Soil Moisture Affects the Rapid Response of Microbes to Labile Organic C Addition

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Pulsed inputs of labile organic carbon (LOC) are common in soils and significantly affect carbon cycling. However, it remains unclear how soil moisture content affects microbial responses to LOC inputs and the relative contributions of native soil organic matter (SOM) and LOC derived from CO₂ emissions during this process. In this study, we aimed to elucidate how moisture content affects microbial response to LOC inputs and native SOM. Here, ¹³C-labeled glucose was added to soils under nine soil moisture treatments [ranging from 10 to 90% of the water holding capacity (WHC)], and the immediate utilization of LOC and native SOM by microbes was measured. We found that the response of soil microbes to LOC was rapid, and promoted native SOM decomposition. Soil moisture content influenced the microbial usage of LOC and native SOM. A soil water content of 60% WHC was the optimal threshold for changes in the proportion of LOC and native SOM utilized by the microbes. Specifically, we found that when the soil moisture content was below 60% WHC, the ratio between LOC and native SOM increased with increasing moisture content levels. It gradually decreased when the soil moisture content was above 60% WHC. Overall, these findings emphasize the important role of moisture and LOC inputs in soil C cycles.

Keywords: soil moisture content, microbial respiration, LOC inputs, utilization strategies, isotopes

INTRODUCTION

Soil organic matter (SOM) is the largest carbon (C) pool in the terrestrial biosphere, and its decomposition is a vital flux between the soil and atmosphere (Davidson and Janssens, 2006). Hence, slight variations in SOM mineralization have a huge impact on atmospheric CO₂ concentration and may result in dramatic climate and global change (Davidson and Janssens, 2006). The size of the soil C pool depends on the combined effects of C input and SOM decomposition.

Abbreviations: \( R_{\text{Total}} \), Total soil respiration rate; \( R_{\text{LOC}} \), Soil respiration rate from glucose; \( R_{\text{SOM}} \), Soil respiration rate from native soil SOM; \( R_{\text{max}} \), the maximum of respiration rate; \( \text{CO}_2^{\text{Total}} \), total accumulation of CO₂ released; \( \text{CO}_2^{\text{LOC}} \), accumulation of CO₂ released from LOC; \( \text{CO}_2^{\text{SOM}} \), accumulation of CO₂ released from native SOM.
However, the input of C does not imply an increase in soil C storage because exogenous C input may significantly alter native SOM decomposition through a process called priming (Kuzyakov et al., 2000). With global change, temperature will increase and LOC input into the soil will increase due to higher plant productivity increases (Wu et al., 2011), leading to uncertainty in soil C pools (Kuzyakov, 2010). Numerous studies have investigated SOM priming; however, the short-term processes involved in the microbial response to LOC inputs remains unclear. This is due to the rapid response of soil microbes to exogenous substrate inputs, as well as the limitations of measuring technologies (Wang et al., 2016a; Xu et al., 2020; Jiang et al., 2021).

Soil moisture is an important medium for various biochemical reactions in soils, but it also affects soil gas exchange and the availability and mobility of soluble SOM (Brockett et al., 2012). Moreover, it plays a crucial role in influencing microbial activity and community composition, which indirectly affects microbial mineralization of SOM and thus soil C balance (Geisseler et al., 2011; Brockett et al., 2012). Precipitation events are closely linked to soil moisture content and play a critical role in regulating SOM turnover (Austin et al., 2004; Li et al., 2021), particularly in arid and semi-arid regions (Kim et al., 2012; Wang et al., 2016a). It is therefore expected that changes in soil moisture content, resulting from variations in precipitation patterns or soil management regimes in managed landscapes (Dore, 2005), influence the availability of substrates (Schimel et al., 2007; Butterfly et al., 2009) and the mineralization of SOM (Navarro-García et al., 2012). However, it is still unclear how moisture content affects the microbial response to LOC inputs, and the relative contributions of native SOM and LOC to microbial respiration influenced by moisture have not been investigated.

