Priming effects in biochar enriched soils using a three-source-partitioning approach: $^{14}$C labelling and $^{13}$C natural abundance

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1. Introduction

The short term increases or decreases in the mineralization of soil organic C caused by the addition of organic substrates are known as positive or negative priming effects (Kuzyakov et al., 2000). Previous studies showed that added biochars could induce either positive or negative priming effects (Luo et al., 2011; Zimmerman et al., 2011). Negative priming effects were attributed to the physical protection of soil organic matter (Maestrini et al., 2014b). However, some studies indicated that physical protection through soil aggregation might be weakly linked to biochar induced priming effects (Kerre et al., 2016). Positive priming effects occur in the early stages following biochar incorporation, and are reported to be over quite rapidly in many studies (Jones et al., 2011a; Maestrini et al., 2014a; Whitman et al., 2014; Zhao et al., 2013; Zimmerman et al., 2011). These short term positive priming effects, due to increased mineralization of soil organic C, are largely attributed to stimulation of microbial activity by the labile C contained within the biochar, or abiotic release of CO$_2$ from carbonates in the ash (Maestrini et al., 2014b; Smith et al., 2010; Zimmerman et al., 2011). The positive priming effects observed following the addition of biochar are of similar magnitude (between 50 and 500 mg CO$_2$-C g$^{-1}$ soil) to other priming effects following the addition of labile organic C, which stimulates microbial activity in the short term (Jones et al., 2011a; Zhao et al., 2013). Thus, biochar induced priming effects are generally believed to be either positive or...
negative in the short term while mainly negative after labile C has been utilized in the long term.

After the short term mineralization of bio-available C from the biochar, the effects of biochar on soil properties, e.g. water-filled pore spaces, habitat, soil aeration, moisture, pH and nutrient availability may persist during the medium (1 year) to long term (above 3 years) period. In addition to environmental factors, e.g. temperature (Benbi et al., 2014; Fang et al., 2014), soil priming effects can also be strongly affected by soil properties, e.g. soil pH, aggregate stability, organic matter and nutrient availability (Blagodatskaya and Kuzyakov, 2008). For example, a highly significant positive correlation between soil primed CO2 (expressed as a percent of added C) and soil pH (range 3–8) was found by Blagodatskaya and Kuzyakov (2008). As soil properties might be changed due to biochar addition, the soil organic C mineralization in biochar free soil and biochar amended soil may respond differently to added substrates. Thus, in addition to the short term effects of biochar on soil organic C mineralization, following addition of substrate to the newly established biochar soil system, the direction or magnitude of the soil priming effect might differ due to the changes in soil properties caused by the biochar.

The first report on biochar causing soil priming effects was criticized because isotopic labelling was not used. It was therefore not possible to separate the different SOM pools involved (Lehmann and Sohi, 2008; Wardle et al., 2008). Since then, stable isotope techniques have been used to distinguish between the mineralization of biochar and soil organic C mineralization in many studies (Jones et al., 2011a; Maestrini et al., 2014a; Maestrini et al., 2014b; Zimmerman et al., 2011). Until now, most studies designed to investigate biochar induced soil priming effects have used two approaches: 1) addition of unlabeled biochar to unlabeled soil (Wardle et al., 2008), and 2) addition of 13C (Jones et al., 2011b; Zimmerman et al., 2011) or 14C labelled biochar to unlabeled soil (Kuzyakov et al., 2009). However, quantitatively partitioning systems using two stable isotopes (13C and 14C) have been limited to only two sources (biochar C and soil organic C). Recently there has been increasing interest in the effects of biochar on priming effects involving three C sources e.g. soil, biochar and plants (Weng et al., 2015; Whitman et al., 2014). Whitman et al. (2014) partitioned the CO2-C evolved from three sources of biochar, plant root exudates and soil organic C using only two isotopes, by assuming an extreme scenario whereby only one C source was mineralized from the combined C sources of biochar plus root exudates, which gave the upper (root exudates) and lower (biochar) values of the mineralization of C4 materials and soil organic C mineralization. Whitman and Lehmann (2015) introduced a dual-isotope approach to partition soil CO2 emissions derived from soil organic C, added biochar and root respiration. However, it still remains challenging for three C sources to be separated by traditional approaches using only two stable isotopes (13C and 14C). More complex methodology approaches that are able to discern three C sources in biochar enriched systems would provide a much needed and valuable research tool in future research (Weng et al., 2015; Whitman et al., 2014).

