Interaction of Biochar Amendment and Nitrogen Deposition on Soil Microbial Biomass Carbon and Enzyme Activity in a Torreya grandis Orchard

Bingwen Chai, Jianhua Lv, Quan Li, Jiasheng Wu*, Xinzhang Song**

State Key Laboratory of Subtropical Silviculture, Zhejiang A&F University, Hangzhou, China

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Abstract

The objective of this study was to test whether biochar amendment could offset the effects of nitrogen (N) deposition on soil microbial biomass carbon (MBC) and enzyme activity. We applied N (low and high rates) and biochar (low and high rates) individually, and in combination (all permutations) to the soil of a Torreya grandis cv. “Merrillii” orchard as a 13-month field experiment during 2015-2016 in Zhejiang Province, China. MBC significantly increased in the low N treatment but decreased in the high N treatment ($P<0.05$). MBC significantly decreased in both biochar-only treatments, and this effect became stronger as biochar amendment rates increased. Biochar amendment amplified the positive effects of low N treatment on MBC and mitigated the negative effects of high N treatment. Catalase, cellulase, and urease activities significantly increased with N addition. Cellulase, nitrite reductase, and urease activities significantly increased in the biochar treatments. The positive effects of low N addition on catalase were amplified in the low biochar amendment treatment, but the positive effects on cellulase decreased significantly. The effects of biochar amendment on MBC and enzyme activities of soil that receives atmospheric N deposition were regulated by the rates of both biochar amendment and N deposition, and varied with enzyme type.

Keywords: biochar, microorganism, nitrogen addition, orchard, Torreya grandis cv. “Merrillii”

Introduction

In recent decades, nitrogen (N) deposition has rapidly increased owing to increases in the burning of fossil fuel and the utilization of fertilizers in agriculture [1]. This phenomenon has strongly impacted subtropical China, which has incurred a maximum annual N deposition rate of 63.53 kg N ha$^{-1}$yr$^{-1}$ and is predicted to become the region with the highest N deposition in the world by 2030 [2]. Soil microbial biomass plays a critical role in soil fertility, and microbial biomass carbon (MBC) can be effectively used as an index to evaluate soil quality [3]. Some studies have found that N input significantly decreases MBC.
Biochar is a byproduct of the thermal combustion (termed pyrolysis) of biomass in the absence of oxygen at relatively low temperatures (300-700°C) [11]. It is thought to be intrinsically resistant to microbial decomposition owing to its conditioned poly-aromatic structures [12]. It is commonly alkaline and is therefore usually applied to neutralize acidic soils and to increase soil pH [13]. Previous studies have shown that biochar amendment increased soil SBC in the plains of northern China during the maize-growing season [14]; in the hilly red soil region of southern China, which has soils that are generally low in fertility [15], it increased microbial activity in highly weathered soils [16]. However, a meta-analysis of studies on this topic showed that a high rate of biochar amendment has a negative effect on MBC [17]. Feng et al. [18] observed that biochar amendment inhibits soil catalase and neutralizes phosphatase activities in potted wheat. However, it is still unknown whether biochar amendment can neutralize the effects of atmospheric N deposition on soil biotic properties due to its alkaline character – especially in orchard environments.

Torreya grandis cv. “Merrillii” is a coniferous tree in the Cephalotaxaceae family that yields a rare and unique dried fruit. The fruit is easily digestible, and has many beneficial effects on human health [19]. Due to its high nutritional value, the planting area of T. grandis has been expanded rapidly, and it therefore has many beneficial effects on human health [19].

Material and Methods

Study Site

The study site was located in the town of Yuqian within Lin’an City, Zhejiang Province, China (30°14′ N, 119°42′ E). The climate at the study site is subtropical monsoonal with a mean annual precipitation of 1613.9 mm. The mean annual temperature is 16°C with a mean annual low of 4.5°C (January) and a mean annual high of 29°C (July). The area has 237 frost-free days a year. The soil is categorized as belonging to the yellow-red soil class (Chinese system of soil classification, which is equivalent to the Hapludult soil class in soil taxonomy).

