

## The response to comments

Dear Dr. Denise M. Akob

We are grateful for your comments and we agree to revise the manuscript in response to the comments. Please find our response to the comments on behalf of all co-authors. Our response to the reviewer's comments are detailed point by point below.

Abstract: there is no mention of slash pine plantation but it is part of the title. Please revise to include and to link your results back to the habitat or if the habitat is not the key part of the story then revise the title.

Response: We accepted your suggestion. And we remove the “slash pine” from the title.

L. 33: change to hydrolysis. The correct term is hydrolysis and cannot be plural

Response: Revised as recommended, please refer to Line 34.

L. 50: change to “is mostly comprised of”

Response: Revised as recommended, please refer to Line 51.

L. 69: define the BG and NAG acronyms

Response: BG is  $\beta$ -1,4-glucosidase, and NAG is  $\beta$ -1,4-N-acetylglucosaminidase. We add the full names to the Line 70. And at the same time, we used the abbreviations of  $\beta$ G and NAG at Line 97-98.

L. 106: “in line with the economic theory” does not fit here. No economic theory has been presented. I would omit it and start with “Microorganisms...” or try to introduce this better

Response: We accepted your suggestion. The sentence has been revised as 'Microorganisms will allocate energy to the relatively absent resources, so that N additions will cause C and P-acquisition enzymes to increase, and N-acquisition enzymes to decrease (Burns et al., 2013)'. (Line 108-110).

L. 114-116: add a reference

Response: The reference of ‘Sinsabaugh et al., 2002’ has been added (Line 119).

L. 134: change to “C-“

Response: Revised as recommended, please refer to Line 137.

L. 135: change to “;”

Response: Revised as recommended, please refer to Line 138.

L. 142: there are so many acronyms in the paper that I would not abbreviate the experimental station, just write it out.

Response: The ‘QYZ’ has been revised as Qianyanzhou (Line 145).

L. 149: define a.s.l.

Response: a.s.l means above sea level (Line 146).

L. 153-165: this is still not clear. How many test plots were there, first you say one control and one test then you state there are 9? Start with how they were divided up then discuss the treatments. On l. 157, it reads as if you added both NH<sub>4</sub> and NO<sub>3</sub> to both test plots. So, how were you comparing effects of each N source? It might be useful to add a schematic to supplemental information. For the application, how much time was there between sprayings? It was only 1 day per month when N was applied?

Response: We have revised the sentence as ‘Nine 20 × 20 m plots were established at the experimental sites, including a control, ammonium only and nitrate only additions plots with three replicates (3 treatments × 3 replicates)’. Furthermore, a schematic was added as supplemental figure S2. The N were applied on a non-rainy day at the interval of about one month, so the N was applied one day per month (Line 168-169).

L. 174: I would not use synthetic here. I suggest changing to seasonal.

Response: Revised ‘synthetic’ to ‘seasonal’ (Line 178).

L. 183-189: was the soil:water shaken before measurement of pH and N? In this section, either provide details on the methods or provide references.

Response: The homogenate was stirring by glass rod for one minute and then was settled for 30 min before measurement of soil pH. And the soil-water mixture was shaken for 2 h before measurement of soil ammonium, nitrate, soil dissolved organic carbon. We have added the references [Bao. (2010)] in the manuscript (Line 187)

L. 187: soil cannot be extracted with soil, revise

Response: The sentence has been revised as ‘Soil DOC was extracted with distilled water at a ratio of 1 g soil : 5 ml water, and was measured with an organic element analyzer (Liquid TOCII, Elementar, Germany)’ (Line 192-195)

L. 205: what concentration of sodium acetate?

Response: The concentration of sodium acetate was 50 mmol L<sup>-1</sup> (Line 212).

L. 241: specify panel A, e.g., Fig 1A

Response: We have specified the panel (a) to (f) for Figure 1-6. Accordingly, we have revised in the results and discussion.

L. 241-242: incorrect usage of “respectively”. Please always refer to the specific figure panels you are referencing.

Response: We have revised the sentence as “The soil nitrate contents were 165% and 129% higher (Fig. 2b), and the soil ammonium contents were 31% and 38% lower in the ammonium and nitrate treatments (Fig. 1b) than in the control for the three sampling events.” (Line 248-251).

L. 258: you are missing an “and”

Response: We added it to Line 265.

L. 264: I caution against using the term “shifted” when you are not showing or referring to time series data.

Response: We have revised the sentence as “The microbial communities was dominated by G<sup>+</sup> in the ammonium-treated plots, meaning that the G<sup>+</sup>/G<sup>-</sup> ratios were higher in the ammonium-treated plots than in the control or nitrate-treated plots.” (Line 272-274)

L. 273-283, Table 2 and elsewhere: are the enzymes supposed to be named with Greek letters? The naming is inconsistent with the intro and methods. Please verify and correct throughout the paper.

Response: We revised aG, BG, BX to  $\alpha$ G,  $\beta$ G,  $\beta$ X throughout the manuscript, the Tables and the Figures.

L. 292-294: this sentence is strangely worded—it is unclear what you are referring to with use of respectively twice in this sentence. What are the values respective of?

Response: We revised the sentence to “The results of RDA between soil properties and absolute enzyme activities showed that the first axis explained 72.0% of the variability (Fig. 6a), while the results of RDA between soil properties and microbial community structures showed that the first axis explained 67.5% of the variability (Fig. 6b).” (Line 301-308)

L. 292-303: refer to your table and figure earlier in the paragraph

Response: We added “Fig.6a” to Line 302, and added “Fig. 6b” to Line 303, and added “Fig.6a,b” to Line 311 and Line 317.

L. 312-313: C cannot be included in the sum of ammonium and nitrate, remove the parenthetical statement

Response: We removed the parenthetical statement. Please refer to Line 326-327.

L. 350: only include the reference once.

Response: We removed “(Li et al., 2016)” from Line 365.

L. 356-357: do you mean microbial communities? Microorganism systems is an unusual choice.

Response: The interaction effects between plant, microbes and soil nutrients were variable across the three sampling events. So we changed the “plant-soil-microorganism systems” to “plant-microorganism competitive relationship”, please refer to Line 372.

L. 560: Bacteria is spelled wrong

Response: Revised as recommended. Please refer to Fig.3b

L. 558: soil dissolved organic carbon is misspelled

Response: Fig.2a was revised as recommended.

Fig. 4f: the “Cb” is not easy to read

Response: Fig.4f was revised as recommended. Capital letters represent significant differences between the treatments ( $P < 0.05$ ), and small letters represent significant differences between the sampling time ( $P < 0.05$ ).

Fig. 6: please define the abbreviations in the legend

Response: The full names of the PLFA biomarkers, enzymes and soil properties were shown in Table 1. We add “The abbreviations are the same as Table 1. SOC: soil organic matter; TN: total nitrogen; C/N: the ratio of soil

organic matter to total nitrogen; SWC: soil water contents.' to the legend of Fig.6. At the same time, we omitted the state of the abbreviations shown in the legend of Fig. 3 and 4 that replicated with Table 1.

## **The list of all relevant changes made in the manuscript**

The line number see the marked-up manuscript that was showed below this section.

1. Title: Delete 'in a slash pine plantation'
2. Line 33-34: Change 'hydrolyses' to 'hydrolysis'
3. Line 51: Change 'comprises' to 'is comprised of'
4. Line 70: Change 'BG and NAG' to ' $\beta$ -1,4-glucosidase ( $\beta$ G) and  $\beta$ -1,4-N-acetylglucosaminidase (NAG)'
5. Line 97-98: Change ' $\beta$ -1,4-glucosidase ( $\beta$ G)' and ' $\beta$ -1,4-N-acetylglucosaminidase (NAG)' to ' $\beta$ G' and 'NAG'
6. Line 108-111: Revised as 'Microorganisms will allocate energy to the relatively absent resources so that N additions will cause C and P-acquisition enzymes to increase, and N-acquisition enzymes to decrease (Burns et al., 2013).'
7. Line 119: Add '(Sinsabaugh et al., 2002)'
8. Line 137: Change 'C and P-hydrolase' to 'C- and P-hydrolase'
9. Line 138: Change ',' to ';'.
10. Line 145: delete the abbreviation '(QYZ)'
11. Line 146-147: Change 'a.s.l' to 'above sea level'
12. Line 158-165: Revised as 'Nine 20  $\times$  20 m plots were established at the experimental sites, including a control, ammonium only and nitrate only treatments with three replicates (3 treatments  $\times$  3 replicates).'
13. Line 178: Change 'synthetic' to 'seasonal'

14. Line 187: We added ‘The measurement of soil chemical properties was followed the method of Bao (2010).’
15. Line 272-274: Revised as ‘The microbial communities shifted from G<sup>-</sup> to was dominated by G<sup>+</sup> in the ammonium-treated plots, meaning that the G<sup>+</sup>/G<sup>-</sup> ratios were higher in the ammonium-treated plots than in the control or nitrate-treated plots (Fig. 3d).’
16. Line 301-308: Revised as ‘The results of RDA between soil properties and absolute enzyme activities showed that the first axis explained 72.0% of the variability (Fig. 6a), while the results of RDA between soil properties and microbial community structures showed that the first axis explained 67.5% of the variability (Fig. 6b).’
17. Line 326-327: Delete ‘(the sum of the ammonium and nitrate concentrations contents)’
18. Results and Discussion section: We have specified the panel (a) to (f) for Figure 1-6.
19. We revised  $\alpha$ G,  $\beta$ G,  $\beta$ X to  $\alpha$ G,  $\beta$ G,  $\beta$ X throughout the manuscript, the Tables and the Figures.
20. We added a schematic as supplemental figure S2.