Based on the biochar induced changes in soil properties, e.g. pH increase, aeration, moisture dynamics, C availability, nutrients status, our hypothesis is that soil organic C mineralization will respond differently to the addition of fresh organic C in unamended and biochar-amended soils. Our main aim was to test the possibility of adopting the three-source-partitioning approach by combining 14C labelling with 13C natural abundance in soil with three C sources (native soil organic C, glucose-C and biochar-C). The second aim was to investigate the differences in amounts of primed CO2 following the addition of 14C glucose to an unamended (soil and glucose) and biochar amended (soil, glucose and biochar) soil.

2. Materials and methods

2.1. Soils, biochar and glucose addition

Soils were sampled (0–23 cm) with a Dutch auger, from Zhejiang province, China (Soil A) and north-west of Gottingen, Germany (Soil B). Both soils were classified as Luvisols according to World Reference Base (WRB) system with soil classification scheme. The two soils were analyzed in different labs but with identical procedures. The soil samples were hand-picked to remove obvious plant debris and roots, sieved at field moisture (<2 mm) and subsequently adjusted to 40% of water holding capacity (WHC). Soil pH was measured at a soil:CaCl2 ratio of 1:2.5 (weight/weight). Air-dry soil (10 g, <2 mm) and 25 ml of CaCl2 (0.01 M) were shaken together for 1 min and left to settle for 30 min, which was repeated once more before pH was determined with a pH electrode. Total C and N of the soils (air-dried, milled <200 μm) were determined by dry combustion (LECO CNS 2000, LECO Corporation, Michigan, USA). The natural δ13C (‰) abundance of the soils (air-dried, milled <200 μm) was determined on an elemental analyser-isotope ratio mass spectrometer (Sercon Ltd, Crewe, UK). All measurements are given on an oven-dry weight basis (o.d., 105°C, 24 h). Soil A had a pH (CaCl2) of 7.0, a bulk density of 1.3 g cm−3, an organic C content of 60.5 g kg−1, a total N content of 3.8 g kg−1 and a δ13C value of −26.7‰ (C3-source). Soil B had a pH (CaCl2) of 6.0, a bulk density of 1.4 g cm−3, an organic C content of 12.4 g kg−1, a total N content of 1.3 g kg−1 and a δ13C value of −27.4‰ (C3-source). More information about soil B including its properties and management were reported by Kramer et al. (2012).

The biochar was made from maize straw (C4-source), previously dried at 105°C for 24 h, milled <1 mm, contained within a sealable retort with N2 flowing to limit O2, and pyrolyzed in a Carbolite CFW 1200 furnace at 400°C for 30 min. Biochar was analyzed in the same way as the soils, except that the pH was measured at a soil:CaCl2 ratio of 1:10 (weight/weight). The biochar analytical values are given in Table 1.

Uniformly labelled 13C glucose as substrate (corresponding to 50 KBq g−1 soil) was prepared 12 h before the incubation study started. The δ13C of the glucose was −9.905‰.

2.2. Experimental design and layout

The experimental design comprised 15 treatments: 2 unamended soils (A and B), 2 biochar amended soils (soils A and B amended with biochar), plus biochar alone supplied with dissolved soil organic C (DOC) solution (1:20 soil: water ratio). All of these 5 treatments were supplied with 3 additions (water and 2 glucose levels). Soils from both treatments (C3 soil plus C4 biochar, and C3 soil) were amended with distilled water as a control to estimate the changes in soil mineralization of C4 materials and soil organic C mineralization. Whitman and Lehmann (2015) introduced a dual-isotope approach to partition soil CO2 emissions derived from soil organic C, added biochar and root respiration. However, it still remains challenging for three C sources to be separated by traditional approaches using only two stable isotopes (13C and 14C). More complex methodology approaches that are able to discern three C sources in biochar enriched systems would provide a much needed and valuable research tool in future research (Weng et al., 2015; Whitman et al., 2014).