In this study, we aimed to investigate the rapid microbial response to LOC and how soil moisture affects microbial respiration derived from LOC and native SOM. Nine soil moisture treatments [10–90% water holding capacity (WHC)] were established in soil from a semi-arid temperate forest, and δ^{13}C-labeled glucose solutions were added to simulate natural LOC inputs. We used a fully automatic system to measure $^{12}$CO$_2$ and $^{13}$CO$_2$ fluxes over a 48-h period, taking measurements at 3-min intervals, to quantify the microbial respiration rate using LOC and native SOM. As the study site is an arid/semi-arid region with a low soil moisture content in its natural state, soil microbes are sensitive to changes in moisture. Lower soil moisture leads to lower microbial activity (Jiang et al., 2021) and higher soil moisture tends to form an anaerobic environment to inhibit microbial activity (Kool et al., 2011). Thus, we hypothesized that, either extremely high or low moisture content is not conducive to microbial decomposition of LOC and SOM; conversely, optimal moisture content conditions (50–70% WHC) would lead to greater CO$_2$ emissions. Moreover, we predicted that soil moisture aects the contributions of LOC and SOM to microbial respiration. We inferred that under low moisture conditions, native SOM would contribute more to microbial respiration than LOC due to the blocking diffusion of LOC (Schjønning et al., 2003), but that the contribution of LOC to microbial respiration would gradually increases with increasing soil moisture content.

**MATERIALS AND METHODS**

**Study Site and Soil Sampling**

Soil samples were collected from a warm-temperate forest in a semi-arid region of northern China (39°58′N, 115°25′E; 1,278 m elevation) in July 2017, where a native slow-growing oak (Quercus liaotungensis) was the dominant species. Mean annual precipitation and temperature in the forest were 638.8 mm and 6.5°C, respectively, and 74% of the precipitation occurred in summer (from June to August). The soil type was classified as Lithosols (Wang et al., 2016a). Three plots (30 m × 40 m) were randomly established at the study site. After removing the surface litter, we randomly collected more than 10 soil samples from 0 to 10 cm soil depth and combined these into a composite sample for each plot. The fresh soil samples were immediately sieved through a 2 mm diameter sieve to remove rocks, roots, and visible organic debris. These were then taken and transported to the laboratory and divided into three parts: one part was stored at 4°C before conducting the incubation experiments, one part was freeze-dried to measure soil microbial phospholipid fatty acid (PLFA), while the last part was air-dried and processed to measure the soil properties.

A soil-water suspension (1:2.5 v/v) was used to measure the soil pH with a pH meter (Myron L. Company, California, United States). Soil organic carbon (SOC) and total nitrogen (TN) content were measured using an elemental analyzer (Vario EL III, Elemental Analysis System GmbH, Germany). Soil texture was analyzed using a Mastersizer-2000 laser particle analyzer (Malvern Company, Worcestershire, England), and the soil was classified into sand (250–2,000 μm), silt (50–250 μm), and clay fractions (<50 μm). Soil microbial community composition was determined using PLFA biomarker analysis to obtain fungal, bacterial, and actinomycete content (Frostegård et al., 1993; Bååth and Anderson, 2003). The soil properties and microbial characteristics are shown in Table 1.

**Incubation Experiment**

Fresh soil (equivalent to 30 g dry soil) was weighed into a 150 mL plastic culture bottles (n = 3), and the headspace volume was approximately 130 mL after placing the soil sample. As the moisture content of fresh soil exceeded 20% WHC, we lightly air-dried the samples, treated with 10 and 20% WHC moisture, and then adjusted the moisture content. The soil samples were then pre-incubated in a 20°C constant temperature incubator for 7 days to activate the microbes and reduce the interference of the “pulse effect” of water on the decomposition of SOM (Li et al., 2022). Additionally, considering that the added LOC contained water would increase the soil moisture, the soil moisture content during pre-culture was less than the predetermined moisture content to ensure that the soil moisture reached 10, 20, 30, 40, 50, 60, 70, 80, and 90% moisture with the addition of LOC. Water loss in the sample bottles was measured daily and corrected by weight.
The automatic temperature control soil flux system (PRI-8800; PRE-ECO, Beijing, China) developed by He et al. (2013) was modified to measure the soil respiration rates. Specifically, this new device with a PICARRO isotope analyzer (G2131-i, PICTARRO Inc., Sunnyvale, California, United States) enabled observations of δ^{13}C and CO₂ concentrations on a minute-scale.

