rhizosphere processes, including spatial distribution of microbes, enzymes and chemical compounds. A common drawback of these tool is the limited capacity of time-points, spots, and analytes they can assess, and their struggle with the heterogeneous soil structure. Moreover, such *in situ* studies based on root exudates present their own unique challenges, as the composition and concentration of root exudates cannot be controlled, and the presence of root respiration further complicates the analysis.

Thus, various studies have added powders or solutions containing compounds found in exudates to sieved soil in an attempt to explore e.g. the mechanisms behind rhizosphere priming effects (Basiliko et al., 2012; Nottingham et al., 2012; Koranda et al., 2013; Wild et al., 2014; Girkin et al., 2018; Jilling et al., 2021) or changes in the microbial community (Shi et al., 2011; Mau et al., 2015; Papp et al., 2020). In contrast to root exudation which creates localized high C concentrations

with strong concentration gradients (Kuzyakov and Razavi, 2019; Vetterlein et al., 2020), the substrate addition results in uniform distribution of moderately enhanced C availability throughout the whole soil volume, which may affect microbial responses to substrate input considerably. In addition, the intricate mechanisms underlying the rhizosphere priming effect critically depend on the physico-chemical and biotic micro-environment of the root exudation hotspot, which is likely disintegrated by the abovementioned approaches.

Microdialysis allows to release artificial root exudates at small spots into undisturbed soil while at the same time collecting metabolites from the surrounding soil solution. In its classic application, metabolites are collected from the soil solution via a diffusion gradient across a small semi-permeable membrane inserted into the soil (Inselsbacher et al., 2011; Oyewole et al., 2014). By reversing the concentration gradient, i. e. by pumping high levels of low-molecular-weight compounds along the inside of the membrane, compounds can be released via passive diffusion at distinct soil spots (Buckley et al., 2022; König et al., 2022). This way it is possible to create a hotspot of high C concentration, akin to passive root exudation. Compared to Rhizon samplers, which also allow to mimic root exudation by releasing a solution (Kuzyakov et al., 2007; Keiluweit et al., 2015; Baumert et al., 2018), microdialysis has the benefit of not generating a mass flow into the soil by relying exclusively on diffusion. Moreover, the 'reverse' microdialysis approach offers the unique advantage of combining the release of organic compounds with the simultaneous collection of (other) soil metabolites (as by 'classic' microdialysis) at high temporal resolution.

The aim of this study was the detailed analysis of the fine-scale temporal dynamics of microbial activity and soil chemistry in response to a simulated root exudation pulse right at the exudation spot in intact soil. Specifically, we addressed the following questions:

- (1) Does a 'root exudation' pulse alter the dynamics of soil metabolites at the exudation hotspot?
- (2) What is the timescale and pattern of substrate- and SOM-derived microbial respiration following a simulated root exudation pulse?
- (3) Which microbial groups are involved in the metabolism of 'root exudates' in the days following the pulse?

To achieve this, we used reverse microdialysis to release a mixture of 13 C-labelled sugars (fructose, glucose) and organic acids (acetate, succinate) into undisturbed soil cores over a period of 9 h while simultaneously collecting metabolites from the soil solution for 12 days. Additionally, we assessed substrate-induced respiration and SOM mineralization, and the uptake of 13 C-labelled substrate into different microbial groups. We observed a sequence of microbial processes over time and their limitation in space around the artificial root, marked by long-lasting effects of simulated root exudation on soil biogeochemistry and microbial activity.

2. Material and methods

2.1. Soil sampling

Soil was sampled in a managed mature beech (*Fagus sylvatica*) forest (average tree age around 90 yrs) in Klausen-Leopoldsdorf, Austria (48°07′ N, 16° 03′ E, 510 m asl). The soil is a Dystric Cambisol (over flysch) with a pH of 4.16 (CaCl₂) and an organic C and total N content of 5.01% and 0.38%, respectively (Kitzler et al., 2006; Brandstätter et al., 2013; König et al., 2022). We sampled soil on June 6th⁻ 2018 at five locations along a horizontal transect at a distance of 60 cm each. After removing the litter layer, we took triplicate soil cores (10 cm diameter × 6 cm depth) right next to each other from the uppermost soil at each of the five locations. From each triplicate, one soil core was sieved, and the gravimetric soil water content was determined, which ranged between 44 and 53%. Per site, one of the two remaining intact soil cores was used to set up the mesocosms for the experiment (the third soil core served as

a backup).

2.2. Experiment design and microdialysis setup

Paired mesocosms (one for pulse simulation and one for control) were set up for each of the five soil sampling sites by coring two smaller cores (2.8 cm diameter \times 3 cm height) from one larger soil core. Intact mesocosm cores were carefully placed in modified polypropylene centrifuge tubes (Greiner Bio-One, 50 ml), which were cut off 4 cm below their screw caps and turned upside-down, so they stood in their closed screw caps (opening screw caps allowed easy collection of the soil at the end of the experiment). One of the two paired mesocosm acted as a control (n = 5), receiving only water.

In order to maximize the soil volume affected by microdialysis within each undisturbed soil core, two microdialysis probes (CMA 20, 20 kDa molecular weight cut-off, 1 cm membrane length, 0.5 mm outer diameter: CMA Microdialysis AB, Kista, Sweeden) were inserted into each mesocosm soil core. To avoid damaging the microdialysis probes, holes were pre-made with steel needles (0.5 mm diameter) to help guide the probes into the soil, which was then gently compressed around it. To ensure that only one sample per hour per mesocosm was collected, the tubes of the two microdialysis probes were joined with a T-connector (Micro T, CMA Microdialysis AB, Kista, Sweden) (Fig. S1). Microdialysis probes were perfused with ultrapure water (Milli-Q) containing low concentrations of KCl (3.848 μ M) and MgSO4 (1.595 μ M) (from now on called 'background solution') at a perfusion flow rate of 2.5 μ l min⁻¹ per microdialysis probe using syringe pumps (CMA 4004, CMA Microdialysis AB, Solna, Sweden). The purpose of adding KCl and MgSO4 to the perfusate was to prevent the formation of an osmotic gradient between the perfusate and the soil solution (Demand et al., 2017; Warren, 2018). Samples (dialysates) were collected in refrigerated fraction collectors (CM4 470, CMA Microdialysis AB, Solna, Sweden) and subsequently stored at -20 °C until analysis.