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mixed in plastic bags to homogenize them. Uniformly labelled \(^{14}\text{C}\)glucose was added to a solution of unlabeled glucose to reach the final concentration of 5.7 KBq g\(^{-1}\) soil. \(^{14}\text{C}\)-labelled glucose was applied to soil at 100 mg C kg\(^{-1}\) (glucose low, GL) or 1000 mg C kg\(^{-1}\) soil (glucose high, GH) in water to provide final soil moisture contents of 50\% WHC. \(\text{NH}_4\text{NO}_3\) was added, in solution, at 170 mg N kg\(^{-1}\) soil (glucose high, GH) in water and dried overnight (80 \(\text{C}\)) glass jars. The soils were adjusted to 40\% WHC and pre-incubated for one week at 20 \(\text{C}\). After pre-incubation, uniformly labelled \(^{14}\text{C}\) glucose and distilled water were added to reach a final soil moisture content of 50\% of WHC. Four empty jars served as blanks. Then the jars were incubated in the dark at 20 \(\text{C}\) for 28 days. During the incubation, the CO\(_2\) evolved from the soils was trapped by 3 ml of 1.0 M NaOH solution in small placed on the soil surface, and exchanged at 1, 3, 5, 7, 14, and 28 days. Aliquots of NaOH from the three randomly chosen replicate vessels from each treatment were used to measure the \(^{13}\text{C}\) activity, \(^{13}\text{C}-\text{CO}_2\) (\(\text{‰}\)) and total amount of trapped CO\(_2\).

2.3. Chemical analysis

2.3.1. Soil CO\(_2\) emission and \(^{13}\text{C}\) (\(\text{‰}\))

The concentrations of CO\(_2\) trapped in the NaOH solutions were measured by titrating 0.5 ml NaOH with 0.1 M HCl with phenolphthalein as an indicator after addition of 0.5 M BaCl\(_2\). To determine the \(^{13}\text{C}\) (\(\text{‰}\)) of the trapped CO\(_2\)-C, 2 ml aliquots of the NaOH were added to 3 ml 1.5 M BaCl\(_2\) in vials (Aoyama et al., 2000). The resulting BaCO\(_3\) precipitates were then filtered and trapped on glass fibre filters (90 mm, Whatman GF/A, UK), carefully rinsed with water and dried overnight (80 \(\text{C}\)). The precipitates were scraped off the filters, weighed (5 mg) into tin capsules and analyzed for \(^{13}\text{C}\) on an elemental analyser-isotope ratio mass spectrometer (DELTA V plus IRMS, Thermo Fisher Scientific, Bremen, Germany).

2.3.2. Glucose derived-\(^{14}\text{C}\) in CO\(_2\) pool

The \(^{14}\text{C}\) activity of CO\(_2\) trapped in NaOH was measured in a scintillation cocktail (Rotiszint Eco PlusCarl Roth, Germany) after decay of the chemiluminescence using a 1450 LSC & Luminescence Counter MicroBetaTriLux (Perkin Elmer Inc., USA). The \(^{14}\text{C}\) counting efficiency was 87\% and the \(^{14}\text{C}\) activity measurement error did not exceed 2\%.

2.4. Calculation

To partition three sources of CO\(_2\)-C, an approach combining \(^{14}\text{C}\) labelling with \(^{13}\text{C}\) natural abundance, the calculation of Blagodatskaya et al. (2011) and Tian et al. (2016) was used. Initially, the amount of glucose-derived C (\(\text{C}_{\text{G\text{-derived}}}\)) was calculated based on the radioactivity of the evolved \(^{14}\text{CO}_2\) (\(\text{C}_{\text{G\text{-cur}}}, \text{DPM}\)), the amount of added glucose (\(\text{C}_{\text{G}}\)), and the radioactivity of the applied glucose (\(\text{C}_{\text{G\text{-DPM}}}\)):

\[
\text{C}_{\text{G - derived}} = 14\text{C}_{\text{cur}} \times \text{CG}/14\text{C}_{\text{G}}
\]  

(1)

Then, the amount of SOM-derived C was calculated as:

\[
\text{C}_{\text{SOM - derived}} = \text{C}_{\text{total}} - \text{C}_{\text{G - derived}}
\]  

(2)

where \(\text{C}_{\text{total}}\) is the total amount of C in the evolved CO\(_2\).