Experimental Design

The T. grandis orchard was established in 1986. In April 2015, 27 T. grandis trees with similar height, basal diameter, and canopy breadth were selected for use in the study, and a corresponding 4 × 4 m plot was established around each sample tree. The initial stand and soil characteristics of the orchard are summarized in Table 1. The experiments were conducted following published methods, with some modifications, for simulating N deposition [20, 21] and determining local N deposition rates (30.9 kg N ha⁻¹·yr⁻¹) [22, 23]; the low-N treatments used 30 kg N ha⁻¹·yr⁻¹ (N30), and the high N treatments used 60 kg N ha⁻¹·yr⁻¹ (N60). Biochar amendment treatments were 20 tonnes·ha⁻¹ (BC20) and 40 tonnes·ha⁻¹ (BC40) treatments. The four combined treatments consisted of all permutations of the two N and two biochar treatments (N30+BC20, N30+BC40, N60+BC20, N60+BC40), and a control (without N or biochar addition) was also implemented. There were three replicate plots per treatment.

In May 2015, biochar was added to the soil, and the top 30 cm was mixed uniformly by manual ploughing [24]. Nitrogen addition also began in May 2015. NH₄⁺ and NO₃⁻ were reported to account for 56.1% and 43.9%, respectively, of local wet nitrogen deposition. NH₄NO₃ was used as the N source to simulate atmospheric N deposition, as it is considered to be the closest available to the chemistry of atmospheric N deposition [21]. Quantitative NH₄NO₃ was dissolved in water and evenly sprayed from the top of the T. grandis canopy using an electric sprayer once per month onto the plots that received the N treatment. The plots that did not receive N were sprayed with an equal amount of N-free water.

Biochar Characteristics

The biochar used in this study was produced by Sanli New Energy Company (Shangqiu, China) from wheat straw at a temperature of about 450°C under anoxic conditions. The original biochar mass was ground so that it could pass through a 2 mm sieve, and then mixed thoroughly to obtain a fine granular consistency. The characteristics of the biochar obtained were as follows: pH (H₂O): 9.8, bulk density: 0.5 g·cm⁻³, surface area: 9.7 m²·g⁻¹, cation exchange capacity (CEC): 189.3 c·mol·kg⁻¹, organic carbon content: 425.3 g·kg⁻¹, total N content: 5.2 g·kg⁻¹, total phosphorus (P) content: 3.4 g·kg⁻¹, and ash content: 18.6%.
Soil Sampling

In June 2016, three soil samples were randomly collected as soil cores from each plot. Samples were taken from the top 20 cm of the soil, 80 cm away from the tree trunk. The samples were mixed until homogenous, and then transported to the laboratory in an incubator. The samples were sieved through 2 mm mesh to remove plant residues, stones, and roots. Part of the fresh sample was then used to determine the MBC. The remaining sample was then used to measure enzyme activity and the physicochemical properties of the soil [25].

Analysis of Soil Microbial Biomass and Soil Physicochemical Properties

MBC was measured using the chloroform fumigation extraction method. Soil organic matter was estimated using dichromate oxidation method. Total nitrogen was determined using an automatic Kjeldahl distillation-titration unit (Foss, Hillerød, Denmark); available nitrogen was determined by titration against a standard 0.01 mol L\(^{-1}\) H\(_2\)SO\(_4\) solution. Total phosphorus and available phosphorus content were measured by the molybdenum blue method, using a spectrophotometer (UV2550, Shimadzu, Kyoto, Japan). Total potassium was determined by NaOH fusion method. The available potassium was measured using the 2 mol·L\(^{-1}\) cold HNO\(_3\) method. The pH was determined in a soil-water extract (1:2.5 w/v) using a pH meter (FE20, Mettler Toledo, Switzerland) after shaking for 30 minutes. The activities of soil enzymes were estimated following the methods suggested by Guan [26].