**The marked-up manuscript see below**

1 **Contrasting effects of ammonium and nitrate additions on the biomass of soil microbial**  
2 **communities and enzyme activities ~~in a slash pine plantation~~ in subtropical China**

3 Chuang Zhang<sup>a,b,c</sup>, Xin-Yu Zhang<sup>b,c</sup>, Hong-Tao Zou<sup>a</sup>, Liang Kou<sup>b</sup>, Yang Yang<sup>b,c</sup>, Xue-Fa Wen<sup>b,c</sup>,  
4 Sheng-Gong Li<sup>b,c</sup>, Hui-Min Wang<sup>b,c</sup>, Xiao-Min Sun<sup>b,c</sup>

5  
6 <sup>a</sup>College of Land and Environment, Shenyang Agricultural University, Shenyang 110866, China;

7 <sup>b</sup>Key Laboratory of Ecosystem Network Observation and Modeling, Institute of Geographic  
8 Sciences and Natural Resources Research, Chinese Academy of Sciences, Beijing 100101, China;

9 <sup>c</sup>College of Resources and Environment, University of Chinese Academy of Sciences Beijing,  
10 100190, China

11 Corresponding author: X. Y. Zhang ([zhangxy@igsnrr.ac.cn](mailto:zhangxy@igsnrr.ac.cn)), H.T. Zou  
12 ([zouhongtao2001@163.com](mailto:zouhongtao2001@163.com))

13

14 **Abstract**

15

16 The nitrate to ammonium ratios in nitrogen (N) compounds in wet atmospheric deposits have  
17 increased over the recent past, which is a cause for some concern as the individual effects of nitrate and  
18 ammonium deposition on the biomass of different soil microbial communities and enzyme activities are  
19 still poorly defined. We established a field experiment and applied ammonium (NH<sub>4</sub>Cl) and nitrate  
20 (NaNO<sub>3</sub>) at monthly intervals over a period of four years. We collected soil samples from the  
21 ammonium and nitrate treatments and control plots in three different seasons, namely spring, summer,  
22 and autumn, to evaluate the how the biomass of different soil microbial communities and enzyme  
23 activities responded to the ammonium (NH<sub>4</sub>Cl) and nitrate (NaNO<sub>3</sub>) applications. Our results showed  
24 that the total contents of phospholipid fatty acids (PLFA) decreased by 24% and 11% in the ammonium  
25 and nitrate treatments, respectively. The inhibitory effects of ammonium on gram positive bacteria (G<sup>+</sup>)  
26 and bacteria, fungi, actinomycetes, and arbuscular mycorrhizal fungi (AMF) PLFA contents ranged  
27 from 14% to 40% across the three seasons. We also observed that the absolute activities of C, N, and P  
28 hydrolyses and oxidases were inhibited by ammonium and nitrate, but that nitrate had stronger  
29 inhibitory effects on the activities of acid phosphatase (AP) than ammonium. The activities of  
30 N-acquisition specific enzymes (enzyme activities normalized by total PLFA contents) were about 21%



31 and 43% lower in the ammonium and nitrate treatments than in the control, respectively. However, the  
32 activities of P-acquisition specific enzymes were about 19% higher in the ammonium treatment than in  
33 the control. Using redundancy analysis (RDA), we found that the measured C, N, and P hydrolysis  
34 ~~hydrolyses~~ and polyphenol oxidase (PPO) activities were positively correlated with the soil pH and  
35 ammonium contents, but were negatively correlated with the nitrate contents. The PLFA biomarker  
36 contents were positively correlated with soil pH, soil organic carbon (SOC), and total N contents, but  
37 were negatively correlated with the ammonium contents. The soil enzyme activities varied seasonally,  
38 and were highest in March and lowest in October. In contrast, the contents of the microbial PLFA  
39 biomarkers were higher in October than in March and June. Ammonium may inhibit the contents of  
40 PLFA biomarkers more strongly than nitrate because of acidification. This study has provided useful  
41 information about the effects of ammonium and nitrate on soil microbial communities and enzyme  
42 activities.

43

44

## 45 **1. Introduction**

46

47 Studies have reported increases of 25% in wet atmospheric nitrogen (N) deposition over the past  
48 decade (Jia et al., 2014), which has resulted in a range of problems in forest ecosystems, such as  
49 induced soil acidification, aggravation of cation and nitrate leaching, and decreased microbial biomass  
50 (Liu et al., 2011; Huang et al., 2014; Gao et al., 2015; Liu et al., 2013). While wet atmospheric N  
51 deposition is mostly comprised of ammonium, nitrate deposition has increased over recent years, so  
52 that the ratio of ammonium to nitrate has decreased from 5 to 2 (Liu et al., 2013). It is therefore  
53 important to study the individual influences of these two forms of N on soil microorganisms to support  
54 improved predictions of C, N, and P cycling under increased nitrate deposition.

55 Soil microorganisms supply nutrients to forests by producing enzymes that catalyze the degradation  
56 of soil organic matter, and drive carbon (C), nitrogen (N), and phosphorus (P) cycling, with  
57 consequences for forest productivity and sustainability (Heijden et al., 2008). The soil microbial  
58 biomass of different communities may be quantified by phospholipid fatty acid (PLFA) biomarkers.  
59 Even though the PLFA signature method is not as advanced as genomic technology, it has been used  
60 extensively with good results to analyze the biomass and structures of microbial communities

61 (Frostegeård et al., 2011). Bacteria, including gram positive ( $G^+$ ) and negative ( $G^-$ ) bacteria, generally  
62 degrade labile compounds by excreting hydrolase, while fungi, including arbuscular mycorrhizal fungi  
63 (AMF) and saprophytes (SAP), are responsible for degrading recalcitrant compounds by secreting  
64 oxidase (Burns et al., 2013; Sinsabaugh et al., 2010; Willers et al., 2015).

65 To date, most studies have considered the influence of organic N on microbial communities (Guo et  
66 al., 2010; Hobbie et al., 2012) and few studies have reported how ammonium and nitrate individually  
67 influence microbial communities in forest soils. Positively charged ammonium is more easily absorbed  
68 by negatively charged soil colloids than nitrate, meaning that ammonium is more available to  
69 microorganisms than nitrate. In our previous study, we showed that ammonium promoted the activities  
70 of BG- $\beta$ -1,4-glucosidase (BG) and NAG- $\beta$ -1,4-N-acetylglucosaminidase (NAG) in soil aggregates were  
71 strongly than nitrate (Yan et al., 2017). However, the process of nitrification, i.e. where ammonium is  
72 rapidly transformed to nitrate when it enters soil, may sterilize microorganisms in the soil (Dail et al.,  
73 2001). Ammonium and nitrate have different effects on the microbial decomposition rate and microbial  
74 respiration of soil organic matter. For example, substrate respiration in peatlands increased when  
75 ammonium was added, but did not change when nitrate was added (Currey et al., 2010). Nitrate  
76 additions strongly promoted the decomposition rates of soil organic matter of fir plantations in the early  
77 incubation phase (0–15 d; Zhang et al., 2012). However, from a laboratory incubation experiment,  
78 Ramirez et al. (2010) showed that nitrate and ammonium had similar inhibitory effects on soil  
79 microbial respiration.

80 It is well known that microorganisms and enzymes are sensitive to soil pH. Tian and Niu (2015),  
81 from their meta-analysis of soil acidification caused by N additions, suggested that ammonium nitrate  
82 ( $NH_4NO_3$ ) contributed more to soil acidification than ammonium. Further, most studies have not  
83 separated the individual effects of additions of different nitrogen forms on PLFAs and microbial  
84 biomass carbon (MBC) in forest ecosystems. From their meta-analysis, Treseder et al. (2008) reported  
85 that N additions caused MBC to decrease by 15%, and that fungi were more sensitive to N additions  
86 than other microbial communities. The responses of microbial biomass to N additions may be  
87 influenced by a wide range of factors, including forest type and geographical location. For example, in  
88 temperate regions, the total PLFA contents decreased in American beech (*Fagus grandifolia* Ehrh) and  
89 yellow birch (*Betula alleghaniensis* Britton), but increased in eastern hemlock (*Tsuga Canadensis* (L.)  
90 Carr) and red oak (*Quercus rubra* (L.) Britton) forests when  $NH_4NO_3$  was added, with variable

91 responses from bacteria and fungi (Weand et al., 2010). In subtropical forests,  $\text{NH}_4\text{NO}_3$  additions  
92 resulted in an increase in total PLFA contents in a Chinese fir forest (Dong et al., 2015), a decrease in  
93 soil MBC contents in an evergreen broad leaved forests, but no change in the pine broad-leaved mixed  
94 forest (Wang et al., 2008).

95 Soil enzymes catalyze the decomposition of soil organic matter (Burns et al., 2013). Enzymes  
96 involved in labile C breakdown that can decompose starch, cellulose, and hemicellulose include  
97  $\alpha$ -1,4-glucosidase ( $\alpha$ G),  ~~$\beta$ -1,4-glucosidase ( $\beta$ G)~~, cellobiohydrolase (CBH), ~~and~~  $\beta$ -1,4-xylosidase ( $\beta$ X)  
98 ~~and  $\beta$ G~~.  ~~$\beta$ -1,4-N-acetylglucosaminidase (NAG)~~, a nitrogen-degradation enzyme, can decompose  
99 oligosaccharides. Acid phosphatase (AP), a phosphorus-degradation enzyme, can decompose chitin  
100 lipophosphoglycan (Stone et al., 2014). Recalcitrant C-degradation enzymes that can decompose lignin,  
101 and aromatic and phenolic compounds including peroxidase and phenol oxidase (Sinsabaugh et al.,  
102 2010). When added to peatland, Currey et al. (2010) found that ammonium and nitrate had different  
103 effects on carbon- and phosphorus-enzyme activities (CBH and AP) but had similar effects on  
104 polyphenol oxidase (PPO) activities, while Tian et al. (2014) found that the effects of ammonium and  
105 nitrate were not significantly different when added to an alpine meadow. To date, few studies have  
106 reported how ammonium and nitrate additions individually influence soil enzyme activities in forest  
107 ecosystems.

108 ~~In line with the economic theory, m~~Microorganisms will allocate ~~enzymes following economic~~  
109 ~~theory that microbes allocate energy to the~~ relatively absent resources according to economic theory  
110 ~~that are absent~~, so that N additions will cause C and P-acquisition enzymes to increase, and  
111 N-acquisition enzymes to decrease (Burns et al., 2013). It has been reported that, when inorganic N  
112 forms were not considered, N additions caused C-degradation enzymes ( $\alpha$ G,  $\beta$ G, CBH and  $\beta$ X) and  
113 P-degradation enzymes (AP) to increase, restricted oxidase (PPO and PER), but did not inhibit  
114 N-degradation enzymes (NAG) (Jian et al., 2016; Marklein and Houlton, 2012), which suggests that the  
115 allocation of enzyme activities does not always correspond exactly with the economic theory.

116 The responses of enzyme activities to N additions are influenced by a range of factors including  
117 environmental conditions, plant types, and N background values. For example, in temperate regions,  
118 the soil activities of  ~~$\beta$ G~~, CBH, NAG, and PPO increased in a dogwood forest, decreased in an oak  
119 forest, and did not change in a maple forest when  $\text{NH}_4\text{NO}_3$  was added (Sinsabaugh et al., 2002); ~~while~~  
120 ~~The~~ AP activities increased in dogwood and maple forests, but were invariant in an oak forest after

121 NH<sub>4</sub>NO<sub>3</sub> additions (Sinsabaugh et al., 2002). However, in acidified temperate regions, the soil ~~βGBG~~  
122 activities increased in a maple forest, but the soil ~~βGBG~~, NAG, and AP activities did not change in  
123 yellow birch, oak, hemlock, and beech forests, when NH<sub>4</sub>NO<sub>3</sub> was added (Weand et al., 2010). In  
124 subtropical and tropical forests, the ~~βGBG~~, NAG, and AP activities increased, and oxidase (PPO and  
125 PER) activities decreased, after NH<sub>4</sub>NO<sub>3</sub> additions (Dong et al., 2015; Guo et al., 2011; Cusack et al.,  
126 2011). To date, we are still not sure if ammonium and nitrate additions have different effects on the soil  
127 microbial biomass of different communities and on enzyme activities. To support improved predictions  
128 of the effects of elevated N deposition on C, N, and P cycling in soil, we therefore need to evaluate the  
129 individual effects of ammonium and nitrate additions on the soil microbial biomass of different  
130 communities and enzyme activities.