As shown in Supplementary Figure 1, the device consists of five parts: an analysis, sampling, control, temperature control, and calibration system. Specifically, the soil sample bottles were placed in a 16-hole electric water bath controlled by an automatic temperature regulator and held at 20°C. When the instrument began testing, a collection needle controlled by the control system was automatically inserted into the sample bottle and completely covered the mouth of the bottle, creating an enclosed space. During this time, the instrument was connected to the PICTACRO G2131-i isotope analyzer through a control system to record the δ^{13}C and CO₂ concentrations every second. After measuring one sample, the collection needle was automatically pulled out, and the next sample was measured. During the measurements, each sample bottle was sealed with a preservative film to reduce water loss. Small holes were poked in the preservative film for ventilation, and the sampling needle was inserted through the holes to collect samples from the vials. The dynamics of the respiration rate (R) were measured at 3-min intervals for each sample, and each sample was measured 290 times over a 48-h period. In addition, after every 30 samples, the instrument was automatically calibrated using the calibration system.

We added glucose as a simple analog for LOC input. Specifically, three incubation bottles were removed from the instrument. Then, 1 mL ¹³C-uniformly labeled glucose (δ^{13}C = 3,864‰) solution was added slowly and uniformly to the soil by pipetting the glucose onto the soil surface at a controlled and consistent rate. This was done to ensure uniform distribution in this soil environment should be minimal, because inorganic C is relatively stable. Therefore, we assumed that the released CO₂ is derived directly from microbial respiration and did not consider the C emissions potentially caused by abiotic factors. Additionally, the fractionation during the biodegradation processes was negligible (Mary et al., 1992). The proportions of CO₂ derived from LOC and SOM components were calculated using a mass balance model:

\[
f_{\text{LOC}} = \frac{(\delta_{\text{Total}} - \delta_{\text{SOM}})}{(\delta_{\text{LOC}} - \delta_{\text{SOM}})} \tag{1}
\]

where \(f_{\text{LOC}}\) is the fraction of total CO₂ derived from glucose, \(\delta_{\text{Total}}\) is the measured isotopic value of the sample obtained at each measurement point, \(\delta_{\text{SOM}}\) is the mean of the isotopic value of the control treatment, and \(\delta_{\text{LOC}}\) is the isotopic value of the added glucose treatment.

### Calculation of the Microbial Respiration Rate

The total soil microbial respiration rate (\(R_{\text{Total}}\); \(\mu g \ C \ g^{-1} \ soil \ h^{-1}\)) was calculated using Eq. 2:

\[
R_{\text{Total}} = \frac{C \times V \times \alpha \times \beta}{m} \tag{2}
\]

where \(C\) is the slope of the CO₂ concentration, \(V\) is the volume of the incubation bottle and gas tube, \(m\) is the soil dry weight (g), \(\alpha\) is the conversion coefficient of the CO₂ mass, and \(\beta\) is the conversion coefficient of time (He et al., 2013).

\(R_{\text{LOC}}\) and \(R_{\text{SOM}}\) are associated with LOC and native SOM decomposition, respectively. \(R_{\text{LOC}}\) and \(R_{\text{SOM}}\) were calculated using Eqs 3 and 4, respectively (Li et al., 2022):

\[
R_{\text{LOC}} = R_{\text{Total}} \times f_{\text{LOC}} \tag{3}
\]

\[
R_{\text{SOM}} = R_{\text{Total}} - R_{\text{LOC}} \tag{4}
\]

### Calculation of the CO₂ Released

During the 48-h incubation, the accumulation of CO₂ release (\(\mu g \ C \ g^{-1} \ soil\)) was calculated as follows:

\[
CO₂ = \sum_{i=1}^{n} \frac{(R_{t_i} + R_{t_i+1})}{2} \times (t_{i+1} - t_i) \tag{5}
\]

where \(R_{t_i}\) and \(R_{t_i+1}\) are the respiration rates at \(t_i\) and \(t_{i+1}\), respectively. The cumulative CO₂ released from LOC, native SOM, and the total CO₂ present was expressed as \(CO₂_{\text{LOC}}, \ CO₂_{\text{SOM}},\) and \(CO₂_{\text{Total}},\) respectively.