During an initial 3-h period, all microdialysis membranes were perfused with the background solution. This was done to facilitate the formation of water bridges within the soil and to analyse initial concentrations of soil compounds collected in the dialysates. In the following 9 h, we implemented a pulse of labile substrate using reverse microdialysis to mimic root exudation, similar to the diurnal carbon release dynamics observed in natural hotspots with heightened exudation during daylight hours (Murray et al., 2004; Cardon and Gage, 2006). A labile substrate mixture consisting of ¹³C-labelled acetate, succinate, fructose and glucose (each 125 µmol C l⁻¹, 98 at% ¹³C, uniformly labelled, Sigma-Aldrich) was added to the background solutions of one mesocosms of each pair (n = 5). These compounds were selected based on their prevalence in natural root exudates of beech trees (Smith, 1976) and to represent a range of chemical structures (sugar, organic acid) commonly found in root exudates (Vives-Peris et al., 2020). We deliberately refrained from including nitrogen-containing compounds, despite acknowledging their significance within root exudate (Drake et al., 2013; Edwards et al., 2018; Chari and Taylor, 2022), in order to focus specifically on the impact of carbon alone - a key element in the root exudate-microbe-soil interaction.

After this pulse, the perfusate was switched back to pure background solution. Control mesocosms received background solution throughout the whole experimental as perfusate. For the first 3 days (72 h), microdialysis was kept running constantly in all mesocosms and dialysates were collected hourly. Subsequently, the pumps were turned off and only switched on for 4 h on day 5, 7, and 12 to collect additional dialysate samples. Throughout the experiment, we monitored the water content by weighing the mesocosms and adjusting it, if necessary, by pipetting Milli-Q onto the surface.

2.3. Microbial responses to simulated root exudation

2.3.1. CO₂ analysis – gas sampling and estimation of respiration rates

Mesocosms were placed in airtight jars fitted with septa for taking gas samples (see Fig. S1). The microdialysis tubing was inserted through the septa to allow simultaneous dialysate collection. We took headspace gas samples at multiple time points during the experiment: on day 1 (hours 2, 6, 7, 8, 9, 10, 13), on days 2 and 3 (twice), and once on days 5, 7 and 12. The ¹²C–CO₂ and ¹³C–CO₂ concentration of gas samples were measured by a headspace gas sampler (GasBench II, Thermo Fisher Scientific, Bremen, Germany) couple to a isotope ratio mass spectrometry (Delta V Advantage, Thermo Electron, Bremen, Germany). When taking gas samples, the mesocosm jars (105 ml volume) were closed airtight, the first gas sample (15 ml) was taken and the removed air was replaced with artificial air (200 ppm CO₂) to avoid sub-atmospheric pressure. The second gas sample was taken after 60 min incubation time and the mesocosm jars were opened. The CO₂ concentration and ¹³C signature of the second sample was corrected for the added artificial air. The total respiration rate and its atom percent ¹³C (at% ¹³C) signature were calculated from ¹²C- and ¹³C-CO₂ concentration differences between the initial gas sample and the second, corrected, gas sample. The percentage and total amount of respiration originating from the added ¹³C-labelled substrate and from the native SOM were calculated using a two-pool mixing model with the ¹³C signature of the added substrate (98 at%¹³C) and natural abundance of ¹³C in the respiration of the control mesocosms (1.08 at% ¹³C) as end members. For a detailed outline of the calculations see the supplementary materials.

It should be noted that the mesocosms that were to receive the labile substrate pulse exhibited a lower average respiration per gram of mesocosm-soil prior to the substrate pulse and lower SOM-derived respiration persisted throughout the first part of the experiment (Fig. S2). We attribute this to an uneven distribution of stones and organic matter across the mesocosms observed during soil harvest, which we could not control for before the experiment as we did not sieve the soil. To account for this initial variability, we calculated a normalized SOM-derived respiration, in which all respiration measurements were put in relation to the initial respiration (% of initial respiration, Fig. 1). This allows a more accurate comparison of the effect of the labile substrate addition on respiration (Fig. 1). For the original data in nmol $CO_2 g^{-1}$ soil hour⁻¹, we refer the reader to Supplementary Fig. S2.

Additionally, we estimated the 'background' SOM-derived respiration in the immediate surrounding of each microdialysis probe, with the assumption that SOM-derived respiration was not locally affected by the substrate input. We calculated the fraction of the SOM-derived respiration obtained per soil core around the microdialysis probe as a cylindrical volume as:

$$R_{background} = R_{SOM} \times DW_{mesocosm} \times \frac{V_{cylinder}}{V_{mesocosm}}$$

where $R_{background}$ is the SOM-derived respiration (nmol CO₂ h^{-1} $V_{cylinder}^{-1}$) within $V_{cylinder}$, the cylindrical soil volume around the microdialysis probe, R_{SOM} is the SOM-derived respiration of the mesocosm (nmol CO₂ h^{-1} g^{-1}), DW_{mesocosm} is the soil dry weight of the mesocosm (g), and V_{mesocosm} is the volume of the mesocosm (mm³).

We selected a 1-mm radius for our calculations as we aimed to estimate substrate- and SOM-derived respiration over a short 9-h period during which substrate was released. This falls at the lower end of range observed in several studies, which have reported plant root exudates occurring between 0.2 and 2 mm away from the surface of the roots (Falchini et al., 2003; Herman et al., 2006; zu Schweinsberg-Mickan et al., 2010). Occasionally, however, studies have found exudates to disperse up to 5–12 mm (Sauer et al., 2006; Dessureault-Rompré et al., 2007; Finzi et al., 2015).