Secondly, the \(^{13}\text{C}\) (\(\text{‰}\)) values of SOM-originated C in each pool (\(^{13}\text{C}_{\text{SOM-derived}}\)) were calculated based on a mass balance equation. The \(^{13}\text{C}\) signature of glucose-derived C (see below) was subtracted from the total \(^{13}\text{C}\) signature, considering the contribution of the amount of glucose-originated C estimated in the first step based on \(^{14}\text{C}\):

\[
^{13}\text{C}_{\text{SOM-derived}} = \left( \text{C}_{\text{total}} - \text{C}_{\text{G-derived}} \right) ^{13}\text{C} - \left( \text{C}_{\text{G-derived}} \right) ^{13}\text{C}
\]  

(3)

where \(^{13}\text{C}_{\text{total}}\) and \(^{13}\text{C}_{\text{G-derived}}\) are the \(^{13}\text{C}\)-values of the total and glucose-derived C. The former value was measured experimentally as described in section 2.3.1. The \(^{13}\text{C}\) of \(\text{G\text{-derived}}\) was assumed to be equal to the \(^{13}\text{C}\) of glucose (\(-9.90\%\)).

The contributions of soil organic C (C3) and biochar C (C4) were then calculated based on the glucose-corrected \(^{13}\text{C}\) signature (\(^{13}\text{C}_{\text{SOM-derived}}\)) at each corresponding date. The amount of C4-derived CO\(_2\)-C was calculated from:

\[
\begin{align*}
\text{C}_{\text{C4-derived}} & = \frac{^{13}\text{C}_{\text{SOM-derived}} \times ^{13}\text{C}_{\text{G-derived}} - ^{13}\text{C}_{\text{C3-ref}}}{^{13}\text{C}_{\text{C4-material}} - ^{13}\text{C}_{\text{C3-materia}}} \\
& = \frac{^{13}\text{C}_{\text{SOM-derived}} \times ^{13}\text{C}_{\text{G-derived}} - ^{13}\text{C}_{\text{C3-ref}}}{^{13}\text{C}_{\text{C4-material}} - ^{13}\text{C}_{\text{C3-materia}}}
\end{align*}
\]  

(5)

where \(^{13}\text{C}_{\text{C3-ref}}\) is the \(^{13}\text{C}\) value in the reference C3 soil at the corresponding sampling date.

The \(\text{C}_{\text{C4-derived}}\) was then calculated by subtracting the \(\text{C}_{\text{C4-derived}}\) from the total amount of C.

The glucose induced priming effects in the biochar amended soils, with and without glucose addition (C3+C4) were calculated based on the changes in the \(^{13}\text{C}\) signature and the amount of extra CO\(_2\) evolved after \(^{14}\text{C}\)-glucose addition compared with the treatment without glucose.

\[
\begin{align*}
\text{PE} & = \text{Biochar soil - C_{glucose amended} - Biochar soil} \\
& - \text{C_{without glucose}}
\end{align*}
\]  

(6)

Then the glucose induced priming effects in unamended soils (C3) were calculated from

\[
\text{PE} = \text{soil derived C_{glucose amended} - soil derived C_{without glucose}}
\]  

(7)

The changes in the \(^{13}\text{C}\) signature due to preferential substrate utilization of labile and \(^{13}\text{C}\)-enriched biochar C (compared with \(^{13}\text{C}\) depleted soil C), were considered to provide the correct assessment of the priming effect (Blagodatskaya et al., 2011). The dynamic changes in \(^{13}\text{C}\) due to preferential utilization were estimated in biochar enriched soil (C3+C4) treated solely with H\(_2\)O. Therefore, the priming effects were calculated separately for C3 and C4 carbon sources (C3-PE and C4-PE, respectively) considering the changes in