131 The N-rich subtropical soils in southern China have experienced increased nitrate deposition in the  
132 recent past. To facilitate an exploration of the different effects of ammonium and nitrate additions on  
133 soil microbial communities and enzyme activities, we established a long-term ammonium and nitrate  
134 trial in a slash pine (*Pinus elliottii*) plantation in a subtropical area. We hypothesized that (1)  
135 ammonium would have stronger inhibitory effects on total PLFA, fungi PLFA contents, and enzyme  
136 activities than nitrate because of its strong negative effect on soil pH; and (2) that ammonium and  
137 nitrate additions would result in increased C- and P-hydrolase activities, and decreased N-hydrolase  
138 activities in line with the economic theory;<sup>35</sup> and (3) that oxidase activities would be restricted due to  
139 their inhibitory effects on fungi.

140

## 141 **2. Materials and methods**

142

### 143 **2.1. Study site**

144

145 The study was conducted in the Qianyanzhou (~~QYZ~~) Experimental Station, in the hilly red soil  
146 region of Taihe County, Jiang Xi Province, China (26°44'29.1"N, 115°03'29.2"E, 102 m above sea  
147 level a.s.l.). The region has a subtropical monsoon climate, a mean annual temperature of 17.9 °C, and  
148 a mean annual precipitation of 1475 mm. The soil formed because of weathering of red sandstone and  
149 mudstone, and, based on the US soil taxonomy (Soil Survey Staff, 2010), is classified as a Typical  
150 Dystrudepts Udepts Inceptisol. The slash pine (*Pinus elliottii*), one of the dominant species in this hilly

151 red soil region, was planted in 1985 under a vegetation restoration program. *Woodwardia japonica*,  
152 *Dicranopteris dichotoma* and *Loropetalum chinense* dominate the understory (Kou et al., 2015).

153

## 154 2.2. Experimental design

155

156 As described by Kou et al. (2015), the plots were established in November 2011 using a randomized  
157 complete block design. Background atmospheric wet N deposition of about 33 kg N ha<sup>-1</sup> yr<sup>-1</sup>  
158 comprises 11 kg N ha<sup>-1</sup> yr<sup>-1</sup> as ammonium and 8 kg N ha<sup>-1</sup> yr<sup>-1</sup> as nitrate (Zhu et al., 2014). ~~Nine 20 ×~~  
159 ~~20 m plots were established at the experimental sites, including a control, ammonium only and nitrate~~  
160 ~~only additions plot treatments with three replicates (3 treatments × 3 replicates). We established a~~  
161 ~~control and test plots at the experimental sites.~~ We equally added two types of N to the test plots, i.e.  
162 ammonium (N<sub>ammonium</sub>) as ammonium chloride (NH<sub>4</sub>Cl) and nitrate (N<sub>nitrate</sub>) as sodium nitrate (NaNO<sub>3</sub>),  
163 at an annual rate of 40 kg N ha<sup>-1</sup> yr<sup>-1</sup>. This rate was about double the background N wet deposition.

164 ~~Each treatment had three replicates, so the experiment comprised a total of nine plots, which each~~  
165 ~~measured 20 × 20 m.~~ The plots had slope angles of less than 15° and were separated by buffer zones of  
166 more than 10 m. The NH<sub>4</sub>Cl or NaNO<sub>3</sub> were dissolved in 30 L of tap water and evenly sprayed onto the  
167 plots once a month, i.e. 12 times per year. The equivalent amount of tap water was sprayed onto the  
168 control plots. Nitrogen additions commenced in May 2012 and were applied each month on non-rainy  
169 days until March 2015. A total of 113 kg N ha<sup>-1</sup> was applied over the course of this study.

170

## 171 2.3. Sampling and analysis

172

173 We collected soil samples in March, June, and October of 2015, to represent spring, summer, and  
174 fall. We removed the surface litter, and extracted soil cores with a diameter of 5 cm from between 0  
175 and 10 cm deep from 5 randomly selected locations in each plot, which we then mixed together as one  
176 composite sample. The atmospheric conditions and plant-derived litters differed between the three  
177 seasons, and so indirectly affected the soil microbial biomass and enzyme activities of different  
178 communities. We collected soils from three seasons so that we could investigate the ~~synthetic seasonal~~  
179 responses of soil microbial biomass and enzyme activities to ammonium and nitrate additions and to  
180 obtain improved information to support predictions of the effects of elevated N depositions on C, N,

181 and P cycling. Field-fresh samples were sieved through a 2 mm mesh after being mixed evenly.  
182 Samples were stored at 4 °C until analysis for PLFA biomarkers, enzyme activities, soil pH,  
183 ammonium, nitrate, and soil dissolved organic carbon (DOC). The PLFA biomarker and enzyme  
184 activity assays were performed on return to the laboratory. Subsamples of each soil were air-dried, and  
185 then sieved through a 0.25 mm mesh before soil organic C (SOC) and total N (TN) concentrations were  
186 determined.

187 The measurement of soil chemical properties was followed the method of Bao (2010). Soil pH was  
188 measured in a soil-water suspension by glass electrode at a soil to water ratio of 1g fresh soil:2.5  
189 volume of water after stirring evenly and leaving to rest for 30 min. Soil water contents (SWC) were  
190 measured by the oven drying method (105 °C). After extraction with 1mol L<sup>-1</sup> KCl and shaking for 2h,  
191 the ammonium and nitrate concentrations in the fresh soils were measured by a continuous flow  
192 auto-analyzer (Bran Lubbe, AA3, Germany). ~~Another portion of the s~~Soil ~~sample~~ DOC was extracted  
193 with distilled water at soil and distilled water at a ratio of 1 g soil : 5 ml distilled water, and ~~soil DOC~~  
194 ~~concentrations were~~ was measured with an organic element analyzer (Liquid TOCII, Elementar,  
195 Germany) after shaking for 2h. Soil TN and SOC were measured with a carbon/nitrogen analyzer  
196 (Vario Max, Elementar, Germany).

197 Phospholipid fatty acid (PLFA) biomarkers were measured as outlined by Bossio and Scow (1998).  
198 In brief, field-fresh soil equal to 8 g of dry soil was subjected to mild alkaline methanolysis to form  
199 fatty acid methyl esters (FAMES). The extracted PLFAs were dissolved in hexane and measured by gas  
200 chromatography (Agilent 6890N) with MIDI peak identification software (version 4.5; MIDI Inc.  
201 Newark, DE) and a DB-5 column. The abundances of the PLFA biomarkers were calculated as nmol  
202 PLFA g<sup>-1</sup> dry soil. The total amounts of the different PLFA biomarkers were used to represent different  
203 groups of soil microorganisms, i.e. gram-positive bacteria (G<sup>+</sup>) by i14:0, i15:0, a15:0, i16:0, i17:0,  
204 a17:0; gram-negative bacteria (G<sup>-</sup>) by 16:1 $\omega$ 7c, cy17:0, 18:1 $\omega$ 7c, cy19:0; arbuscular mycorrhizal fungi  
205 (AMF) by 16:1 $\omega$ 5; saprophytic fungi (SAP) by 18:1 $\omega$ 9c, 18:2 $\omega$ 6c, 18:2 $\omega$ 9c 18:3 $\omega$ 6c; actinomycete (A)  
206 by 10Me16:0, 10Me17:0, 10Me18:0 (Bradley et al., 2007; Deneff et al., 2009). Bacterial biomass was  
207 calculated as the sum of G<sup>+</sup> and G<sup>-</sup>, and fungi biomass were calculated as the sum of AMF and SAP,  
208 respectively.

209 We measured four C-acquisition hydrolases (i.e.  $\alpha$ G,  $\beta$ G, CBH, and  $\beta$ X), one N-acquisition  
210 hydrolase (NAG), and one P-acquisition hydrolase (AP) following the methods of Saiya-Cork et al.

211 (2002), and have provided information about their corresponding substrates and functions in Table S1.  
212 In brief, 1 g of field-fresh soil was homogenized in a 50 mmol L<sup>-1</sup> sodium acetate buffer (125 ml). We  
213 then added 200 µl of homogenate and 50 µl of substrate to black microplates with 96 wells with eight  
214 replicates for each soil sample. The microplates were then incubated at 20 °C for 4 h. After incubation,  
215 10 µl of 1 mol L<sup>-1</sup> NaOH was added to each well to terminate the reactions, and fluorescence values  
216 were measured at an excitation of 365 nm and emission of 450 nm with a microplate fluorometer  
217 (Synergy H4, BioTek). The absolute hydrolase activities were expressed in units of nmol g<sup>-1</sup> soil h<sup>-1</sup>.  
218 We compared the stoichiometry of C and P to N-acquisition enzyme activities by  
219  $\ln(\alpha\text{G}\alpha\text{G}+\beta\text{G}\text{B}\text{G}+\text{CBH}+\beta\text{X}\text{B}\text{X})$  and  $\ln\text{aP}$  to  $\ln\text{NAG}$ , respectively (n=27).

220 Two oxidases, i.e. PER and PPO, were measured using 96-well transparent microplates as outlined  
221 by Saiya-Cork et al. (2002). We added 600 µl of homogenate and 150 µl of substrate to deep  
222 microplates with 96 wells. To measure the PER activities, we added 10 µl of 0.3% H<sub>2</sub>O<sub>2</sub> to the  
223 homogenate and substrates mixtures. After incubation at 20 °C for 5 h, the microplates were  
224 centrifuged at 3000 r for 3 minutes, then 250 µl of liquid supernatant was transferred to a 96-well  
225 transparent microplate. The absorbance values were measured at 460 nm by microplate  
226 spectrophotometer (Synergy H4, BioTek). We calculated the specific activities of the enzymes by  
227 dividing the enzyme activities by the PLFA values to normalize the activity to the size of the microbial  
228 active biomass (Cusack et al. 2011).

229

#### 230 **2.4. Statistical analyses**

231

232 We used a two factor randomized block analysis of variance and Duncan's multiple comparisons to  
233 test the differences between the treatments and sampling time (n=9). To evaluate the effects of  
234 ammonium and nitrate additions, the treatment differences of time-dependent indexes were tested by  
235 one-way analysis of variance (ANOVA) and Duncan's multiple comparisons for each sampling event or  
236 season (n=3). Analyses were performed with SPSS 17.0. Relationships among the soil physical and  
237 chemical properties, soil PLFA biomarker contents, and the soil enzyme activities were tested by  
238 redundancy analysis (RDA) in CANOCO 4.5 (n=27). Results were statistically significant when  $P <$   
239 0.05. The figures were plotted in Sigmaplot 10.0.

240

241 **3. Results**

242

243 **3.1. Soil physical and chemical properties**

244

245 The soil pH and ammonium contents were either treatment- or time-independent. There were  
246 interaction effects between the treatments and the sampling time on the soil DOC and nitrate contents  
247 ( $P < 0.01$ , Table 1). The soil pH decreased by 0.7 of a unit across the three sampling events in the  
248 ammonium-treated plots, but did not change significantly in the nitrate-treated plots (Fig. 1a). The soil  
249 nitrate contents were 165% and 129% higher (Fig. 2b), and the soil ammonium contents were 31% and  
250 38% lower, ~~respectively~~, in the ammonium and nitrate treatments (Fig. 1b & 2) than in the control for  
251 the three sampling events. Compared with the control, the soil DOC concentrations were 17% higher in  
252 the nitrate-treated plots across the three sampling events, but did not change significantly in the  
253 ammonium-treated plots (Fig. 2a). Ammonium contents were higher in March than in June and October  
254 (Fig. 2, Table S2), while DOC and nitrate concentrations were highest in October and lowest in March  
255 (Table 2 Fig. 2a,b).