### Statistical Analysis

As shown in Figure 1, the corresponding parameters were defined to better determine the microbial response to the LOC input. Tukey’s HSD test was used to identify differences in \(R_{\text{max}}\) time for \(R_{\text{max}}\), absolute change in \(R\), and accumulation of CO₂ released among different soil moisture treatments. Curve fitting was used to explore the relationships between the soil moisture and the above parameters. Statistical analyses were performed using the SPSS software (SPSS for Windows, Version 13.0; SPSS Inc., Chicago, Illinois, USA).

### Table 1: Soil properties used in this study.

| δ^{13}C (%) | Soil pH  | SOC (%) | TN (%) | Sand (%) | Silt (%) | Clay (%) | PLFA (nmol g⁻¹) | Bacterial (nmol g⁻¹) | Fungi (nmol g⁻¹) | Actinomycete (nmol g⁻¹) |
|------------|---------|---------|-------|---------|--------|--------|--------------|----------------|----------------|------------------------|
| -25.2 ± 0.4 | 6.58 ± 0.49 | 3.9 ± 0.1 | 0.3 ± 0.01 | 25.22 ± 11.92 | 68.79 ± 10.81 | 5.98 ± 1.49 | 13.47 ± 1.87 | 5.39 ± 1.36 | 2.38 ± 0.52 | 0.34 ± 0.05 |

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\]

where \(f_{\text{LOC}}\) is the fraction of total CO₂ derived from glucose, \(\delta_{\text{Total}}\) is the measured isotopic value of the sample obtained at each measurement point, \(\delta_{\text{SOM}}\) is the mean of the isotopic value of the control treatment, and \(\delta_{\text{LOC}}\) is the isotopic value of the added glucose treatment.

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the ratio of $\text{CO}_2^{\text{SOM}}/\text{CO}_2^{\text{Total}}$ exhibited a decreasing and then increasing relationship (Figures 5D,E). Additionally, the ratio of $\text{CO}_2^{\text{LOC}}/\text{CO}_2^{\text{SOM}}$ also revealed an increasing and then decreasing relationship with soil moisture (Figure 5F) and a soil moisture content of approximately 60% WHC. $\text{CO}_2^{\text{LOC}}/\text{CO}_2^{\text{Total}}, \text{CO}_2^{\text{SOM}}/\text{CO}_2^{\text{Total}},$ and $\text{CO}_2^{\text{LOC}}$ reached their extreme values when the soil moisture content was approximately 60% WHC.

**DISCUSSION**

**Soil Moisture Content Affects the Response of Soil Microbes to Labile Organic Carbon Addition and the Decomposition of Native Soil Organic Matter**

Soil moisture content is an important environmental factor that affects microbial activity, the diffusion rate of gases, and the availability and mobility of soluble SOM (Casals et al., 2000; Liu et al., 2009). Therefore, changes in soil moisture content have a profound impact on soil microbial respiration and SOM mineralization. In general, aerobic microbial respiration is optimal when the soil moisture content is approximately 60% WHC (Das et al., 2019). Here, we observed that extremely high or low moisture content was not conducive to microbial utilization of both LOC and native SOM. However, when soil moisture content was 50–70% of WHC, the responses of soil microbes to added LOC were stronger than those observed under other soil moisture treatments; that is, $R_{\text{max}}$ and $\text{CO}_2$ released were greater. This finding supports our hypothesis, and consistent with findings in related studies (Geisseler et al., 2011; Wang et al., 2016b). We found that soil microbes with lower moisture content (<30%WHC) were less responsive to added LOC (lower $R_{\text{max}}$ and $\text{CO}_2$ LOC). The movement of the glucose solution strongly depends on the soil moisture content (Casals et al., 2000; Liu et al., 2009). During low soil moisture treatment, the downward movement of the glucose solution was
much slower than that during high soil moisture treatment. Microbes and their enzymes responsible for macromolecule degradation can be affected by moisture (Šnajdr et al., 2008; Geisseler et al., 2011); low soil moisture content affects the utilization of added LOC by microbes (Xue et al., 2017). That is because the diffusion of substrate may slow down due to lack of water which limits the substrate supply to soil microbes (Schjønning et al., 2003). Additionally, as the matrix potential of soil moisture decreases, soil microbes need to consume additional energy to maintain osmotic balance with their surroundings (Schimel et al., 2007). They also need to produce extracellular substances to buffer soil moisture changes and improve diffusivity (Or et al., 2007). Together, these factors lead to a reduced metabolic activity of soil microbes and a limited rate of reproduction and respiration. The utilization of LOC by microorganisms is also reduced when the moisture content is too high (> 70% WHC). High moisture content creates an anaerobic environment (Hackl et al., 2005), which inhibits microbial activity and is unfavorable for the utilization of LOC by microorganisms.