Fig. 1. Temporal dynamics of normalized SOM-derived respiration rate (percentage of initial respiration) and substrate-derived respiration rate. The time of the 9-h long pulse from labile substrate (4–12 h) is highlighted as an orange background and the long-term response (5–12 days) is highlighted as a grey background. The short-term response is depicted in hours and the long-term effects in days and are separated by an x-axis break at hour 72. Top: The normalized SOM-derived respiration rate normalized by dividing by the initial respiration rate in response to the exudation pulse (orange) compared to a water control (blue). Bottom: The respired labile substrate (nmol $CO_2 g^{-1} h^{-1}$) is shown in red. Depicted are the means and standard error of the mean (n = 5) with asterisks indicating significant differences in response to exudation pulse.

2.3.2. Dialysate analysis (soil chemistry & metabolites)

We measured organic and inorganic anions and sugars in the dialysates using HPLC (Dionex ICS 5000+, Thermo Fisher, Germany). To optimize the chromatographic analyses, we pooled the consecutive hourly dialysates and combined 2, 3 or 4 dialysates for periods 1–20 h, 21–29 h and remaining dialysates, respectively.

Anions were measured on a Dionex IonPac AS11-HC (2×250 mm) column with a Dionex IonPac AG11-HC (2×50 mm) guard column at a constant flow rate of 0.38 ml min⁻¹ with a KOH solvent. We analysed acetate, propionate, formate, butyrate, chloride, nitrate, succinate, malate, sulfate, oxalate, phosphate, and citrate. The HPLC run started at 1 mM KOH (10 min), then increased to 15 mM (1 mM min⁻¹), then to 60 mM (5 mM/min) and kept at this concentration for 5 min and finally the run was ended with 7 min at 1 mM. Sugars (glucose, fructose, galactose, sucrose) were measured on a Thermo CarboPac PA20 (0.4 × 150 mm) column with a Thermo CarboPac PA20G (0.4 × 35 mm) guard column at a constant flow rate of 8 µl min⁻¹ with a KOH solvent. We started the run with 10 min of 10 mM KOH, increased to 50 mM for 5 min, increased to 200 mM for 5 min and ended the run with 5 min at 20 mM. The ammonium concentration in dialysates was determined colorimetrically (Kandeler and Gerber, 1988; Hood-Nowotny et al., 2010).

2.3.3. PLFA and NLFA analyses

At the end of the experiment (day 12), we harvested the soil to investigate the ¹³C enrichment of different microbial groups in soils that received labile ¹³C-labelled substrate. We harvested soil in the surrounding (<5 mm) and distant (>5 mm, Fig. S3) radius of the microdialysis probes by coring the soil with a rectangular tube (1 cm \times 1.5 cm) with the two microdialysis probes in the centre. Afterwards the soil was homogenised by mixing and stones were removed with a tweezer. To determine which microbial groups took up the substrate, we extracted the phospholipid and neutral lipid fatty acids (PLFAs, NLFAs) and used them as biomarkers for different microbial groups and fungal storage, respectively.

Lipids were extracted from lyophilized soils with a mixture of

chloroform, methanol, and citrate buffer (v/v/v = 1:2:0.8; (Bligh and Dyer, 1959; Frostegård et al., 1991). Neutral lipids and phospholipids were consecutively eluted on silica solid phase extraction columns with chloroform and methanol, with an acetone washing step in-between. The extracts were subsequently derivatized via mild alkaline methanolysis. The resulting fatty acid methyl esters were analysed on a gas chromatograph (Trace GC Ultra, Thermo Scientific, Germany) coupled to a mass spectrometer (ISO, Thermo Scientific, Germany) for fatty acid identification and quantification, and on a GC-Ultra (Thermo Fisher Scientific, Milan, Italy) coupled to an isotope ratio MS (Finnigan Delta-V, Thermo Fisher Scientific, Bremen, Germany) via a GC IsoLink (Thermo Fisher Scientific, Bremen, Germany) for determination of isotopic ¹³C/¹²C ratios. We used nonadecanoic acid methyl ester as an internal standard for quantification, and bacterial and fungal fatty acid methyl esters (BAME CP mix, Supelco; 37 Component FAME mix, Supelco) as qualitative external standards.

Fatty acids of the phospholipid and neutral lipid fractions were assigned to the same phylogenetic groups. Unless otherwise noted, fatty acids were assigned according to (Willers et al., 2015). We assigned the unsaturated fatty acids 14:0, 15:0, 16:0, 17:0 and 18:0 as general microbial markers (data not shown), the monoand cyclopropyl-unsaturated fatty acids 16:1w5 (Phillips et al., 2002), 16:107, 17:107, cy17:0 and cy19:0 as markers for gram-negative bacteria, the terminally branched fatty acids i15:0, a15:0, i16:0, i17:0 and a17:0 for gram-positive bacteria (excluding Actinobacteria) (Joergensen and Wichern, 2008), the methyl-branched fatty acids 10Me18:0 and 10Me19:0 for Actinobacteria, and the fatty acids 18:109c, 18:109t and 18:2 ω 6,9 for fungi. However, we must note that although the 18:1 ω 9t is reported as specific for fungi (Balser et al., 2005) it can also show patterns similar to gram-negative bacteria biomarkers.

We calculated isotopic enrichment in individual PLFAs and NLFAs as the difference in relative ¹³C-enrichment of labelled samples to unlabelled control as at% ¹³C excess (Kaiser et al., 2015). The relative ¹³C content (μg ¹³C excess) of individual PLFAs and NLFAs was calculated as relative ¹³C-enrichment weighted by the mean abundance (in μg C in

FAs g^{-1} dry soil).

2.4. Statistical analysis

All statistical analysis were performed using R (R Core Team, 2022) and all graphs were plotted using ggplot2 (Wickham, 2016). We removed outliers that had values that were higher or lower three times the interquartile range. We tested the normality of the data using Shapiro-Wilk test, and homogeneity of variances using the Levene test.

We tested significant differences of compound concentrations per time point using Mann-Whitney *U* test (Table 1) since the data were neither normal distribution nor showed homogeneity of variances, regardless of data transformation. We log-transformed the respiration data (SOM-derived respiration and normalized SOM-derived respiration) and tested significant differences from control for each time point with Student's two sample *t*-test. The μ g C and at% ¹³C values of PLFA and NLFA biomarker groups were tested for differences between amended mesocosms receiving an exudation pulse compared to the water controls using a Student's two sample *t*-test (Table 2). Moreover, the μ g C, μ g ¹³C excess and at% ¹³C excess of fatty acid biomarker groups were tested for differences between PLFAs and NLFAs using a Student's two sample *t*-test or Mann-Whitney *U* test (Table 3).