256

257 **3.2. Soil microbial biomass of different communities**

258

259 Both the treatment and the time of sampling significantly influenced the soil microbial biomass of  
260 the different communities ( $P < 0.01$ ). Total PLFAs, bacteria,  $G^-$ , and  $G^+/G^-$  were either treatment- or  
261 time-independent. There were also interaction effects between treatments and sampling time on fungi,  
262 actinomycetes,  $G^+$ , AMF, SAP, and the fungi/bacteria ratio (Table 1). The inhibition effects of  
263 ammonium additions on total PLFA contents were stronger than those of nitrate additions and the total  
264 PLFA contents were 24% and 11% less in the ammonium- and nitrate-treated plots across the three  
265 sampling events than in the control (Fig. 3a). The PLFA contents of  $G^+$ , AMF, bacteria, fungi, and  
266 actinomycetes were between 14% and 40%, and 7% and 24%, lower in the plots treated with  
267 ammonium and nitrate, respectively, than in the control across the three sampling events (Fig. 3b,c and  
268 Fig. 4a,b,c,d,e). The soil PLFA contents also showed seasonal variation (Table 1). Total PLFA  
269 biomarker contents and bacterium, fungi,  $G^+$ ,  $G^-$ , AMF, and SAP PLFA biomarker contents were  
270 highest in March and lowest in October, while actinomycete PLFA biomarker contents were highest in



271 June and lowest in October ([Fig. 4a,b,c,d,e](#)~~Fig. 4~~, Table S2).

272 The microbial communities ~~shifted from G<sup>-</sup> to~~ was dominated by G<sup>+</sup> in the ammonium-treated plots,  
273 meaning that the G<sup>+</sup>/G<sup>-</sup> ratios were higher in the ammonium-treated plots than in the control or  
274 nitrate-treated plots ([Fig. 3d](#)). The fungi/bacteria ratios were lower in both the ammonium- and  
275 nitrate-treated plots than in the control, but were much lower in the nitrate-treated plots than in the  
276 ammonium-treated plots ([Figs. 3 and Fig. 4f](#)).

### 278 3.3. Soil enzyme activities

279  
280 There were significant influences from both treatment and sampling time on the measured absolute  
281 enzyme activities (P<0.01). Activities of [βGBG](#), AP, and PPO were either treatment- or  
282 time-independent, and there were interaction effects between the treatments and sampling time on  
283 activities of [αGaG](#), [βXBX](#), CBH, NAG, and PER (Table 1). Ammonium and nitrate had similar  
284 inhibition effects on [αGaG](#), [βGBG](#), [βXBX](#), CBH, NAG, PPO, and PER activities, which decreased by  
285 between 6% and 50% across the three sampling events (~~Table 2~~). The AP absolute activities were about  
286 9% lower in the nitrate treatment than in the ammonium treatment (Table 2). When compared to control,  
287 the ratios of C to N-acquisition enzyme activities were about 0.2 higher, the ratios of N to P acquisition  
288 enzyme activities were about 0.1 lower, and there were no obvious differences in the ratios of C to P  
289 acquisition enzyme activities in the ammonium and nitrate treatments. The measured enzyme activities  
290 varied seasonally (Table 2). Activities of [βGBG](#), [βXBX](#), CBH, NAG, AP, and PPO were lowest in  
291 March and highest in October; [αGaG](#) activities were highest in March and lowest in June, and PER  
292 activities were highest in March and lowest in October (Table 2).

293 The treatments had a significant influence on the activities of N- and P-acquisition specific enzymes  
294 (P<0.01), but not on the activities of C and oxidase specific enzymes (Table 1). The inhibitory effects  
295 of nitrate on the activities of N-acquisition specific enzymes were stronger (about 43%) than those of  
296 ammonium (about 21%, [Fig. 5a](#)). When compared with the control, the AP specific activities were about  
297 19% higher in the ammonium-treated plots across the three sampling events ([Fig. 5b](#)).

### 299 3.4. Redundancy analyses

301 The results of RDA between soil properties and absolute enzyme activities showed that the first axis  
302 explained 72.0% of the variability (Fig. 6a), while the results of RDA between soil properties and  
303 microbial community structures showed that the first axis explained 67.5% of the variability (Fig. 6b).  
304 The results of RDA between soil properties and absolute enzyme activities, PLFA biomarker contents  
305 showed that the first and the second axis explained 72.0% and 11.5% (Fig. 6a), 67.5% and 14.3% (Fig.  
306 6b) of the variation.~~The results of RDA between soil properties, absolute enzyme activities, and PLFA~~  
307 ~~biomarker contents showed that the first ordination RDA axis explained 72.0% and 67.5%, respectively,~~  
308 ~~and the second axis explained 11.5% and 14.3%, respectively, of the variation.~~ The RD1 for soil  
309 absolute enzyme activities and PLFA biomarkers was correlated with DOC/SOC, DOC, ammonium,  
310 and SOC. However, nitrate was only correlated with the RD1 of the absolute enzyme activities but not  
311 the PLFA biomarker contents (Fig. 6 a, b). Most of the measured absolute soil enzyme activities and the  
312 PLFA biomarker contents were positively correlated with soil pH, but G<sup>+</sup>/G<sup>-</sup> and F/B were negatively  
313 correlated with soil pH. Ammonium and DOC contents were positively correlated with all the soil  
314 absolute enzyme activities except PER, but were negatively correlated with PLFA biomarker contents.  
315 Nitrate contents were negatively correlated with soil absolute enzyme activities, but were barely  
316 correlated with the PLFA biomarker contents. SWC were positively correlated with soil PLFA  
317 biomarker contents, but were not correlated with the absolute enzyme activities (Fig. 6 a, b).

#### 319 4. Discussion

320  
321 Our results agree with our first hypothesis and show that the inhibition effects on soil PLFA contents  
322 of bacteria, fungi, and actinomycetes across the three sampling events or seasons were stronger when  
323 ammonium was added than when nitrate was added (Figures 3b and Fig. 4a,b, Table 1). Results from  
324 RDA suggest that acidification because of the ammonium additions triggered the decrease in the  
325 microbial biomarkers-PLFA contents (Fig. 6b). Soil microbial biomass may be inhibited by resource  
326 availability and acidification (Sinsabaugh et al., 2014; Moorhead et al., 2006). However, C and N  
327 availability and N availability ~~(the sum of the ammonium and nitrate concentrations)~~ either  
328 increased or stayed the same over the three sampling events when ammonium and nitrate were added  
329 (Figs. 1b and Fig. 2a,b). Ammonium additions may aggravate nitrification in subtropical soils (Tang et  
330 al. 2016), and nitrification may be toxic to microorganisms (Dail et al., 2001), which may then lead to a

331 decrease in the microbial PLFA contents.

332 The soil pH did not change when nitrate was added (Fig. 1a), which may explain why nitrate had  
333 weaker inhibition effects on PLFA biomarker contents than ammonium. Nitrate additions may inhibit  
334 the PLFA biomarker contents because of accelerated leaching of  $\text{Ca}^{2+}$  and  $\text{Mg}^{2+}$  (Qian et al., 2007),  
335 increases in the soil osmotic potential, and activation of  $\text{Al}^{3+}$  absorbed by soil colloids (Treseder et al.,  
336 2008). The PER activity was lower when ammonium and nitrate were added (Table 2), which may  
337 eventually result in polyphenol accumulation in soil. Accumulated polyphenol may be toxic to  
338 microorganisms (Sinsabaugh et al., 2010) and may have contributed to the decrease in the contents of  
339 the PLFA biomarkers. Moreover, the higher soil DOC concentrations observed in the nitrate-addition  
340 treatments (Fig. 2a) may be attributed to changes in the diversity of the composition of saprophytic  
341 bacteria (Freedman and Zak, 2014; Freedman et al., 2016).

342 In our study, the fungi /bacteria ratios were lower in the ammonium and nitrate treatments than in the  
343 control, which suggests that fungi were more sensitive to N additions than bacteria. In an earlier study,  
344 we found that the fine root biomass decreased after N additions (Kou et al., 2015), which suggests that  
345 N might upset the symbiosis between AMF and plants, thereby restricting the AMF-PLFA contents.

346 Our study showed that the absolute activities of C, N, and P-hydrolases and oxidase were inhibited  
347 by ammonium and nitrate in the three seasons (Table 2). This agrees with our second and third  
348 hypothesis, i.e., that N additions caused the absolute activities of the N-acquisition enzyme (NAG) to  
349 decrease, in line with the microbial economic theory; and that N additions reduced the absolute  
350 activities of the oxidase by decreasing the PLFA contents of fungi. However, we did not expect the C-  
351 or P-acquisition enzymes to decrease. As main producers of soil enzymes, the microbial biomass would  
352 decrease in response to ammonium and nitrate additions, resulting in lower absolute enzyme activities  
353 in the treated plots than in untreated plots (Allison et al., 2005).

354 The ratios of C or P to N acquisition enzyme activities were higher in the ammonium and nitrate  
355 treatments than in the control plots, and the N-acquisition enzyme activities per unit of microbial  
356 biomass were lower in the ammonium and nitrate treatments than in the control (Fig. 5a), indicating  
357 that microorganisms secreted enzymes in line with the economic theory. Measured absolute enzyme  
358 activities were positively correlated with soil pH and ammonium contents, and negatively correlated  
359 with nitrate contents (Fig. 6a). The inhibitory effects of N on the soil absolute enzyme activities may be  
360 more closely related to abiotic factors, i.e. soil pH and nitrification, than biotic factors (Kivlin et al.,

361 2016).

362 We also found that ammonium and nitrate additions inhibited AP activities (Table 2). However,  
363 P-acquisition enzyme activities per unit of microbial biomass increased in the ammonium treatments  
364 (Fig. 5b). Li et al. (2016) reported that N applications aggravated the P-limitations on biomass  
365 production (Li et al., 2016). In line with the microbial economic theory, when the P-availability was  
366 low, the activities of P-acquisition enzymes were higher. The decreased AP activities that resulted from  
367 ammonium additions may be more strongly related to abiotic inhibition caused by the ammonium, such  
368 as acidification, aggravated nitrification, and leaching of cations and nitrate, than biotic inhibition.