Although we found that microbes preferred the utilization of LOC over native SOM, the simultaneous effect of LOC on microbial activity further promoted the decomposition of native SOM. This decomposition was also influenced by soil moisture. Generally, microbes are limited by the availability of C substrates (Cleveland et al., 2007), and LOC inputs increase substrate availability, relieve microbial C limitations, and increase microbial activity. This in turn promotes the decomposition of native SOM (Weedon et al., 2013). The microbial nitrogen (N)-mining hypothesis can be used to explain this phenomenon (Craine et al., 2007). Although the addition of LOC increases the availability of C substrate to microbes, microbial growth and metabolism will be limited by N, when microbes need to decompose more native SOM to obtain nutrients to meet growth requirements, and thus stimulate the decomposition of native SOM (Chen et al., 2014).

Our results show an interesting phenomenon: when the soil moisture was below 60% WHC, the absolute change in $R_{SOM}$ was less than 0 but that of $R_{LOC}$ was greater than 0, after 1 h of LOC input (Figures 3G,H). This implies the microbial...
preferential utilization of LOC and an inhibition of SOM decomposition (Li et al., 2022). Generally, microbial availability of organic matter depends on its chemical recalcitrance, mineral association, and accessibility to decomposers (Deng et al., 2020). Microbes are limited by the availability of C substrates, and thus soil microbes prefer to utilize labile substrate with a low molecular composition when compared to recalcitrant SOM, as less energy is required for mineralization (Deng et al., 2020). Additionally, it is difficult for microorganisms to access and decompose the dissolved soil C due to the microbial activity is limited by the low moisture content (Curiel Yuste et al., 2007). This allows them to preferentially use LOC, thereby inhibit microbial decomposition of native SOM (Blagodatskaya and Kuzyakov, 2008). In contrast to lower soil moisture content, optimal soil moisture content allows the cleavage and dispersion of soil aggregates, which leads to the dissolution and release of SOM encapsulated within the aggregates, thereby allowing more and better access to and decomposition of native SOM by soil microbes (Marín-Spiotta et al., 2014). In contrast, when soil moisture is saturated (>70% WHC), the oxidation-reduction decreases and anaerobic conditions limit the activity of soil microbial enzymes (Hackl et al., 2005). Furthermore, any soil moisture content above 70% WHC also reduces the diffusion of O₂ and forms a flooded anaerobic environment. This environment inhibits the activity of aerobic microbes and various oxidative enzymes, thereby affecting the mineralization of C by microbes (Schjønning et al., 2003).