3. Results

3.1. Transfer rates and retrieval rates of added labile substrate

The transfer rate of compounds i.e., the percentage of compounds in the perfusate that were actually released into the soil (König et al., 2022), were 31.84% (±0.87 se) for acetate, 15.06% (±1.08) for succinate, 9.15% (\pm 1.21) for glucose and 5.65% (\pm 1.7) for fructose, averaged over the 9-h pulse (Fig. 2: acetate, Fig. S4: succinate, glucose, fructose). This transfer rates correspond to an average $113.43 (\pm 4.89 \text{ se})$ nmol C for acetate, 53.1 (±8.31) nmol C of succinate, 27.92 (±7.41) nmol C of glucose and 18.23 (±5.24) nmol C of fructose, or in sum 212.68 (\pm 22.17) nmol C per mesocosm. The 'retrieval' rates i.e., the percentage of initially exuded compounds, which were retrieved back in the first 2 h after the pulse after switching back to background solution were 6.32% (±0.28 se) for acetate, 16.51% (±4.26) for succinate, 27.67% (±2.99) for glucose and 47.62% (±10.78) for fructose. This retrieval rates correspond to an average 7.14 (± 0.33) nmol C acetate, 7.68 (±1.15) nmol C succinate, 9.38 (±0.36) nmol C glucose, and 9.63 (± 0.88) nmol C fructose, or in sum 34.53 (± 2.30) nmol C per mesocosm. Note that for sugars in particular, the transfer rates dropped to nearly zero percent after 4 h with simultaneously high retrieval rates (Fig. S4), indicating a saturation in sugars around the microdialysis probe after not being removed quickly enough by biological or physical processes.

3.2. Temporal pattern of several soil metabolites is altered after a pulse of labile C compounds over the following 2-12 days

Overall, we found that several soil metabolites, such as acetate, propionate, formate and ammonium, displayed distinct temporal patterns in the hours following the labile substrate pulse, but we also observed longer-term changes in acetate, formate, nitrate and sulfate concentrations several days later (Fig. 2, Table 1). The concentrations of certain small chain organic acids peaked about 24 h after the labile C input (Fig. 2, Table 1). Specifically, acetate concentrations increased significantly 9 h after the end of the input pulse (hour 20) and remained elevated for 34 h (i.e., until hour 53) (Fig. 2, Table 1). Similarly, in the same period, ammonium and propionate concentrations significantly increased compared to the control (hours 26–45 and hours 29–45, respectively). This increase in organic acids and ammonium was accompanied by a significant increase of sulfate during hours 29–33. Following the organic acid peak (acetate, ammonium, propionate) we detected a trend towards higher formate and sulfate concentration. We

	2	exudá	ttion puls	ë																				day	
hour		4	9	8	10	12	14	16	18	20	23	26	29	33	37	41	45	49	53	57	61	65	69	ß	7
acetate	I	*	+ ^{4 4}	**+	+** ↓	+ ^{4 4}	^{**} ↑	I	*	**+	*	*	^{**} ↑	+** ↓	*	*1	*	+** ↓	*	I	*	*	1 ^{**}	*	*↓
succinate	I	*↓	*	*↓	**↓	**↓	I			I		I	I	*↓	I	I	I	I	I			I	I	I	I
fructose		*	**†	*↓	**↓	**†	*↓																		
glucose		*	**†	*↓	**↓	**†	*↓																		
propionate	I	I	I	I	I	I	*	*→	*→	I	I	I	**↓	**↓	**↓	*↓	*↓	I	I	I	I	I	I	I	I
formate	I	I	* →	I	I	I	**	* *→	* *	*↓	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I
ammonium	I	I	I	I	I	I	I	I	*→	*→	I	+** ↓	+×× ↓	I	+×× ↓	+×+ ↓	*↓	I	I	I	I	I	I	I	I
nitrate	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I
sulfate	I	I	I	I	I	I	I	I	I	I	I	I	**↓	*	I	I	*	*↓	I	I	I	I	I	I	I
citrate	I	I	I	I	I	I	I	I		I	I					I						I	I	I	I
oxalate	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	*↓	I	I	I	I	I	I	I	I	I

[able]

Table 2

Response of phospholipid fatty acids (PLFAs) and neutral lipid fatty acids (NLFAs) in the proximity (<5 mm) of the microdialysis probe, 12 days after the simulated root exudation event. The differences of μ g C and at% ¹³C values (PLFA, NLFA) between exudation treatment (n = 5) and water control (n = 5) were tested. Depicted are p-values and asterisks for significant differences (p < 0.05).

	biomarker	fungi			bacteria		
data		18:2ω6,9	18:1ω9c	18:1ω9t	gram -	gram +	actino
µg C	PLFA NI FA	n.s.	n.s.	n.s.	n.s.	0.015 *	n.s.
at% ¹³ C	PLFA NLFA	n.s. 0.004 **	0.009 ** 0.010 *	0.018 * 0.015 *	0.029 * 0.040 *	0.005 ** 0.010 *	0.019 * 0.019 *

Student's *t*-test, p-value: >0.05 n.s., <0.05 *, <0.01 **, <0.001 ***.

Table 3

Differences between phospholipid fatty acids (PLFAs) and neutral lipid fatty acids (NLFAs) in the proximity (<5 mm) of the microdialysis probe, 12 days after the simulated root exudation event (n = 5). The differences between PLFA and NLFA for μ g ¹³C excess, μ g C and at% excess were tested. Depicted are p-values and asterisks for significant differences (p < 0.05).

	fungi			bacteria		
data	18:2ω6,9	18:1ω9c	18:1ω9t	gram -	gram +	actino
µg C	n.s.	0.0 ***	0.0 ***	0.0 ***	0.0 ***	0.006 **
µg ¹³ C excess at% ¹³ C excess	n.s. n.s.	n.s. 0.021 *	0.032 * n.s.	n.s. n.s.	0.016 * n.s.	n.s. n.s.