369 The N treatments also varied significantly on a seasonal basis and there were interaction effects  
370 between N treatments and seasons on the contents of some PLFA biomarkers and enzyme activities  
371 (Table 2). Climate conditions, plant growth, the amount of litter returned, and plant-soil-microorganism  
372 systems-competitive relationship varied across the three seasons. The temperature ranged from 13.5 to  
373 27.6 °C, and precipitation ranged from 88.2 to 176.6 mm, across the three seasons (Fig. S1), and did  
374 not limit the growth of microorganisms. The positive relationships between PLFA biomarker contents  
375 and soil moisture contents indicate that soil moisture had a strong influence on soil microbial  
376 community biomass. There may be interaction effects between plant growth, the mass and quality of  
377 litter, plant-microbe competition, and soil nutrient dynamics. For example, compared with the control  
378 plots, the soil DOC contents were lower, and soil nitrate contents stayed the same in June (the growing  
379 season) in the ammonium treatment, but the soil DOC and nitrate contents were higher in the  
380 ammonium and nitrate treatments in March and October (non-growing season, Fig. 2a). This indicates  
381 that there was stronger competition between plants and microbes for available C and N in June than in  
382 March and October, and that there were interaction effects between plants and microbes on soil C and  
383 N availability. This might explain the interaction effects between N additions and seasons on the  
384 activities of C and N-acquisition enzymes. The effects of interactions between N additions and season  
385 on the AMF PLFA contents, along with available C and N dynamics, may result from plant growth as  
386 plant-AMF symbiotic systems may be influenced by fine root biomass.

387

## 388 5. Conclusions

389

390 The results showed that soil bacteria, fungi, and actinomycetes- PLFA biomarker contents decreased

391 after ammonium and nitrate additions. Ammonium inhibited the biomass of different soil microbial  
392 communities except SAP more strongly than nitrate, perhaps because of acidification caused by  
393 ammonium. The microbial communities were dominated by G<sup>+</sup> and bacteria after ammonium additions,  
394 and were dominated by bacteria under nitrate additions.

395 The absolute activities of C, N, and P-acquisition hydrolases and oxidase decreased after additions of  
396 ammonium and nitrate, and nitrate had a stronger inhibition effects on P-acquisition absolute enzyme  
397 activities than ammonium. However, ammonium improved the P-demand per unit of microbial biomass.  
398 C and P-acquisition absolute enzyme activities were higher than N-acquisition absolute enzyme  
399 activities under ammonium and nitrate additions. Because of the positive correlation between the  
400 measured absolute enzyme activities and soil pH, the decreases in the absolute hydrolase and oxidase  
401 activities reflected abiotic restrictions, i.e. acidification and nitrification caused by ammonium  
402 additions, rather than biotic restrictions.

403 Ammonium and nitrate additions had a range of effects on soil microbial communities and the  
404 activities of specific enzymes. Our results show that the effects of ammonium and nitrate need to be  
405 discussed separately to provide the information that we need to predict the effects of elevated N  
406 deposition on soil microbial biomass and enzyme activities.

407

408 *Author contribution:* Xin-yu Zhang, Xue-Fa Wen, Sheng-Gong Li, Hui-Min Wang, and Xiao-Min Sun  
409 designed the research; Chuang Zhang, Liang Kou, and Yang Yang performed the study and analyzed  
410 data; and Chuang Zhang, Xin-yu Zhang and Hong-tao Zou wrote the paper.

411

412 *Competing interests:* The authors declare no conflict of interest.

413

414 *Acknowledgments*

415

416 This study was jointly financed by the General, State Key and Major Programs of National Natural  
417 Science Foundation of China (Nos. 41571251, 41571130043, 31130009)

418

419 **References**

420

421 Allison S. D., and Vitousek P. M.: Response of extracellular enzymes to simple and complex nutrient  
422 inputs. *Soil Biology and Biochemistry*, 37, 937-943, doi: 10.1016/j.soilbio.2004.09.014, 2005.

423 [Bao, S.D.: Soil and agricultural chemistry analysis. third ed. Agriculture Press, Beijing \(In Chinese\)](#)  
424 [2008.](#)

425 Burns R. G., DeForest J. L., Marxsen J., Sinsabaugh R. L., Stromberger M. E., Wallenstein M. D.,  
426 Weintraub M. N., and Zoppini A.: Soil enzymes in a changing environment: Current knowledge  
427 and future directions. *Soil Biology and Biochemistry*, 58, 216-227, doi:  
428 org/10.1016/j.soilbio.2012.11.009, 2013.

429 Cusack D. F., Silver W. L., Torn M. S., Burton S. D., and Firestone M. K.: Changes in microbial  
430 communities characteristics and soil organic matter with nitrogen additions in two tropical forests.  
431 *Ecology*, 92, 621-630, doi: 10.1890/10-0459.1, 2011.

432 Dail D. B., Davidson E. A., and Chorover J.: Rapid abiotic transformation of nitrate in an acid forest  
433 soil. *Biogeochemistry*, 54, 131-143, doi: 10.1023/A:1010627431722, 2001.

434 Dong W. Y., Zhang X. Y., Liu X. Y., Fu X. L., Chen F. S., Wang H. M., Sun X. M., and Wen X. F.:  
435 Responses of soil microbial communities and enzyme activities to nitrogen and phosphorus  
436 additions in Chinese fir plantations of subtropical China. *Biogeosciences*, 12, 5540-5544, doi:  
437 10.5194/bg-12-5537-2015, 2015.

438 Freedman Z., and Zak D. R.: Atmospheric N Deposition Increases Bacterial Laccase-Like Multicopper  
439 Oxidases: Implications for Organic Matter Decay. *Applied and Environmental Microbiology*, 80:  
440 4460-4468, doi: org/10.1128/AEM.01224-14, 2014.

441 Freedman Z. B, Upchurch R. A, Zak D. R., and Cline L C.: Anthropogenic N Deposition Slows Decay  
442 by Favoring Bacterial Metabolism: Insights from Metagenomic Analyses. *Frontiers in*  
443 *Microbiology*, 7: 1-11, doi: 10.3389/fmicb.2016.00259, 2016.

444 Frostegård A., Tunlid A., and Bååth E.: Use and misuse of PLFA measurements in soils. *Soil Biology*  
445 *and Biochemistry*, 43, 1621–1625, doi: org/10.1016/j.soilbio.2010.11.021, 2011.

446 Gao W. L., Yang H., Kou L., and Li S. G.: Effects of nitrogen deposition and fertilization on N  
447 transformations in forest soils: a review. *Journal of Soil and Sediments*, 15, 863–875, doi:  
448 10.1007/s11368-015-1087-5, 2015.

449 Guo P., Wang C. Y., Jia Y., Wang Q., Han G. M., and Tian X. J.: Response of soil microbial biomass and  
450 enzymatic activities to fertilizations of mixed inorganic and organic nitrogen at a subtropical forest

451 in East China. *Plant and soil*, 338, 357-361, doi: 10.1007/s11104-010-0550-8, 2011.

452 Heijden M. G. A. V. D., Bardgett R. D., and Straalen N. M. V.: The unseen majority: soil microbes as  
453 drivers of plant diversity and productivity in terrestrial ecosystems. *Ecology letters*, 11, 296-310,  
454 doi: 10.1111/j.1461-0248.2007.01139.x, 2008.

455 Hobbie S. E., Eddy W. C., Buyarski C. R., Adair C. A., Ogdahl M. L., and Weisenhorn P.: Response of  
456 decomposing litter and its microbial community to multiple forms of nitrogen enrichment.  
457 *Ecological Monographs*, 82, 389–405, doi: 10.1890/11-1600.1, 2012.

458 Huang J., Mo J. M., Zhang W., and Lu X. K.: Research on acidification in forest soil driven by  
459 atmospheric nitrogen deposition. *Acta Ecologica Sinica*, 34, 304-306, doi:  
460 org/10.1016/j.chnaes.2014.10.002, 2014.

461 Jia Y. L., Yu G. R., He N. P., Zhan X. Y., Fang H. J., Sheng W. P., Zuo Y., Zhang D. Y., and Wang Q. F.:  
462 Spatial and decadal variations in inorganic nitrogen wet deposition in China induced by human  
463 activities. *Scientific reports*, 4, 1-3, doi: 10.1038/srep03763, 2014.

464 Jian S. Y., Li J. W., Chen J., Wang G. S., Mayes M. A., Dzantor K. E., Hui D. F., and Luo Y. Q.: Soil  
465 extracellular enzyme activities, soil carbon and nitrogen storage under nitrogen fertilization: A  
466 meta-analysis. *Soil Biology and Biochemistry*, 101, 32-41, doi: org/10.1016/j.soilbio.2016.07.003,  
467 2016.

468 Kivlin S. N, and Treseder K. K.: Soil extracellular enzyme activities correspond with abiotic factors  
469 more than fungal community composition. *Biogeochemistry*, 117, 24-34, doi:  
470 10.1007/s10533-013-9852-2, 2014.

471 Kou L., Chen W. W., Zhang X. Y., Gao W. L., Yang H., Li D. D, and Li S. G.: Differential responses of  
472 needle and branch order-based root decay to nitrogen additions: dominant effects of  
473 acid-unhydrolyzable residue and microbial enzymes. *Plant and Soil*, 394, 318-319,  
474 doi:10.1007/s11104-015-2517-2, 2015.

475 Kou L., Guo D. L., Yang H., Gao W. L., and Li S. G.: Growth, morphological traits and mycorrhizal  
476 colonization of fine roots respond differently to nitrogen addition in a slash pine plantation in  
477 subtropical China. *Plant and Soil*, 391, 207-218, doi:10.1007/s11104-015-2420-x, 2015.

478 Li Y., Niu S. L., and Yu G. R.: Aggravated phosphorus limitation on biomass production under  
479 increasing nitrogen loading: a meta –analysis. *Global Change Biology*, 22, 934–943, doi:  
480 10.1111/gcb.13125, 2016.

481 Liu X. J., Duan L., Mo J. M., Du E. Z., Shen J. L., Lu X. K., Zhang Y., Zhou X. B., He C. N., and  
482 Zhang F. S.: Nitrogen deposition and its ecological impact in China: An overview. *Environmental*  
483 *Pollution*, 159, 2253-2254, doi: org/10.1016/j.envpol.2010.08.002, 2011.

484 Liu X. J., Zhang Y., Han W. H., Tang A., Shen J. L., Cui Z. L., Vitousek P., Erisman J. W., Goulding K.,  
485 Christie P., Fangmeier A., and Zhang F.: Enhanced nitrogen deposition over China. *Nature*, 494,  
486 459-462, doi:10.1038/nature11917, 2013.

487 Marklein A. R., and Houlton B. Z.: Nitrogen inputs accelerate phosphorus cycling rates across a wide  
488 variety of terrestrial ecosystems. *New Phytologist*, 193, 696-702, doi:  
489 10.1111/j.1469-8137.2011.03967.x, 2012.

490 Moorhead D. L., and Sinsabaugh R. L.: A theoretical model of litter decay and microbial interaction.  
491 *Ecological Monographs*, 76, 151-172, doi: 10.1890/0012-9615, 2006.

492 Paulinem, C., David, J., Lucyj, S., Iand, L., Hannah, T., René, V. D., Lorna A. D and Rebekka R. E A.:  
493 Turnover of labile and recalcitrant soil carbon differ in response to nitrate and ammonium  
494 deposition in an ombrotrophic peatland. *Global Change Biology*, 16, 2307-2321, doi:  
495 10.1111/j.1365-2486.2009.02082.x, 2010.