| Table 1 |
| Biochar characterization. |

<table>
<thead>
<tr>
<th>Total C (%)</th>
<th>C/N</th>
<th>C/H</th>
<th>C/O</th>
<th>Ash content (%)</th>
<th>DOC (mg g(^{-1}))</th>
<th>CEC (c mol/kg)</th>
<th>pH</th>
<th>Total K (mg/g)</th>
<th>(^{13}\text{C})</th>
</tr>
</thead>
<tbody>
<tr>
<td>50.27 (3.99)(^a)</td>
<td>30.31 (0.30)</td>
<td>14.70 (0.06)</td>
<td>2.57 (0.24)</td>
<td>20.2 (0.15)</td>
<td>0.600 (0.012)</td>
<td>116.20 (5.46)</td>
<td>10.06 (0.011)</td>
<td>8.60 (0.02)</td>
<td>-14.29 (0.02)</td>
</tr>
</tbody>
</table>

\(^a\) Means ± standard error.

The \(^{13}\text{C}\) value of biochar, it was supplied with dissolved soil organic C (DOC) (soil free), the measured \(^{13}\text{C}\) value of released CO\(_2\) from biochar plus DOC treatment is the biochar \(^{13}\text{C}\) value. The \(^{13}\text{C}\) of the biochar can be found in Table 1.
the contribution of soil C (C3) and biochar C (C4) in the controls:

\[
C3 - PE = \frac{C_{\text{amended}}}{C_{\text{unamended}}} - C3
\]

\[
C4 - PE = \frac{C_{\text{amended}}}{C_{\text{amended}}} - C4
\]

2.5. Statistics

Data meeting assumptions of normality and equality of variances were analyzed by ANOVA. Data not meeting the assumptions were transformed logarithmically and analyzed by ANOVA. A one-way analysis of variance was undertaken to determine the significance (p < 0.05) of differences between the treatments of glucose induced priming effects with and without biochar (SPSS 19.0). LSD was chosen for the multiple comparisons between different treatments.

3. Results

3.1. CO2 release from soil

Soils that received glucose gave much higher CO2 fluxes than soils not receiving glucose, during the 28 days of incubation (Fig. 1, P < 0.001), with significant differences between the two glucose concentrations. Soils with 100 \(\mu\)g glucose g\(^{-1}\) soil evolved lower concentrations of CO2, (less than 1000 \(\mu\)g C g\(^{-1}\) soil) while the soil receiving glucose 1000 \(\mu\)g C g\(^{-1}\) soil at evolved much more CO2 (more than 1500 \(\mu\)g C g\(^{-1}\) soil) over 28 days of incubation (Fig. 1).

At 100 \(\mu\)g glucose C g\(^{-1}\) soil, rapid C mineralization occurred in the first few days, then glucose mineralization ceased (Fig. 1a and c), while glucose-derived CO2 continued to be evolved after 28 days of incubation at 1000 \(\mu\)g C g\(^{-1}\) soil. In addition, glucose at the low (GL) and high (GH) concentrations evolved different amounts of CO2, at 29.9 and 285.0 \(\mu\)g C g\(^{-1}\) soil, respectively, over 28 days of incubation. Percentage mineralization of added glucose in GL and GH treatments were similar (29.4% and 30.2% of added glucose respectively). Compared to biochar free soil, biochar amended soil gave higher CO2 emissions (P < 0.05), with 300.1 and 298.5 \(\mu\)g C g\(^{-1}\) evolved from soil A and B, respectively (Fig. 1b). The addition of GH resulted in the largest soil CO2 evolution in the biochar enriched soil B, at 1534.2 \(\mu\)g C g\(^{-1}\) soil (Fig. 1d), which was not significantly different from the CO2 evolved from the corresponding treatment in Soil A (1468.3 \(\mu\)g C g\(^{-1}\) soil) (Fig. 1b).