Statistical Analyses

A one-way analysis of variance (ANOVA) was used to determine whether statistically significant differences existed between MBC or enzyme activities between the treatments. Pairwise comparisons were then conducted to identify which of the treatments were significantly different using a least-significant difference (LSD) test. Two-way ANOVAs were performed to assess the combined effects of N deposition and biochar amendment on MBC and enzyme activity. Analyses were conducted using SPSS (Statistical Package for the Social Sciences) version 18.0 for Windows (SPSS Inc., Chicago, Illinois, USA).

Results and Discussion

Data Analysis of Soil Microbial Biomass Carbon and Enzyme Activity

The soil MBC was 7%, 65%, and 33% lower in the BC20, BC40, and N60 treatments, respectively, than in the control; however, soil MBC was 31%
higher in the N30 treatment than in the control (Fig. 1). Soil MBC significantly increased in the N30+BC20 treatment but significantly decreased in the N30+BC40 treatment, compared with the soil MBC in the N30 treatment (Fig. 1). The soil MBC also significantly increased in the N60+BC20 and N60+BC40 treatments compared with the soil MBC in the N60 treatment (Fig. 1). A two-way ANOVA showed that N deposition, biochar amendment, and their interaction all significantly affected soil MBC (Table 2).

In general, catalase activity was significantly higher in the N30, N30+BC20, and N60+BC20 treatments than in the control. Catalase activity significantly increased in the N30 plots with BC20 amendment, compared with that of the N30 plots (Fig. 2a). A two-way ANOVA showed that N and biochar addition individually significantly affected catalase activity, but that their interaction was not significant (Table 2).

Cellulase activity was significantly higher in the BC20, N30, N30+BC20, N60, and N60+BC40 treatments than in the control, but it decreased significantly in the N30+BC40 treatment (Fig. 2b). A two-way ANOVA also showed that the interaction of N and biochar

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Table 2. Results of a two-way ANOVA of the effects of nitrogen deposition and biochar amendment on soil microbial biomass carbon (MBC) and enzyme activities in a *Torreya grandis* orchard.

<table>
<thead>
<tr>
<th>Category</th>
<th>Difference source</th>
<th>SS</th>
<th>F</th>
<th>P-value</th>
</tr>
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<td>MBC</td>
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<td>Interaction</td>
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<td>β-fructofuranosidase</td>
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significantly affected cellulase activity, although N and biochar addition individually did not (Table 2).

Nitrate reductase activity did not vary significantly under any treatment trialed in this study (Fig. 2c; Table 2). Nitrite reductase activity, however, was significantly higher in the BC20, N30+BC20, and N60+BC40 treatments than in the control. Nitrite reductase activity significantly decreased in the N30+BC40 and N60+BC20 treatments (Fig. 2d). A two-way ANOVA showed that biochar amendment (both by itself and combined with N addition) significantly affected nitrite reductase activity (Table 2).

Urease activity was significantly higher in the BC40 and N60 treatments, and their interaction, than in the control (Fig. 2e). In addition, a two-way ANOVA showed that N and biochar addition significantly affected urease activity individually, but that their interaction was not significant (Table 2).

Finally, the activity of β-fructofuranosidase was significantly higher in the N60+BC20 and N60+BC40 treatments...
treatments than in the control (Fig. 2f). A two-way ANOVA showed that N deposition, biochar amendment, and their interaction all significantly affected β-fructofuranosidase activity (Table 2).

Our results showed that soil MBC significantly increased in N30 plots, but significantly decreased in N60 plots. This partially supports our first hypothesis, which was that N deposition increases soil MBC. It is generally known that N addition can alter microbial biomass and activity in several ways [4, 7]. Li et al. [27] found that N deposition (30, 60, or 90 kg N ha⁻¹ yr⁻¹) significantly increased soil MBC in Moso bamboo (Phyllostachys edulis) plantations. However, a meta-analysis conducted by Treseder [6] showed that chronic N deposition significantly decreases MBC. Excessive N deposition can decrease the C:N ratio of the soil organic matter, pH, and soil organic C that are the energy sources for soil microorganisms [28], which leads to the decline in MBC.