Student's *t*-test (Mann-Whitney *U* test: $\mu g^{13}C$ excess), p-value: >0.05 n.s., <0.05 *, <0.01 **, <0.01 ***.

observed a slight increase in nitrate concentrations in the dialysates of all mesocosms over the first three days, but no significant differences between the ones that received an input pulse and the controls.

In the days 5–12, in a longer-term response to the labile substrate input, acetate levels again rose significantly on the third day and remained elevated until day 12. At the end of the experiment, the concentrations of acetate, formate, nitrate and sulfate were significantly higher in the dialysates of the samples that had received the input pulse compared to the control samples (Fig. 2). All other compounds (succinate, oxalate, citrate, glucose, and fructose) were detected in negligible amounts and were not clearly affected by the labile substrate input (Fig. S4). Butyrate, malate, phosphate, chloride, galactose and sucrose were not detected in our samples.

3.3. Rapid microbial respiration of labile substrate during the pulse precedes delayed changes in SOM respiration

Our measurements of ¹³C concentration in the CO₂ produced during the incubation showed that the added mixture of ¹³C-labelled C compounds was readily taken up and rapidly respired (Fig. 1). The substratederived respiration increased sharply 3 h after the start of the labile substrate pulse and remained elevated for the duration of the 9-h long substrate pulse. Out of the total substrate added, the microbes respired 34%, of which 45% was respired during the initial 2–3 h of the pulse (data published in: König et al., 2022). In the hours following the pulse the substrate-derived respiration levelled off (Fig. 1). In contrast, the SOM-derived respiration was unaffected initially (Fig. 1). On day 3, however, soils that received a labile substrate pulse respired significantly more SOM than the control soils (Welch's *t*-test, t(6) = -2.52, p =0.045) when compared in % of their respective initial SOM respiration (Fig. 1). We observed a similar, albeit non-significant, trend of higher normalized SOM-respiration during later time points (day 5, 7, 12).

3.4. Estimated increase in CO_2 respiration at the exudation hotspot

In order to obtain a rough estimate on the magnitude of increase of



Fig. 2. Temporal dynamics of six soil metabolites (μ M; acetate, propionate, formate, ammonium, nitrate, sulfate) in response to a 9-h long substrate pulse (orange) compared to a water control (blue). The time of the exudation pulse (4–12 h) is highlighted as an orange background and the long-term response (5–12 days) is highlighted as a grey background. The short-term response is depicted in hours and the long-term effects in days and are separated by an x-axis break at hour 72. Shown are the mean and standard error of the mean (n = 5) of concentrations in collected dialysate. Acetate was part of the labile substrate pulse and its initial concentration in perfusate is depicted as a black horizontal line. The difference between the black horizontal line and the acetate measurements in dialysates illustrate the net transfer into the soil. Asterisks indicate that the response to labile substrate significantly differs from the water control. Detailed statistics are shown in Table 1. For net transfers and temporal dynamics of citrate and oxalate (n.s.) see Fig. S4.

6

Respiration at exudation spot

(nmol CO₂ h⁻¹) F

microbial respiration right at the exudation hotspot, we calculated SOMderived and substrate-derived respiration for a volume within a 1 mm radius around the microdialysis probes, which we assume to be directly affected by the substrate input. We calculated the fraction of the SOMderived respiration obtained per soil core that occurred in a cylindrical volume of 1 mm radius and 10 mm height, with the aim to estimate the 'background' SOM-derived respiration in the immediate surrounding of each microdialysis probe. We then calculated the cumulative respiration within the hypothetical 1 mm diameter, consisting of the calculated SOM-derived respiration and the measured ¹³C-substratederived respiration (Fig. 3), which we assumed to occur to 100% in the surrounding of the probes (Fig. 4). Based on these assumptions, during the second half of the pulse (hour 7-10), an average additional 10.15 nmol CO₂ (\pm 1.7 se, n = 5) were respired from the added substrate in the immediate surrounding of each microdialysis probe, approximately doubling the respiration within this volume relative to the 'background' SOM-derived respiration (9.36 \pm 1.46 se nmol CO₂) (Fig. 3). These calculations rely on our assumption that root exudates would not go beyond a 1 mm radius within 9 h, and that there is no local increase in SOM-derived respiration triggered by the released C compounds. If we would vary the assumed range of root exudates to 0.5, 2 or 5 mm radius within which 2.34, 37.45 and 234.06 nmol CO₂ would be respired from SOM, substrate-derived respiration would increase CO₂ production in this volume by 434, 27 and 4 %, respectively. If there was a local increase in SOM-derived respiration within the chosen radius (which we were unable to measure), we would underestimate the increase in respiration due to the added substrate, as our calculation only includes the substrate-derived respiration plus a volume-based aliquot of the total SOM-derived respiration of the core.

3.5. Phospholipid and neutral lipid fatty acids were ${}^{13}C$ enriched in vicinity of the exudation hotspot



substrate respiration

SOM-derived respiration



18:2 ω 6,9, were significantly ¹³C enriched in response to the labile substrate pulse within a 5 mm radius around the microdialysis membranes (at% ¹³C: Fig. S5, Table 2). There was no significant enrichment in ¹³C in biomarkers in the soil sampled further away i.e., between 5 and 10 mm around the membranes (Fig. S3). Within the 5 mm radius, the NLFA fungal biomarker 18:1 ω 9c showed the highest relative ¹³C enrichment compared to the natural abundance control and was also significantly higher compared to its corresponding PLFA biomarker (at% ¹³C excess: Fig. 4). The at% ¹³C excess of the PLFAs and NLFAs from other microbial groups showed no significant differences (Table 3, Fig. 4).