3.2. Partitioning the total soil CO2-C evolved

By combining \(^{14}\)C labelling with \(^{13}\)C natural abundance measurements (Blagodatskaya et al., 2011), it is possible to partition the CO2-C from \(^{14}\)C glucose, native soil organic C and biochar (Fig. 2). In both soils with glucose addition, the cumulative C4-derived CO2 was greater than the CO2 efflux derived from C3-soil organic C at the beginning of the incubation +8 (day 3) but there was no significant difference (P > 0.05) by the end of the incubation. The mineralization of C4-biochar C mainly occurred during the first 7 days of incubation. For example, in Soil B, significantly more biochar C (P < 0.001) was mineralized at day 7 (368.8 \(\mu\)g C g\(^{-1}\)) compared to day 3 (272.7 \(\mu\)g C g\(^{-1}\)) with GL addition and from 413.9 \(\mu\)g C g\(^{-1}\) to 550.8 \(\mu\)g C g\(^{-1}\) with GH addition over the same period, but only another 10% (GL) and 4% (GH) of biochar C were mineralized from day 7 to day 28 (Fig. 2). In both soils, biochar mineralized very slowly and no biochar mineralization was detected after day 7. Soil organic C (C3) mineralization depended largely on the glucose addition rate, irrespective of soil type. The GL rate mainly stimulated C3 soil mineralization during the first 14 days (314 \(\mu\)g C g\(^{-1}\) in soil A and 332 \(\mu\)g C g\(^{-1}\) in Soil B, respectively), and produced much lower mineralization rates (49.2 \(\mu\)g C g\(^{-1}\) in Soil A and 27.3 \(\mu\)g C g\(^{-1}\) respectively) in Soil B from day 14–28, while the high concentration of glucose (GH) caused continuous C3 soil organic C mineralization with no decrease in rate, with the CO2 evolved increasing from 346.2 \(\mu\)g C g\(^{-1}\) to 486.9 \(\mu\)g C g\(^{-1}\) (Soil A) and 435.8 \(\mu\)g C g\(^{-1}\) to 574.2 \(\mu\)g C g\(^{-1}\) (Soil B) during the corresponding period from day 14–28 (Fig. 2).

3.3. Glucose induced priming effect

Following the addition of glucose at the high rate (GH), different trends of soil organic C mineralization occurred in unamended and biochar amended soil. The soil organic C loss was hardly detectable in any treatments apart from the GH plus biochar enriched soil,
where the continuous mineralization of soil organic C extended after the 28 day incubation period (Fig. 3). The GH induced soil primed C losses in the biochar enriched soil during 28 days of incubation were 457.0 and 423.5 μg CO2-C g⁻¹, respectively, in soil A and B (Fig. 3 b, d and Fig. 4), which were 140% and 53% higher respectively than the primed soil CO2 in biochar free soil (Fig. 3b,d). Following GL addition, no significant differences occurred between biochar free and biochar enriched Soil A (Fig. 3a) during the 28 day

Fig. 2. Contribution of three C sources to cumulative CO2 evolved from soil A with low at 100 mg C kg⁻¹ soil (a) and high concentration of glucose addition at 1000 mg C kg⁻¹ soil (b), and soil B with low (c) and high (d) concentration of glucose addition. The three sources of CO2-C were: 1) Soil derived CO2-C (C3-Soil), 2) Biochar derived CO2-C (C4-Biochar), and 3) Glucose derived CO2-C (14C). Error bars indicate standard errors of the means (n = 3).

Fig. 3. Glucose induced soil priming effects (PE) in unamended (grey bar) and biochar amended soil (black) from soil A with low at 100 mg C kg⁻¹ soil (a) and high concentration of glucose addition at 1000 mg C kg⁻¹ soil (b), and soil B with low (c) and high (d) concentration of glucose addition. Error bars indicate standard errors of the means (n = 3).
incubation period. The combined addition of biochar and GL caused 169.0 μg CO₂-C g⁻¹ soil organic C mineralization while GL alone produced 160.9 μg CO₂-C g⁻¹ of soil primed CO₂ during the 28 days of incubation (Fig. 3a).

Glucose alone also caused biochar priming effects. GL gave an additional C₄ biochar loss of 270.5 μg CO₂-C g⁻¹ soil (Soil A) and 300.9 μg CO₂-C g⁻¹ soil (Soil B), while GH caused C₄ biochar losses of 539.8 μg CO₂-C g⁻¹ soil (Soil A) and 482.6 μg CO₂-C g⁻¹ soil (Soil B), respectively (Fig. 4).