496 Qian C., and Cai Z. C.: Leaching of nitrogen from subtropical soils as affected by nitrification potential  
497 and base cations. *Plant and Soil*, 300, 199–204, doi:10.1007/s11104-007-9404-4, 2007.

498 Ramirez K. S., Craine J. M., and Fierer N.: Nitrogen fertilization inhibits soil microbial respiration  
499 regardless of the form of nitrogen applied. *Soil Biology and Biochemistry*, 42, 2336-2338, doi:  
500 org/10.1016/j.soilbio.2010.08.032. 2010.

501 Saiya-Cork K. R., Sinsabaugh R. L., and Zak D. R.: The effects of long term nitrogen deposition on  
502 extracellular enzyme activities in an *Acer saccharum* forest soil. *Soil Biology and Biochemistry*,  
503 34, 1309–1314, doi: org/10.1016/S0038-0717(02)00074-3, 2002.

504 Sinsabaugh R. L.: Phenol oxidase, peroxidase and organic matter dynamics of soil. *Soil Biology and*  
505 *Biochemistry*, 24, 391-401, doi: org/10.1016/j.soilbio.2009.10.014, 2010.

506 Sinsabaugh R. L., Belnap J., Findlay S. G., Follstad Shah J. J., Hill B. H., Kuehn K. A., Kushe C. R.,  
507 Litvak M. E., Martinez N. G., Moorhead D. L., and Warnock D. D.: Extracellular enzyme kinetics  
508 scale with resource availability. *Biogeochemistry*, 121, 287-301, doi:10.1007/s10533-014-0030-y,  
509 2014.

510 Sinsabaugh R. L., Carreiro M. M., and Repert D. A.: Allocation of extracellular enzymatic activities in



511 relation to litter composition, N deposition, and mass loss. *Biogeochemistry*, 60, 6–22, doi:  
512 10.1023/A:1016541114786, 2002.

513 Soil Survey Staff, 2010. *Keys to Soil Taxonomy*, 11th ed. USDA Natural Resources Conservation  
514 Service, Washington, DC.

515 Stone M. M., DeForest J. L., and Plante A. F.: Changes in extracellular enzyme activity and microbial  
516 community structure with soil depth at the Luquillo Critical Zone Observatory. *Soil Biology and*  
517 *Biochemistry*, 75, 240-241, doi: org/10.1016/j.soilbio.2014.04.017, 2014.

518 Tang Y. Q., Zhang X. Y., Li D. D., Wang H. M., Chen F. S., Fu X. L., Fang J. M., Sun X. M., and Yu G.  
519 R.: Impacts of nitrogen and phosphorus additions on the abundance and community structure of  
520 ammonia oxidizers and denitrifying bacteria in Chinese fir plantations. *Soil Biology and*  
521 *Biochemistry*, 103, 284-293, doi: org/10.1016/j.soilbio.2016.09.001, 2016.

522 Tian D., and Niu S.: A global analysis of soil acidification caused by nitrogen addition. *Environmental*  
523 *Research Letters*, 10, doi: 10.1088/1748-9326/10/2/024019, 2015.

524 Tian X. F., Hu H. W., Ding Q., Song M. H., Xu X. L., Zheng Y., and Guo L. D.: Influence of nitrogen  
525 fertilization on soil ammonia oxidizer and denitrifier abundance, microbial biomass, and enzyme  
526 activities in an alpine meadow. *Biology and Fertility of Soils*, 50, 703-713, doi:  
527 10.1007/s00374-013-0889-0, 2014.

528 Treseder K. K.: Nitrogen additions and microbial biomass: a meta-analysis of ecosystem studies.  
529 *Ecology Letters*, 11, 1114-1118, doi: 10.1111/j.1461-0248.2008.01230.x, 2008.

530 Wang H., Mo J. M., Lu X. K., Xue J. H., Li J., and Fang Y. T.: Effects of elevated nitrogen deposition  
531 on soil microbial biomass carbon in major subtropical forests of southern China. *Acta Ecologica*  
532 *Sinica*, 4, 21-27, doi:10.1007/s11461-009-0013-7, 2008.

533 Weand M. P., Arthur M. A., Lovett G. M., McCulley R. L., and Weathers K. C.: Effect of tree species  
534 and N additions on forest floor microbial communities and extracellular enzyme activities. *Soil*  
535 *Biology and Biochemistry*, 42, 2161-2171, doi: org/10.1016/j.soilbio.2010.08.012, 2010.

536 Wei Y., Wang Z. Q., Zhang X. Y., Yang H., Liu X. Y., and Liu W. J.: Enzyme activities and microbial  
537 communities in subtropical forest soil aggregates to Ammonium and Nitrate-Nitrogen additions.  
538 *Journal of Resources and Ecology*, 8, 258-267, doi: 10.5814/j.issn.1674-764x.2017.03.006, 2017.

539 Willers C., Jansen van Rensburg P. J., and Claassens S.: Phospholipid fatty acid profiling of microbial  
540 communities—a review of interpretations and recent applications. *Journal of Applied Microbiology*,

541 119, 1207-1213, doi:10.1111/jam.12902, 2015.

542 Zhang W. D., and Wang S. L.: Effects of  $\text{NH}_4^+$  and  $\text{NO}_3^-$  on litter and soil organic carbon decomposition  
543 in a Chinese fir plantation forest in South China. *Soil Biology and Biochemistry*, 47, 116-121, doi:  
544 org/10.1016/j.soilbio.2011.12.004, 2012.

545 Zhu J. X., He N. P., Wang Q. F., Yuan G. F., Wen D., Yu G. R., and Jia Y. L.: The composition, spatial  
546 patterns, and influencing factors of atmospheric wet nitrogen deposition in Chinese terrestrial  
547 ecosystems. *Science of the Total Environment*, 511, 777-784, doi: org/ 10.1016/ j.scitotenv.2014.  
548 12. 038, 2015.

549 **Figure Legends**

550

551 **Fig. 1.** The effects of ammonium and nitrate additions on soil pH and ammonium contents. Small  
552 letters represent significant differences between treatments ( $P < 0.05$ ), error bars represent means  $\pm$   
553 standard errors (n=9).

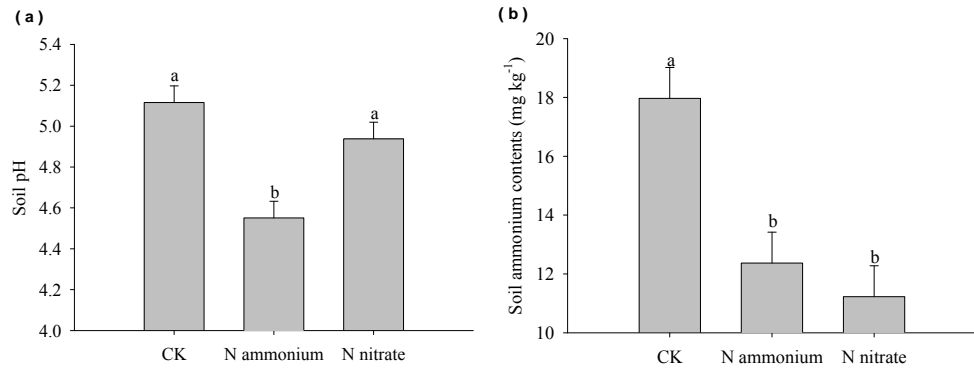
554 **Fig. 2.** The effects of ammonium and nitrate additions on soil nitrate and ~~DOC~~ soil dissolved organic  
555 carbon contents for each sampling event. Capital letters represent significant differences between the  
556 treatments ( $P < 0.05$ ), and small letters represent significant differences between the sampling events ( $P$   
557  $< 0.05$ ), error bars represent means  $\pm$  standard errors (n=3).

558 **Fig. 3.** The effects of ammonium and nitrate additions on Total PLFAs, PLFA contents of bacteria, G<sup>-</sup>  
559 and G<sup>+</sup>/G<sup>-</sup>. Small letters represent significant differences between treatments ( $P < 0.05$ ), error bars  
560 represent means  $\pm$  standard errors (n=9). The abbreviations are the same as Table 1. G<sup>+</sup> represents  
561 gram positive bacteria; and G<sup>-</sup> represents gram negative bacteria.

562 **Fig. 4.** The effects of ammonium and nitrate additions on PLFA contents of fungi, actinomycetes, AMF,  
563 SAP, G<sup>+</sup>, and fungi/bacteria ratio for each sampling event. Capital letters represent significant  
564 differences between the treatments ( $P < 0.05$ ), and small letters represent significant differences  
565 between the sampling time ( $P < 0.05$ ), error bars represent means  $\pm$  standard errors (n=3). The  
566 abbreviations are the same as Table 1. G<sup>+</sup> is gram positive bacteria, AMF is arbuscular mycorrhizal  
567 fungi, and SAP is saprophytic fungi.

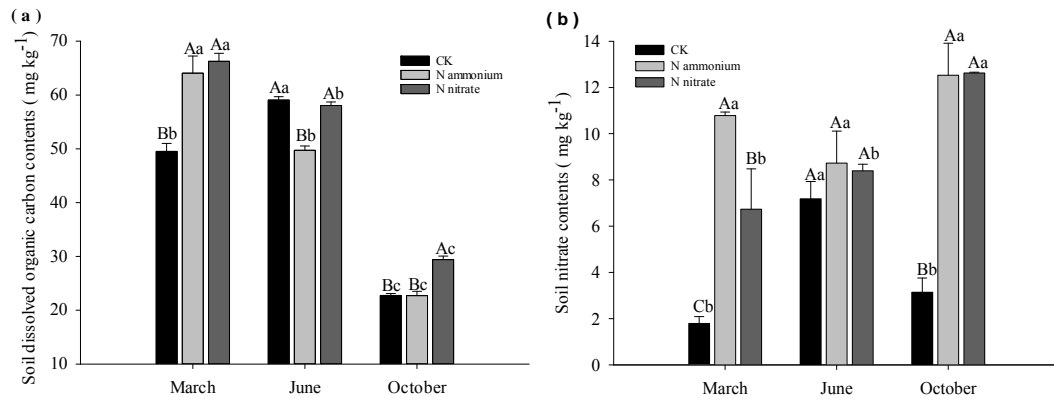
568 **Fig. 5.** The effects of ammonium and nitrate additions on N, P-acquisition specific enzyme activities  
569 for each sampling event. Capital letters represent significant differences between the treatments ( $P$   
570  $< 0.05$ ), and small letters represent significant differences between the sampling time ( $P < 0.05$ ), error  
571 bars represent means  $\pm$  standard errors (n=3).

572 **Fig. 6.** Redundancy analyses between (a) soil properties and enzyme activities, and (b) soil properties  
573 and PLFA-biomarker contents.  $\alpha$ G:  $\alpha$  1,4 glucosidase;  $\beta$ G:  $\beta$  1,4 glucosidase; CBH: Cellobiohydrolase;  
574  $\beta$ X:  $\beta$  1,4 xylosidase; NAG:  $\beta$  1,4 N acetylglucosaminidase; AP: Acid phosphatase; PER: Peroxidase;  
575 PPO: Phenol oxidase. ~~DOC and the abbreviation of PLFA biomarkers were showedn before. The~~  
576 abbreviations are the same as Table 1. SOC: soil organic matter; TN: total nitrogen; C/N: the ratio of  
577 soil organic matter to total nitrogen; SWC: soil water contents.