When comparing absolute amounts of μg ¹³C excess of PLFAs and NLFAs, we found significantly more ¹³C in PLFA than in NLFA of 18:1 ω 9t (fungi), gram-positive bacteria, and gram-negative bacteria (not significant), while the fungal marker 18:1 ω 9c had a tendency (albeit not significant) for higher amounts of ¹³C excess in NLFAs compared to PLFAs (μg ¹³C excess: Fig. 4, Table 3). Since we did not find differences in the relative ¹³C enrichments (at% ¹³C excess) between PLFAs and NLFAs for 18:1 ω 9t, gram-negative and gram-positive bacteria, the observed differences in absolute ¹³C enrichments need to be attributed to the higher absolute amounts (μg C) of PLFAs compared to NLFAs (μg C: Fig. S5).

The labile substrate input significantly increased the microbial biomass (i.e., μ g C in PLFAs) of the gram-positive bacteria, but did not affect the biomass of any of the other groups. It also did not affect NLFAs of any microbial group (μ g C: Fig. S5). NLFA concentrations were lower compared to PLFA concentrations for all groups except for the fungal biomarker 18:2 ω 6,9.

4. Discussion

Our results show that a pulse of organic C compounds released via micro-diffusion at discrete locations in intact soil alters the temporal dynamics of soil metabolites right at these spots over the following hours and days. Interestingly, the concentration of several small-chain fatty acids increased in response to the C input, indicating changes in microbial metabolic pathways. This went along with a strong increase of ammonium availability, indicating microbial mineralization of soil organic N. Our measurements thus support the widely observed phenomenon that labile C compounds exuded by roots lead to enhanced nutrient availability in the rhizosphere (Zhu et al., 2014; Murphy et al., 2015).

4.1. Impact of labile C pulse on microbial activity

Soil microbes respired the released substrate with a 2-3 h delay, which might reflect a physiological lag phase, caused by the need to adapt their internal cellular machinery to the availability of a new substrate (Bertrand, 2019). Had the delayed respiration of the added substrate been due to a slow diffusion of the substrate, we would have seen a steady increase in substrate-derived respiration and no lag phase followed by the very high respiration rates observed (Fig. 1, bottom). It was hypothesized that soil microbes, whose activity is often constrained by C and energy limitations (Blagodatskaya and Kuzyakov, 2013), maintain a state of 'metabolic alertness' to quickly utilize infrequently arriving resources in an otherwise scarce environment (De Nobili et al., 2001; Mondini et al., 2006). The observed 2-3 h delay is consistent with our previous study in which reverse microdialysis was performed with various organic C compounds in different soils (König et al., 2022). There, we observed - after a quick initial transfer - a rapid decline of transfer rates of all compounds into the soil. This indicates (as transfer rates depend on the concentration gradient between the perfusate and the soil), that the released substances accumulated in the soil solution outside the membrane. Transfer rates abruptly increased 3 h later, implying a sudden uptake of the accumulated compounds by microbes after a time delay. These findings, along with the here observed delay in microbial respiration, collectively suggest that microbes in intact soil



Fig. 4. Enrichment in labile-substrate derived ¹³C in phospholipid fatty acids (PLFAs, yellow) and neutral lipid fatty acids (NLFAs, pink) in close proximity (<5 mm radius) to the microdialysis probe 12 days after a 9-h pulse of ¹³C-labelled substrate (n = 5). Due to the significant enrichment observed in all PLFA and NLFA biomarkers (at% ¹³C: Fig. 3S), except for 18:2 ω 6,9 PLFA, we have opted to exclusively present the ¹³C excess data here (μ g ¹³C excess, at% ¹³C excess). The fungi-specific fatty acid biomarkers 18:1 ω 9c, 18:1 ω 9t and 18:2 ω 6,9 are depicted ungrouped. The remaining fatty acids were grouped into gram-negative bacteria, grampositive bacteria (*excluding *Actinobacteria*) and *Actinobacteria*. Asterisks indicate significant differences (p < 0.05) between PLFA and NLFA for μ g ¹³C excess and at % ¹³C excess. Detailed statistics are shown in Table 2 and Table 3.

environments undergo a lag phase before taking up and respiring a new substrate.

Organic acids, in contrast to sugars, exhibited higher transfer rates (substrate release) during the pulse and lower back-retrieval rates afterwards, consistent with our previous study in which transfer and retrieval rates were negatively correlated (König et al., 2022). This suggests that sugars accumulated around the membrane after their initial release and were not effectively removed by biotic or abiotic processes like microbial uptake or adsorption to soil surfaces (Jones et al., 2003). Sugars, while widely regarded as ubiquitous substrates that undergo intensive recycling in microbial biomass (Apostel et al., 2015), appeared to be metabolised to a lesser extent by soil microbes compared to organic acids. The relatively quicker removal of organic acids from the membrane surrounding, as indicated by their higher transfer and lower back-retrieval rates, might indicate a more rapid adsorption to soil minerals compared to sugars. However, low-molecular-weight organic substances like glucose and acetate were reported to be consumed within a few hours (Jones et al., 2009), faster than their physico-chemical sorption to the soil (Fischer et al., 2010). To understand the underlying causes for the different behaviour of sugars and organic acids, future experiments should examine each compound class separately. Especially since glucose and acetate may enter different metabolic pathways, such as glycolysis and gluconeogenesis, respectively (Schink et al., 2022).

4.2. Rhizosphere priming effect and nitrogen mineralization

While the strong substrate respiration on the first day did not coincide with an accelerated SOM mineralization, we observed a significant increase in SOM-derived respiration compared to the control starting at day 3 (Fig. 1, top). This 'rhizosphere' priming effect may be attributed to a gradual activation and increase in microbial biomass in response to the added substrate (De Nobili et al., 2001), leading to an increased production of extracellular enzymes that boosted SOM decomposition (Kuzyakov, 2010; Dijkstra et al., 2013). This was further reflected in the up-to-threefold increase in ammonium concentrations within 24 h of substrate input. This indicates N mineralization from SOM and hence accelerated SOM decomposition, which was not yet matched by an increase in SOM-derived respiration (Fig. 1).