Most enzyme activities were higher in the N treatments than they were in the control. The exception was nitrate reductase, which was lower in the N30 treatment than the control (Fig. 2), indicating that N addition significantly increased enzyme activity. These results also partially support our first hypothesis: that N deposition increases enzyme activities. Catalase is one of the major lignin-degrading enzymes found in the soil [29]. Waldrop et al. [30] found that catalase activity increased after one year of N addition in sugar maple (Acer saccharum Marshall) forests in northern Lower Michigan, USA. This is consistent with the results of the present study, suggesting that N deposition is beneficial for the promotion of lignin degradation and litter decomposition, and can accelerate nutrient cycling in T. grandis orchards. However, Deforest et al. [31] observed that catalase activity declines after nine years of N addition (30 kg N ha⁻¹ yr⁻¹) in a mature northern hardwood forest, and other studies have not found any significant effects [32]. Chronic N addition can reduce catechol and vanillin degradation, lignocellulose depolymerization, and soil microbial metabolism capacity; this may potentially explain why catalase activity was observed to decrease. These contradictory results imply that the effect of N deposition on catalase activity can vary with the timing of N deposition events.

Cellulase is produced by a great number of bacterial and fungal species [29]. A previous study found that the addition of N for seven years generally stimulated cellulase activity in eight forests and grassland sites with sandy, well drained, and poorly developed soils in Minnesota, USA [33], which is consistent with the results of the present study (Fig. 2b). In contrast, Liu et al. [34] observed that two levels of N addition (100 and 150 kg N ha⁻¹ yr⁻¹) significantly decreased cellulase activity in natural Pinus tabuliformis Carrière forests on Taiyue Mountain, China. Cellulase activity is sensitive to alterations in soil pH [35]. Excessive N input can decrease soil pH (Fig. 3) and inhibit litter decomposition – especially the numbers and activity of microorganisms [36], leading to a decline in cellulase activity.

Nitrification play important roles in soil denitrification, which are closely related to NO₃⁻-N concentrations [29]. In the present study, N deposition did not significantly affect the activities of these two enzymes (Fig. 2c, d; Table 2). Nonetheless, additional study is needed to fully understand these effects.

Urease is a type of N-acquiring enzyme [37]. In the present study, N addition significantly increased urease activity (Fig. 2e; Table 2). Similar results were also obtained by Saiya-Cork et al. and Guan et al. [37, 38]. Saiya-Cork et al. [37] observed an increase in urease activity that suggested the potential for an increase in gross N mineralization rates during a study conducted in a sugar maple-dominated forest with simulated N (30 kg N ha⁻¹ yr⁻¹) deposition in northern Michigan, USA in the 1998-2000 growing seasons [37]. N deposition may improve the effectiveness of nitrogen, which strongly stimulates the transformation of urease activity. However, Ajwa et al. [39] observed that urease activity significantly decreased with N addition (100 kg N ha⁻¹ yr⁻¹) to Irwin silty clay loam soil in a tall grass prairie ecosystem in Manhattan, USA, which is inconsistent with the results of the present study. This apparent contradiction is perhaps due to the high rate of N addition used in Ajwa et al. [39], who induced a negative effect on urease activity.

β-Fructofuranosidase serves not only as an indicator of the release of low molecular weight sugars, but also as an energy source for microorganisms [40]. Tu et al. [41] observed that β-fructofuranosidase activity significantly increased after six months of N addition in a Pleioblastus amarus (Keng) Keng f. plantation in a high-rainfall area of western China, which is inconsistent with the results of the present study (Fig. 2f). This is partly due to the large amount of fresh
litter that contained soluble organic matter and cellulose present in the *Pleioleptus amarus* plantation, which increased β-fructofuranosidase activity.