**Soil Moisture Affects the Proportion of Labile Organic Carbon and Native Soil Organic Matter Used by Microbes**

The proportions of LOC and native SOM utilized by microbes differed in response to different soil moisture content. This provides quantitative evidence that soil moisture content can
affect how microbes utilize LOC and native SOM, through $R_{\text{LOC}}/R_{\text{SOM}}$, CO$_2$−LOC/CO$_2$−Total, CO$_2$−SOM/CO$_2$−Total, and CO$_2$−LOC/CO$_2$−SOM. Our study found that soils have greater CO$_2$ fluxes under suitable moisture conditions (approximately 60% WHC). However, this is inconsistent with our alternative hypothesis that microbes would utilize more LOC under higher moisture conditions, as we found that microbes have different strategies for utilizing LOC and native SOM under different moisture conditions. We found that the relative proportions of the metabolism of soil microbes using the two C sources to total respiration were different under different moisture conditions (Figure 5). Under lower moisture conditions (<30% WHC), more native SOM was used for microbial respiration. This may occur because low moisture conditions are not conducive to the diffusion of LOC; thus, microbes predominantly use native SOM (Davidson et al., 2010). Furthermore, the decrease in soil water potentially may stimulate microbial responses to water stress. This stimulation results in an increase in soil microbial C-use efficiency (Herron et al., 2009), which can lead to lower LOC for respiration. However, as soil water content increased, more microbes had access to LOC, resulting in more LOC for respiration. Thus, values of CO$_2$−LOC/CO$_2$−Total increased and those of CO$_2$−SOM/CO$_2$−Total decreased. Moreover, when the soil moisture content was too high (>60% WHC), the proportion of native SOM utilization by microbes gradually increased again (Figure 5B). This increase occurs because moisture content affects the microbial C use efficiency. Many studies have found that the native SOM mineralization rate is highest under higher moisture content conditions after LOC input, which is consistent with our results (Guenet et al., 2013; 2014).
Bian et al. (2016). When the water content is too high, it contributes to the disruption of the soil aggregate structure and SOM leaching and increases the contact area between native SOM and decomposers, thus promoting native SOM mineralization (Marín-Spiotta et al., 2014). Moreover, it is possible that LOC input resulted in local energy excesses and nutrient deficits caused by the highly heterogeneous SOM forming many relatively independent microhabitats. Meanwhile, soil flooding resulted in a homogeneous mixing of decomposing substrates and adequate utilization of LOC, which promoted the utilization of native SOM by microbes (Guenet et al., 2013).

Here, we provide quantitative evidence that soil moisture content can influence the way microbes utilize LOC and native SOM and propose potential mechanisms of influence (Figure 6). However, microbial taxa determine the composition and functional properties of microbial communities, which organize and structure their responses to resource availability and changes in environmental conditions and play a key role in the C cycle (Banerjee et al., 2018). Theoretically, soil microbes are divided into $r$- and $K$-strategists according to their biological characteristics and utilization of resources. Among them, $r$-strategists (mainly bacteria) grow faster and generally use LOC, whereas $K$-strategists grow slowly and can use more resistant organic matter (Fontaine et al., 2003). Therefore, future studies should examine the dynamics of soil microbes. This will help us to fundamentally understand the microbial strategies for the utilization of LOC and native SOM under different moisture conditions.

**CONCLUSION**

In this study, we demonstrated that soil moisture content influences the microbial response to LOC and native SOM. We show that a soil moisture content of 60% WHC is the optimal and allows for changes in the proportion of LOC and native SOM utilized by microbes. Additionally, we found that when the soil moisture content was below 60% WHC, the proportion of microbial respiration derived from LOC increased with increasing soil moisture content. In contrast, this proportion gradually decreased when the soil moisture content was above 60% WHC. Moreover, under suitable moisture conditions (approximately 60% WHC), the soil exhibited greater CO$_2$ emissions. Overall, our results emphasized that microbial utilization of LOC and native SOM was significantly affected by soil moisture content. These findings may be important considering climate induced changes in precipitation and increased C inputs.

**DATA AVAILABILITY STATEMENT**

The raw data supporting the conclusions of this article will be made available by the authors, without undue reservation.
AUTHOR CONTRIBUTIONS

NH and JZ designed the research. HB and SZ conducted the study and analyzed the data. CL drafted the manuscript with input from HB and JZ. ML and LX reviewed and revised the manuscript. All authors contributed to revisions.

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SUPPLEMENTARY MATERIAL

The Supplementary Material for this article can be found online at: https://www.frontiersin.org/articles/10.3389/fevo.2022.857185/full#supplementary-material

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