Priming effects have been reported to increase N mineralization (Dijkstra et al., 2009; Murphy et al., 2015). Adding C-rich substrate without providing additional N may have induced N-limitation and prompted the microbial community to mine for N (Craine et al., 2007; Fontaine et al., 2011; Brzostek et al., 2013). In a separate microdialysis experiment, sucrose addition was proposed to increase the N demand of the microbial community by alleviating C limitations (Buckley et al., 2022). Consequently, the delayed onset of increased SOM-derived respiration (priming effect) by three days could be partially ascribed to the lack of N in our exudate, which is recognized for its pivotal role in enzyme production (Drake et al., 2013) and, thus, the degradation of SOM. The sequence of events, ranging from substrate respiration to the subsequent increased ammonium concentrations (N-mining) and accelerated SOM-derived respiration (priming effect), can be attributed to the time, energy and N resources required for enzyme production (Liu et al., 2019).

Alternatively, the increase in ammonium concentrations could result from a liberation of previously clay-bound ammonium through organomineral interactions with the released organic acids (Keiluweit et al., 2015). We think, however, that this is an unlikely explanation, as the steep ammonium peak occurred only 24 h after the release of organic acids, at a time when both acetate and succinate were only present at much smaller than initial (acetate) or even negligible (succinate) concentrations (Fig. 2). From day 5 onwards, the increased nitrate concentration compared to the controls indicates microbial nitrification, probably triggered by the high amounts of ammonium. Besides nitrate, the elevated levels of acetate (days 3–12), formate (day 12), and sulfate (day 12; Fig. 2, Table 1), imply long lasting changes in soils that received labile substrate.

It should be noted that our experimental design placed limitations on the measurement sensitivity of a spatially constrained priming event. Specifically, the high background of SOM-derived respiration of the whole soil core may have masked the local increase of SOM-derived respiration within the small volume influenced around the two microdialysis membranes. However, this 294:1 vol ratio (assuming a 1 mm radius of influenced soil around the two membranes) implies that the actual priming effect at the exudation hotspot on the third day was likely considerably higher than our observation suggests.

4.3. Altered metabolic pathways in response to labile C

Simultaneous with the onset of the 24-h long ammonium peak, other organic and inorganic compounds increased: acetate concentrations increased 6 h prior, while propionate, sulfate, and formate concentrations increased 3, 3 and 15 h after the onset of the ammonium peak (Fig. 2), suggesting a complex interplay of metabolic pathways. The organic acids could be fermentation products, indicating potential anaerobic conditions. However, in high glucose environments, it has been shown that E. coli cultures favour fermentation over respiration. leading to the release of high concentrations of acetate, even when oxvgen is available (Luli and Strohl, 1990; Chacón et al., 2018; Mori et al., 2019). Remarkably, many bacteria employ these respiro-fermentative overflow metabolism and still attain high growth rates (Wortel et al., 2018). For thermo-dynamical reasons, heterotrophic microbes face a trade-off between growth rate and yield (Pfeiffer et al., 2001; Kreft and Bonhoeffer, 2005), which becomes particularly relevant under conditions of abundant resource or limited oxygen supply. Under such conditions the ATP production rate of respiration saturates rapidly and microbes can maximize growth rate at the expense of yield by switching at least partly to incomplete metabolic pathways. In E. coli, the growth-rate dependent acetate overflow was explained by fermentation being favoured at high growth rates due to its lower biosynthesis cost relative to the electron transport chain (Basan et al., 2015). Alternatively, the membrane real estate hypothesis proposes that at high growth rates, cells increase in size, reducing their available membrane area for respiratory proteins, prompting a shift towards less efficient but faster ATP production through fermentation (Szenk et al., 2017). Additionally, it was suggested that under nutrient limitations excess C may be routed to overflow respiration or excreted, to accommodate the decomposers stoichiometric requirements (Schimel and Weintraub, 2003; Manzoni and Porporato, 2009).

Interestingly, this would also be a viable strategy for microbes under aerobe conditions, given that microbes face a surplus of sugars (Lipson, 2015). Incomplete metabolic pathways, such as fermentation, allow a higher substrate flux per unit of time but go along with the release of metabolic intermediates such as acetate (Pfeiffer and Bonhoeffer, 2003). The release of high glucose levels at local exudation hotspots could thus have created the right conditions for this glucose-mediated aerobe acidogenesis to take place. Moreover, nutrient limitations induced by the absence of N in the exudate could have resulted in the redirection of surplus C towards excretion or potentially overflow respiration to meet the decomposers stoichiometric requirements (Schimel and Weintraub, 2003; Manzoni and Porporato, 2009). The reduction in the organic acid peaks, e.g., acetate starting at hour 33, signifies a transition from a period of rapid growth with organic acid production to a slower growth phase involving the assimilation of the excreted organic acids. This switch is likely driven by the depletion of the organic acid-producing carbon sources (e.g., sugars) (Wolfe, 2005). Collectively, our findings suggest altered metabolic pathways and microbial activity in the days following the substrate input. Going forward, it should be taken into account that high concentrations of organic acids, like acetate and formate measured in the rhizosphere solution of beach trees (Fagus sylvatica) (Fender et al., 2013), might originate from microbial excretion just as likely as from root exudates, highlighting the need for careful interpretation in future studies.

4.4. Oxygen depletion at the exudation hotspot

It is, however, also possible that microbial respiration in the immediate vicinity of the exudation spot caused a temporary depletion of oxygen (O₂) at the spot due to the additional respiration of the released substances: If we assume a pore space of approx. 50% of the soil volume (Voltolini et al., 2017), the 1 mm cylinder around each microdialysis probe will contain a pore volume of 15.71 mm³. We calculated that, based on oxygen saturation levels of 9.1 mg l^{-1} (20 °C) in water and a concentration of 20.95% O_2 in (atmospheric) air the total amount of available O2 within such a cylinder would be around 75.7 nmol O2 (assuming a 50% water saturation of the soil pores), with only 2.24 nmol O₂ being dissolved and thus immediately available in water-filled soil pores (Supplementary material). Comparing this potential O₂ availability with the estimated increased O2 demand for respiration (SOM-derived plus substrate-derived) of around 19.52 nmol CO₂ (nmol O₂, hour 7-10) during the 9-h long labile substrate pulse (Fig. 3) suggests that a temporary depletion of O₂ concentrations in the vicinity of the substrate input is possible. While it is unknown whether the release of high concentrations of organic compounds leads to temporary oxygen depletion around roots, the formation of anoxic hotspots has been shown in an artificial pore system caused by increased bacterial oxygen consumption at locations with high C availability (Borer et al., 2018). If temporary oxygen depletion occurs around root exudation hotspots it would have consequences for microbial metabolism. Microbes could, for instance, switch to an anaerobic metabolism, which would also go along with shorter, incomplete catabolic pathways that lead to a release of intermediates, such as acetate (Kreft et al., 2020). Moreover, when oxvgen concentrations are low, nitrate can serve as an alternative electron acceptor, to allow the oxidation of SOM to CO₂ (Nannipieri and Eldor, 2009; Song et al., 2023). In this case, nitrate is reduced to ammonium in a process called dissimilatory nitrate reduction (Takaya, 2002; Nannipieri and Eldor, 2009), which might also have contributed, to the increased ammonium concentrations we observed at around hour 26 - 45