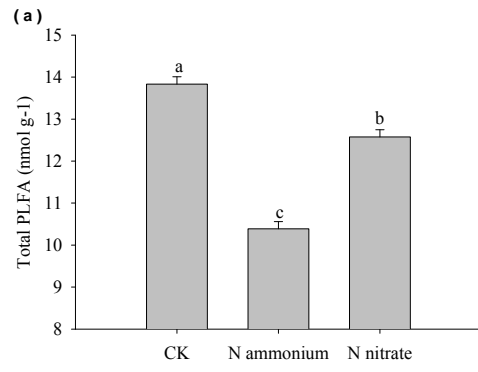
578

579 **Fig.1**

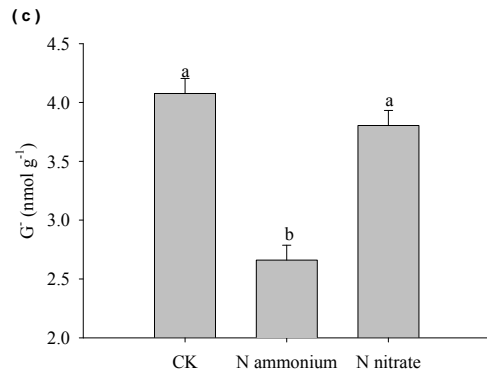
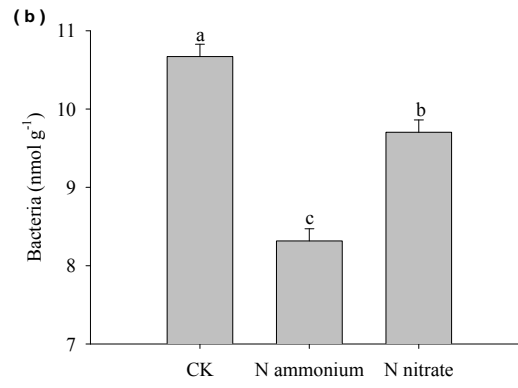


580

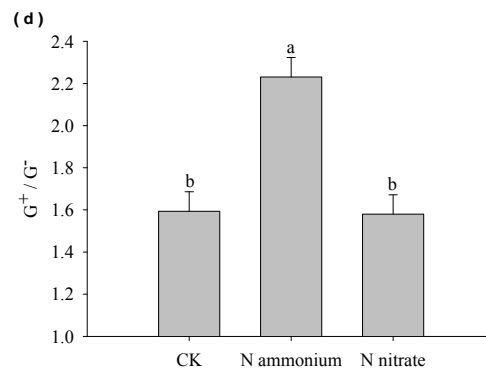
581 **Fig.2**



582

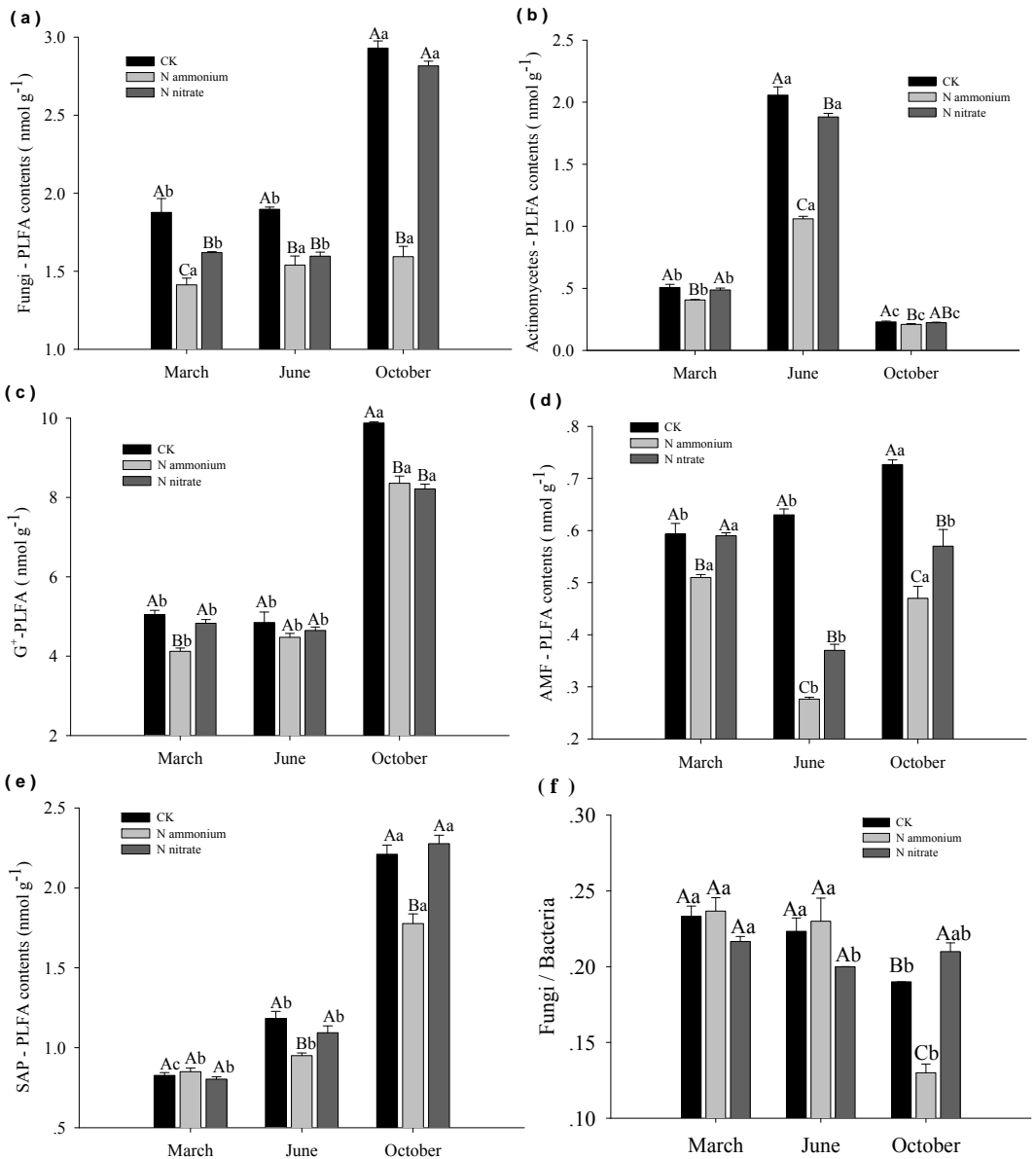


583



584

**Fig.3**

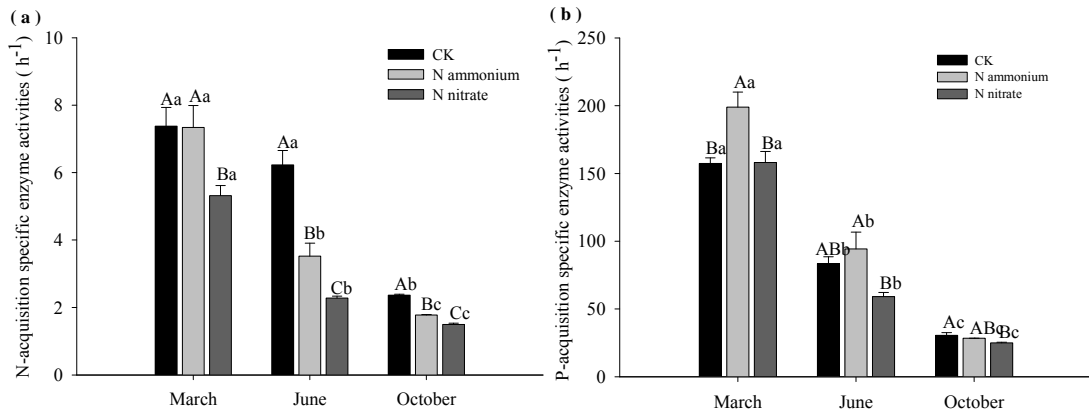


585

586

587  
588

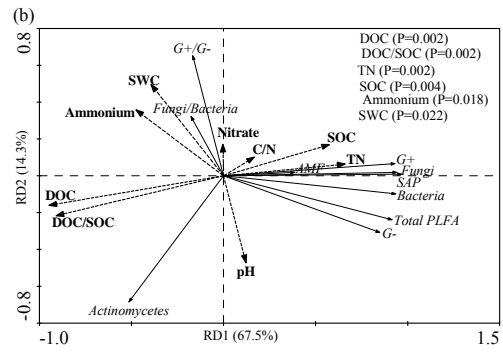
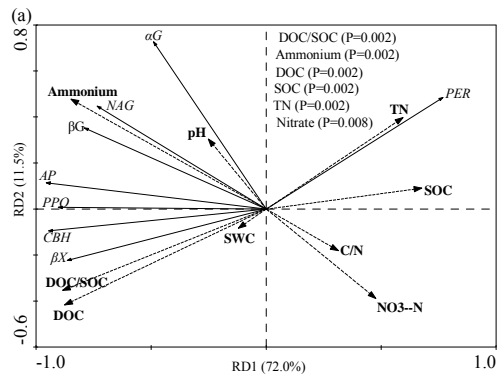
**Fig. 4**



589  
590

**Fig.5**





591

592 Fig. 6

593 **Table 1** Summary statistics (F ratio) for the two factor randomized block analysis of variance (ANOVA)  
 594 applied to soil variables, enzyme activities and PLFA biomarkers. The bold numbers are significant ( $P$   
 595  $< 0.05$ ).