**Effects of Biochar on Soil Microbial Biomass Carbon and Enzyme Activity**

MBC significantly decreased with an increasing rate of biochar amendment (Fig. 1), which supports our second hypothesis, which was that biochar amendment decreases soil MBC and enzyme activities. Similarly, Huang et al. [42] observed that a high rate of biochar amendment (200 g·kg⁻¹) over the course of one year resulted in a decrease in MBC in high-fertility soil. A possible reason for this may be that biochar can adsorb low molecular weight organic matter, which can reduce soil MBC. The high C:N ratio that can result from an excessive biochar amendment may cause soil microbial nitrogen immobilization, resulting in low microbial activity [43, 44]. Dempster et al. [45] found that soil MBC decreased with increasing rates of the addition of jarrah (*Eucalyptus marginata* Donn ex Sm.) wood biochar; one possible explanation for this observation is that the biochar input decreased the mineralization rate of the soil organic carbon. In addition, other studies have reported that soil organic carbon showed a significant positive correlation with soil MBC [46, 47]. Li et al. [27] showed that high rates of biochar amendment (40 kg·ha⁻¹) significantly decreased soil MBC, but that moderate rates of biochar amendment (20 kg·ha⁻¹) significantly increased soil MBC. The effects of biochar on soil microbes are controlled by multiple physicochemical and environmental factors.

Biochar amendment significantly increased cellulase, nitrite reductase, and urease activity, but had no significant effect on the other three enzymes examined in this study (Fig. 2). This contradicts our second hypothesis, that biochar amendment decreases enzyme activities. Catalase is a type of oxidoreductase, which indicates soil metabolic capacity [29]. In the present study, biochar amendment had no effect on catalase activity (Fig. 2a). Masto et al. [48] demonstrated that catalase activity significantly increased with increasing rates of biochar amendment (0, 1, 3, 5, 10 and 20 g·kg⁻¹) produced from water hyacinth in a red soil in India. In addition, catalase activity significantly increased with intermediate (10 g·kg⁻¹) or low (5 g·kg⁻¹) rates of biochar amendment, when compared with a high (50 g·kg⁻¹) rate. An excessive rate of biochar amendment may inhibit enzyme catalysis binding sites via the absorption of enzyme molecules, which inhibits catalase activity [11].

Biochar is generally an alkaline material that can raise soil pH [35] (Fig. 3). In this study, it significantly increased cellulase activity (Fig. 2b). Similar results were observed by Bailey et al., Baminger et al., and Paz-Ferreiro et al. [40, 49, 50]. However, Demisie et al. [51] found that three rates (5 g·kg⁻¹, 10 g·kg⁻¹, and 20 g·kg⁻¹) of two kinds of biochar amendment (produced from oak wood and bamboo) had no effect on cellulase activity in a highly weathered, erosion-susceptible Chinese red soil.

In the present study, biochar amendment had no effect on nitrate reductase activity but significantly increased nitrite reductase activity (Fig. 2c, d; Table 2). In contrast, Zhang et al. [52] observed that the addition of biochar produced from apple tree branches increased nitrate reductase activity, but did not affect nitrite reductase, in the soil of a winter wheat system. These differences may be due to differences in biochar and ecosystem types.

Urease is sensitive to land use and agricultural management practices [40]. Previous studies [40, 51, 53, 54] have shown that biochar amendment has a positive effect on urease activity, which is consistent with our findings (Fig. 2e; Table 2). However, Gu et al. [55] found that biochar amendment produced from wheat straw significantly decreased urease activity in a grey desert soil, but did not decrease the activity of urease in an aeolian sandy soil. Wu et al. [56] observed that urease activity decreased with increasing rates (0, 9.9, and 24.4 g·kg⁻¹) of wheat straw biochar amendment in a chernozemic soil. The differences in these results imply that the effects of biochar on urease activity are complex.

β-Fructofuranosidase absorbs, utilizes, and provides energy for microorganisms via sucrose hydrolysis. The activity of this enzyme reflects its utilization of easily soluble substances, and the accumulation and transformation of microorganisms in the soil. In the present study, the high biochar amendment rate (40 g·kg⁻¹) significantly increased the activity of β-fructofuranosidase, but the low biochar amendment rate did not (Fig. 2f). Similarly, Zou et al. [54] observed that a high biochar amendment rate (50 g·kg⁻¹) had a positive effect on the activity of β-fructofuranosidase in the soil around cucumber roots.