4.5. Substrate utilization by microbial groups and fungal strategies

All microbial groups were significantly ¹³C enriched exclusively within a 5 mm radius of the exudation pulse (at% ¹³C: Fig. S5, Table 2). This suggests that the volume of influence of the exudation pulse, and thus the uptake of the ¹³C-labelled substrate, was limited to the immediate vicinity of the input. This is consistent with the observations that most methods find the rhizosphere to extend between 0.5 and 4 mm (Kuzyakov and Razavi, 2019). Notably, among the microbial groups investigated, only the biomass of gram-positive bacteria significantly increased in response to the labile substrate pulse (µg C: Fig. 4S and Table 2). This was surprising, as we would instead have expected a biomass increase in gram-negative bacteria biomass, which are believed to primarily rely on simple (plant-derived) C compounds (Butler et al., 2003; Kramer and Gleixner, 2008). For instance, in the rhizosphere of young beech trees exudates were preferentially utilized by gram-negative bacteria and fungi, with gram-positive bacteria exhibiting little reliance on rhizodeposits (Esperschütz et al., 2009). Gram-positive bacteria, thought to prefer more complex C forms, were proposed to utilize older soil organic carbon upon fresh root exudate inputs (Bird et al., 2011). Consequently, the increase in gram-positive biomass seen 12 days after the labile substrate pulse may be attributed to them benefitting from priming SOM degradation.

PLFAs and NLFAs of the microbial groups (except 18:1 ω 9c) did not differ in their relative ¹³C enrichment (at% ¹³C excess: Fig. 4, Table 3). Still, gram-negative bacteria and gram-positive bacteria incorporated more ¹³C into PLFAs than NLFAs in absolute amounts (μ g ¹³C excess: Fig. 4) which can be attributed to their higher concentrations of PLFAs compared to NLFAs (μ g C: Fig. S5). Given that the extraction of fatty acids occurred 12 days after the substrate pulse, the ¹³C incorporation

does not discern groups that initially utilized ¹³C; rather, it identifies microbial groups that benefitted directly (initial uptake) or indirectly (cross-feeding) from the added substrate over the experimental duration.

Interestingly, we found that the three fungal fatty acid biomarkers responded very differently to the labile substrate input. The fungispecific biomarker 18:1009c exhibited significantly higher at% 13C excess in NLFAs than PLFAs (Fig. 4). In contrast, the 18:109t fungal biomarker did not differ in relative ¹³C enrichment between PLFAs and NLFAs (Fig. 4, bottom), but had higher absolute amounts of 13 C (µg 13 C excess) in PLFAs (Fig. 4, top). The fungal biomarker 18:206,9 was the least responsive to the labile substrate input, with overall low fatty acid biomass, no significant enrichment of ¹³C in the PLFAs, and only little ¹³C incorporated into NLFAs (Fig. 4, Fig. S5). Under the assumption that these fatty acids are representative for different fungal groups or species (Joergensen, 2022), our data may reflect different fungal strategies. While PLFA are part of cell membranes, NLFA originate form fungal storage compounds (Bååth, 2003). Some fungi (represented by 18:1ω9c) may thus have invested into storage compounds whereas others (represented by 18:109t) could have invested into biomass growth (PLFA). Alternatively, it could also be that fungi use the 18:109c NLFA as a storage compound and incorporate the 18:109t PLFA into cell membranes.

5. Conclusion

Our experiment demonstrated the complex microbial response to a spatially constrained input of labile C compounds, i.e. simulating root exudation, in an undisturbed soil. Our findings suggest that such an input pulse has altered microbial metabolism and increased nutrient availability around the exudation spot, revealing the time frame in which this might occur. We were able to identify a three-stage microbial response to artificial root exudation: First, a rapid local substrate respiration (2-3 h), likely depleting oxygen at the exudation spot; secondly, the initiation of (anaerobic) metabolic pathways, accompanied by the production of fermentation byproducts like organic acids and the release of N-compounds (ammonium) (onset approx. 24 h); and thirdly, the accelerated respiration of organic matter (day 3). Regardless, there are several aspects of root exudation we could not capture with our experimental design, such as the release of a more complex mixture of chemical compounds (including N and P), or the presence of an already specialised rhizosphere community inhabiting the root's surface. Our results provide valuable insights into the potential cascade of effects following a substrate pulse, thereby advancing our understanding of the underlying mechanisms of rhizosphere priming.

CRediT authorship contribution statement

Julia Wiesenbauer: Formal analysis, Investigation, Methodology, Software, Validation, Visualization, Writing – original draft, Writing – review & editing. Alexander König: Investigation, Methodology, Software, Visualization. Stefan Gorka: Formal analysis, Investigation, Methodology, Software, Validation, Writing – original draft, Writing – review & editing. Lilian Marchand: Investigation, Methodology. Naoise Nunan: Writing – review & editing, Funding acquisition. Barbara Kitzler: Resources. Erich Inselsbacher: Methodology, Resources, Writing – review & editing, Funding acquisition. Christina Kaiser: Conceptualization, Funding acquisition, Methodology, Project administration, Resources, Supervision, Validation, Writing – review & editing, Writing – original draft.

Declaration of competing 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 to this article can be found online at https://doi.org/10.5281/zenodo.8109601.

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Appendix A. Supplementary data

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