Factors ( <u>Abbreviation</u> )	Treatments	Months	Treatments $\times$ Months
<u>Soil acidity</u> $\text{pH}$ (pH)	<b>12.43</b>	0.31	0.09
<u>DOC</u> <u>Soil dissolved organic carbon</u> (DOC)	<b>23.53</b>	<b>561.25</b>	<b>20.11</b>
Nitrate	<b>43.19</b>	<b>7.96</b>	<b>8.21</b>
Ammonium	<b>11.84</b>	<b>65.46</b>	0.42
<u>TPLFA</u> <u>Total phospholipid fatty acid</u> (TPLFA)	<b>102.51</b>	<b>477.77</b>	2.68
<u>Bacteria</u>	<b>56.94</b>	<b>555.14</b>	2.73
<u>Fungi</u>	<b>180.49</b>	<b>277.81</b>	<b>52.16</b>
<u>Actinomyces</u>	<b>172.230</b>	<b>2627.61</b>	<b>123.12</b>
<u>G<sup>+</sup></u> <u>Gram positive bacteria</u> (G <sup>+</sup> )	<b>50.30</b>	<b>1221.19</b>	<b>14.39</b>
<u>G<sup>-</sup></u> <u>Gram negative bacteria</u> (G <sup>-</sup> )	<b>34.33</b>	<b>105.59</b>	0.45
<u>AMF</u> <u>Arbuscular mycorrhizal fungi</u> (AMF)	<b>147.77</b>	<b>83.55</b>	<b>21.64</b>
<u>SAP</u> <u>Saprophytic fungi</u> (SAP)	<b>24.70</b>	<b>781.67</b>	<b>13.08</b>
G <sup>+</sup> /G <sup>-</sup>	<b>16.24</b>	2.38	0.94
Fungi/Bacteria	<b>3.82</b>	<b>56.42</b>	<b>21.67</b>
<u><math>\alpha</math>G</u> <u><math>\alpha</math>-1,4-glucosidase</u> ( $\alpha$ G)	<b>30.24</b>	<b>53.17</b>	<b>3.47</b>
<u><math>\beta</math>G</u> <u><math>\beta</math>-1,4-glucosidase</u> ( $\beta$ G)	<b>3.26</b>	<b>72.90</b>	0.58
<u><math>\beta</math>X</u> <u><math>\beta</math>-1,4-xylosidase</u> ( $\beta$ X)	<b>9.86</b>	<b>79.08</b>	<b>3.86</b>
<u>CBH</u> <u>Cellulohydrolase</u> (CBH)	<b>28.51</b>	<b>194.75</b>	<b>4.39</b>
<u>NAG</u> <u><math>\beta</math>-1,4-N- acetylglucosaminidase</u> (NAG)	<b>100.42</b>	<b>67.49</b>	<b>8.47</b>
<u>AP</u> <u>Acid phosphatase</u> (AP)	<b>22.81</b>	<b>467.77</b>	1.73
<u>PPO</u> <u>Peroxidase</u> (PPO)	<b>6.87</b>	<b>64.40</b>	1.98
<u>PER</u> <u>Phenol oxidase</u> (PER)	<b>6.27</b>	<b>194.30</b>	<b>3.07</b>
C-acquisition specific enzyme	2.82	<b>334.41</b>	2.07
N-acquisition specific enzyme	<b>29.10</b>	<b>128.31</b>	<b>6.36</b>
P-acquisition specific enzyme	<b>13.42</b>	<b>397.19</b>	<b>4.53</b>
Oxidase specific enzyme	1.68	<b>89.04</b>	1.84

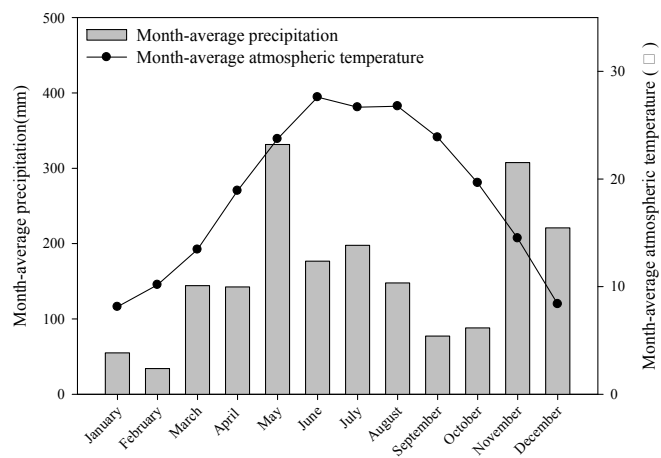
596 **Table 2** Summary statistics (means  $\pm$  standard errors, n=3) for one way analyses of variance (ANOVA) and Duncan multiple  
 597 comparisons applied to soil absolute enzyme activities. ~~Capital letters represent significant differences between the treatments ( $P$~~   
 598  ~~$<0.05$ ), and small letters represent significant differences between the sampling events ( $P < 0.05$ ).~~

Months	Treatments	<del>aGcG</del> nmol g <sup>-1</sup> h <sup>-1</sup>	<del>BGβG</del> nmol g <sup>-1</sup> h <sup>-1</sup>	<del>BXβX</del> nmol g <sup>-1</sup> h <sup>-1</sup>	CBH nmol g <sup>-1</sup> h <sup>-1</sup>	NAG nmol g <sup>-1</sup> h <sup>-1</sup>	AP nmol g <sup>-1</sup> h <sup>-1</sup>	PPO μmol g <sup>-1</sup> h <sup>-1</sup>	PER μmol g <sup>-1</sup> h <sup>-1</sup>
March	CK	7.0±0.1Aa	160.9±15.6 Aa	36.4±3.4Aa	30.±2.1A a	77.5±4.7 Aa	1658.7±59.1 Aa	7.9±0.9Aa	1.4±0.1A b
	N <sup>-</sup> ammonium	4.5±0.2Ba	143.5±4.0A a	26.8±3.2Aa	27.3±1.5 Aa	56.1±5.2B a	1520.7±78.2 Aa	8.9±0.0Aa	1.5±0.1A b
	N <sup>-</sup> nitrate	4.5±0.2Ba	157.1±10.9 Aa	33.4±1.0Aa	21.0±0.8 Ba	49.7±2.6B a	1475.2±53.2 Aa	9.9±1.4Aa	1.6±0.1A b
June	CK	4.0±0.9Ab 2.2±0.1A	83.2±13.0A b	37.2±1.6Aa	28.6±2.5 Aa	77.0±4.7 Aa	1030.3±41.2 Ab	7.7±1.2Aa	1.4±0.1A b
	N <sup>-</sup> ammonium	Bc	70.6±0.9Ab 89.4±10.3A	25.9±1.8Ba	17.9±0.2 Bb	31.8±1.7B b	848.5±62.1B b	4.0±0.0Bb	0.9±0.1B b
	N <sup>-</sup> nitrate	1.7±0.3Bb	b	28.7±1.2Bb 15.2±0.4A	Ba 9.7±0.3A	b 44.7±0.2	b 578.0±38.1A	Bb	4.8±0.9A 7.6±0.1A
October	CK	3.7±0.4Ab	89.1±0.9Ab	Bb	b	Ab 26.5±0.2B	c	2.9±0.2Ab	a 5.5±0.8A
	N <sup>-</sup> ammonium	3.7±0.1Ab	64.0±4.2Ab 68.3±11.5A	16.2±0.9Ab	5.2±0.1Bc 5.3±0.1B	b 24.5±0.2C	423.4±1.6Bc	2.8±0.1Ab	a 5.6±0.8A
	N <sup>-</sup> nitrate	2.2±0.0Bb	b	13.5±0.1Bc	b	b	409.8±4.7Bc	1.9±0.1Bc	a

599 Note: Capital letters represent significant differences between the treatments ( $P < 0.05$ ), and small letters represent significant  
 600 differences between the sampling events ( $P < 0.05$ ). The abbreviations are the same as Table 1.  
 601

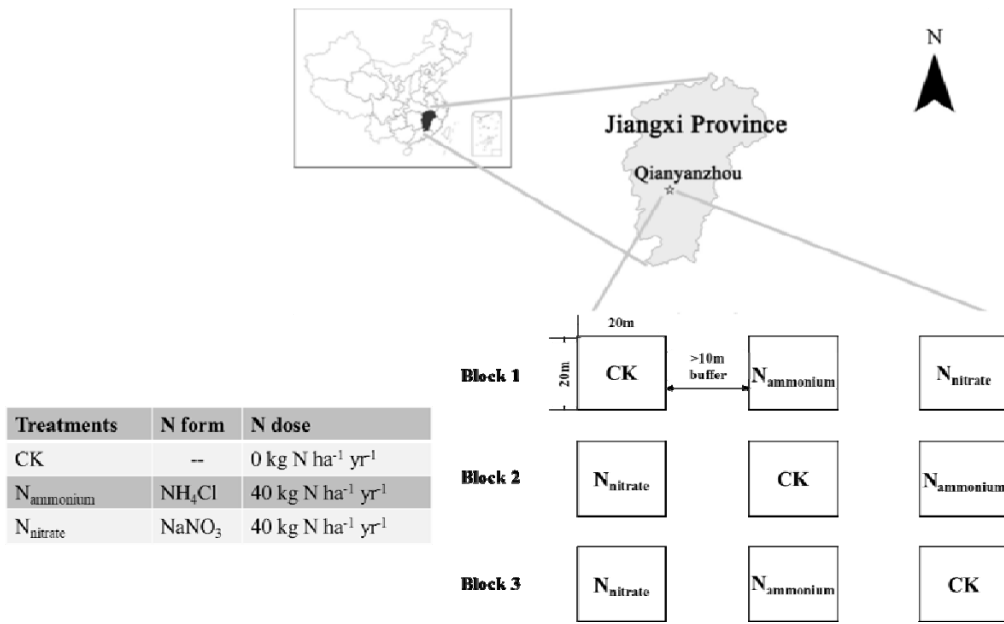
602 **Supplementary materials**

603



604

605 **Fig S1.** Average monthly atmospheric temperature and precipitation at the study site during 2015.



606

607

**Fig S2. The schematic diagram of our the experimental design treatments.**

608 **Table S 1** Enzymes and their corresponding substrates and functions.

Enzyme	Ec	Abbrevia tion	Substrate	Function
Peroxidase	1.11.1.7	PER	L-DOPA	Oxidize lignin and aromatic compounds using H <sub>2</sub> O <sub>2</sub> or secondary oxidants as an electron acceptor
Phenol oxidase	1.10.3.2	PPO	L-DOPA	Oxidize phenolic compounds using oxygen as an electron acceptor
$\alpha$ -1,4-glucosidase	3.2.1.20	<del><math>\alpha</math>GaG</del>	4-MUB- $\alpha$ -D-glucoside	Releases glucose from starch
$\beta$ -1,4-glucosidase	3.2.1.21	<del><math>\beta</math>GBG</del>	4-MUB- $\beta$ -D-glucoside	Releases glucose from cellulose
Cellobiohydrolase	3.2.1.91	CBH	4-MUB- $\beta$ -D-cellobioside	Releases disaccharides from cellulose
$\beta$ -1,4-xylosidase	3.2.1.37	<del><math>\beta</math>XBX</del>	4-MUB- $\beta$ -D-xyloside	Releases xylose from hemicellulose
$\beta$ -1,4-N-acetylglucosaminidase	3.2.1.14	NAG	4-MUB-N-acetyl- $\beta$ -D-glucosaminide	Releases N-acetyl glucosamine from oligosaccharides
Acid phosphatase	3.1.3.1	AP	4-MUB-phosphate	Releases phosphate groups

609 **Table S2** Time-independent seasonal variations in ammonium and PLFAs. Small letters represent  
610 significant differences between the sampling time ( $P < 0.05$ ), error bars represent means  $\pm$  standard  
611 errors (n=9).

Months	Ammonium mg kg <sup>-1</sup>	Total PLFA nmol g <sup>-1</sup>	Bacteria nmol g <sup>-1</sup>	G <sup>-</sup> nmol g <sup>-1</sup>
March	23.5 $\pm$ 1.0a	9.2 $\pm$ 0.2c	7.1 $\pm$ 0.2c	2.5 $\pm$ 0.1c
June	10.6 $\pm$ 1.0b	11.0 $\pm$ 0.2b	7.7 $\pm$ 0.2b	3.1 $\pm$ 0.1b
October	7.5 $\pm$ 1.0b	16.7 $\pm$ 0.2a	13.8 $\pm$ 0.2a	5.0 $\pm$ 0.1a

612