**Interactive Effects of N Deposition and Biochar on Soil Microbial Biomass Carbon and Enzyme Activity**

Our results indicate that the low rate of biochar (20 tonnes·ha⁻¹) amendment of the N30 plots amplified the positive effect of N deposition on the MBC (Fig. 1). In contrast, the high rate of biochar (40 tonnes·ha⁻¹) amendment of the N30 plots inhibited this positive effect (Fig. 1). In the N60 plots, biochar amendment significantly alleviated the negative effect of N deposition on the soil MBC, and even transformed it to a positive effect (Fig. 1). This pattern partly supports the third hypothesis – that biochar amendment offsets the negative effect of N deposition – and indicates that biochar amendment can change the effects of N deposition on enzyme activity. The results of the present study also suggest that soil pH affected MBC. Low biochar amendment neutralized the acidity of
the soil in the N30 plots, and therefore increased the soil MBC (Fig. 1). However, high pH values induced by high rates of biochar amendment (40 tonnes·ha$^{-1}$) resulted in a negative effect on soil MBC (Figs 1, 3). In the N60 plots, excessive N deposition largely decreased soil pH, while biochar amendment significantly increased the pH of acidic soils (Fig. 3) and therefore significantly increased soil MBC (Fig. 1). Biochar amendment was also observed to reverse the reduction in the soil MBC caused by the high rate of N addition (Fig. 1). One reason for this may be that the physicochemical properties of biochar, for example its alkaline pH, could not only improve soil pH, but also the water-holding capacity and CEC of the soil. Biochar can create a favorable habitat for microbial colonization due to its large surface area and porous structure [57]. Our findings suggest that the direction and magnitude of variation of soil MBC depends on the rates of both biochar and N addition.

Nitrogen addition significantly increased catalase and cellulase activities. A low rate of biochar amendment significantly amplified the positive effects of the low rate of N addition on catalase, but significantly offset the positive effects on cellulase. Biochar amendment of the N60 plots significantly increased β-fructofuranosidase activity, although N addition did not have a significant effect. Our findings indicate that the effects of biochar amendment of soil receiving atmospheric N deposition were regulated by the rates of both biochar amendment and N deposition, and varied with enzyme type.

Conclusions

Overall, low N deposition (30 kg N ha$^{-1}$·yr$^{-1}$) increased soil MBC, whereas high N deposition (60 kg N ha$^{-1}$·yr$^{-1}$) decreased it. The addition of N increased enzyme activity with the exception of nitrate reductase activity, which decreased slightly in the N30 plots. Biochar amendment significantly decreased MBC, but increased enzyme activity with the exception of nitrate reductase activity that was observed to decline after biochar amendment. When combined with the addition of N, biochar amendment could significantly alter the effects of N deposition on MBC and enzyme activity. Our findings indicated that N deposition and biochar amendment, both individually and combined, significantly affected soil MBC and enzyme activity in a T. grandis orchard. The direction and magnitude of these effects depended on the rates of both biochar amendment and N addition.

Acknowledgements

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Conflict of Interest

The authors have not declared any conflict of interest.

References

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33. KEELER B.L., HOBBIE S.E., KELLOGG I.E. Effects of long-term nitrogen addition on microbial enzyme activity in eight forested and grassland sites: Implications for litter and soil organic matter decomposition. Ecosystems 12, 1, 2009.
42. HUANG C., LIU L., ZHANG M. Effects of biochar on properties of red soil and ryegrass growth. J. Zhejiang Univ. 37, 439, 2011.
55. GU M., XU W., TANG G., GE C., MA H. Effects of biochar on soil microbial diversity and function related with N transformation in grey desert soil and aeolian sandy soil in Xinjiang. Xinjiang Agricultural Sciences 51, 926, 2014.