4. Discussion

4.1. Approaches to partition three C sources in a biochar rich soil system

A three-source-partitioning approach of combining ¹⁴C labelling with ¹³C natural abundance has been adopted in some studies to partition three C sources in soil systems (Blagodatskaya et al., 2011). This yielded advantages. Compared with the natural abundance of ¹³C, making the analyses difficult, the use of highly labelled ¹⁴C substrates e.g. glucose-C permits the precise and accurate estimation of the different soil pools which contribute to soil CO₂ fluxes (Blagodatskaya et al., 2011; Whitman and Lehmann, 2015). One less treatment is necessary to partition sources, compare to the usage of dual treatments with different isotope ratios for the same component (Whitman and Lehmann, 2015). Finally, the ¹⁴C substrates permit direct estimation of interactions between the soil C pools, without complicated 3-way subtraction (Blagodatskaya et al., 2011). Our results demonstrated this approach by partitioning soil CO₂ emissions derived from soil organic C, added biochar and substrate (glucose in this study). This helps us to better understand the soil priming effects following substrate addition to biochar enriched soils.

4.2. Priming effects in biochar free and enriched soil

A ‘real’ priming effect reflects an increase in mineralization of soil organic C whereas an ‘apparent’ priming effect is due to the increased turnover of microbial C (Blagodatskaya and Kuyzakov, 2008). In this study, the ‘apparent’ priming effect caused by C pool substitution could be largely excluded as the microbial biomass C concentrations (unpublished data) were less than the primed CO₂ in all of the treatments (Fig. 3).

Previous studies of biochar induced priming effects in unpertrubed soils relied on evaluations in the short term (two weeks) (Jones et al., 2011a; Luo et al., 2011), medium term (half a year) (Zimmerman et al., 2011) and long term (8 years) (Kuyzakov et al., 2014). Based on a review study of the 650 data points from 18 studies of biochar induced priming effects, the most positive priming effect occurred in the first 20 days while negative priming appeared in a later stage (Maestrini et al., 2014b). This might indicate that the initial loss of primed C following biochar incorporation would be compensated by greater C stabilization in the long term. This, however, does not reflect the situation in natural soil ecosystems, as substrates are ubiquitous in all soils under agricultural, grassland and forest managements. Our present work on biochar-induced priming effects considers this process following the incorporation of substrates in biochar enriched soil. We used glucose in this study. However, future study should focus more on naturally occurring substrates e.g. plant residues or root exudates. We found that the added glucose gave higher priming effects (53% and 140% more unlabeled C evolved) from two soils which had previously received biochar, compared to biochar free soil (Fig. 3). In another recent study, the combined addition of maize derived biochar and maize feedstock caused 430 μg CO₂-C g⁻¹ soil primed CO₂ during 180 days of incubation (Kerre et al., 2016). Whitman and Lehmann (2015) also found 23% higher soil organic C mineralization induced by roots when biochar was present. However, Whitman et al. (2014) found that biochar additions counteracted positive priming effect induced by maize residues, eliminating net C losses by decreasing soil organic C decomposition (48% lower). This might be due to the limited amount of C released from roots into the rhizosphere (Wichern et al., 2007). Similarly, in our study, a
4.3 Mechanisms involved in substrate induced priming effects in biochar enriched soil

The presence of living plants or addition of organic substances with various composition e.g. cellulose (Fontaine et al., 2011), and sugar-cane sucrose (Nottingham et al., 2009) alter the soil physical, hydrological and chemical environment and associated biological processes thus altering soil organic C mineralization rates (Blagodatskaya and Kuzymakov, 2008). Compared with the substrates used in the above studies, biochar induces larger and longer significant changes in the physio-chemical characteristics of soil (e.g. soil pH, bulk density, porosity and moisture content) due to its high stability (Chan et al., 2007; Sohi et al., 2010). Thus, in two C source systems, the short term priming effects following biochar addition are mainly due to labile C, while positive priming effects (of varying duration) in three C source systems can be explained by both increases in labile C and changes in soil physical and chemical properties, which might further activate certain groups in the microbial biomass (Fig. 5).

A fresh substrate amendment can induce a shift of the microbial community towards the utilization of more easily decomposable C (Blagodatskaya and Kuzymakov, 2008). Microbial colonization within biochar was observed in several studies, and the most likely area for colonization is referred as the “charsphere”, defined as the interface between biochar and soil that provides a unique and preferred micro-environment for microbial colonization (Luo et al., 2013). Glucose added in our study was soluble and likely interacted with microorganisms in the charsphere, which would be expected to support greater amounts of biomass, higher activity and cause a faster turnover of soil organic C close to that specific area. This notion may be partially substantiated by the faster response of bacteria following biochar application in the early stage and fungal dominance in the later stage (unpublished data). Fresh substrate alters the bacterial and fungal community structure during the incubation period in a “microbial succession”, suggesting that the ecological functions of the bacterial and fungal community changes as soil conditions (mainly soil labile C and physical properties) change over time (Andrews and Harris, 1986; Blagodatskaya et al., 2007). We believe that succession from r-strategists to K-strategists occurred during the 28 days of incubation, as the growth of microorganisms becomes increasingly limited by resource deficiency (most glucose is decomposed in the first 1–3 days). K-strategists, instead of r-strategists, are better able to use other sources of C to produce biomass than r-strategists (Andrews and Harris, 1986; Blagodatskaya et al., 2007; Chen et al., 2016). The microbial succession can be more profound in biochar amended soils as the effects of biochar on soil properties are relatively long term. This can largely explain our results, showing that larger priming effects occurred in the later stages of incubation in biochar amended soil rather than in unamended soil (Fig. 3). However, this needs further research using modern approaches, e.g. Illumina MiSeq sequencing.

4.4. Environmental implications

Soil amendment with biochar will increase total soil organic C and might help offset climate change (Sohi, 2012). Biochar, however, can cause priming effects, which may partially offset its effects on C sequestration (Jones et al., 2011b). In many laboratory incubations, biochar induced priming effects ceased after a short period of incubation, and it is generally concluded that short term priming C losses due to biochar are negligible compared to the extra C incorporated into soil following biochar incorporation. However, this input of biochar into soil provides additional surface area and, therefore, an important new microbiological niche (Noyce et al., 2016) where decomposers are concurrently mineralizing soil organic C and other C sources, e.g. plant residues. Thus, with these fresh organic matter inputs and their repeated additions in a biochar-soil-plant system, we believe that the interactions between non-living and living organic C are more likely to occur and cause faster rates of soil organic C decomposition in the charsphere (Luo et al., 2013). If our results are validated in other systems with different soils, biochars and substrates, this C loss cannot be ignored and the C balance might need to be carefully recalculated in the long term by considering multiple labile organic C sources, e.g. root exudates, plant and animal residues etc., which commonly exist in nature (Zhu et al., 2014). Currently, we still do not know the magnitude of priming effects from substrates with different chemical compositions. We believe this needs further systematic investigation. In natural systems where pulses of substrate are usual, e.g. inputs of litter and root exudates in most terrestrial ecosystems (Qiao et al., 2014), there will be a continuous priming of soil organic C mineralization. Since this ongoing process has the potential to stimulate more soil organic C mineralization than one-time substrate inputs (Hamer and Marschner, 2005), we believe the C losses will be larger under field conditions than thus far estimated from controlled laboratory studies.

5. Conclusions

We tested the possibility of adopting the three-source-partitioning approach by combining 14C labelling with 13C natural abundance in a biochar amended soil system. For the first time we were able to separate CO2 emission from biochar, soil organic C and substrate (glucose) based on a combination of 14C with 13C addition to soil. We verified our hypothesis that soil organic C mineralization
responds differently to the addition of fresh organic C in un-amended and biochar amended soils. This study found that following 14C labelled glucose addition, the accumulated primed soil C loss during 28 days were 140% and 53% higher than the priming soil CO2 in biochar free soils. These results help us to better understand the soil priming effects following substrate addition in biochar amended soils. These posited effects need to be confirmed in long term field experiments with different soil and biochar types. As well, some methodological problems remained to be overcome, for example, how to deploy C source partitioning methods under field conditions, considering the safety issues associated with 14C use in natural environments and the high cost of generating enough 14C labelled plant residue for field experiments.

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