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## Responses of rice paddy micro-food webs to elevated CO<sub>2</sub> are modulated by nitrogen fertilization and crop cultivars

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6 **Responses of rice paddy micro-food webs to elevated CO<sub>2</sub> are modulated by nitrogen**  
7 **fertilization and crop cultivars**

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32

33 **Abstract**

34 Elevated atmospheric CO<sub>2</sub> concentrations (eCO<sub>2</sub>) often increase plant growth but  
35 simultaneously lead to the nitrogen (N) limitation in soil. The corresponding mitigation  
36 strategy such as supplementing N fertilizer and growing high-yielding cultivars at eCO<sub>2</sub>  
37 would further modify soil ecosystem structure and function. Little attention has, however,  
38 been directed toward assessing the responses of soil food web. We report results from a  
39 long-term free air CO<sub>2</sub> enrichment (FACE) experiment in a rice paddy agroecosystem that  
40 examined the responses of soil micro-food webs to eCO<sub>2</sub> and exogenous nitrogen fertilization  
41 (eN) in the rhizosphere of two rice cultivars with distinctly weak and strong responses to  
42 eCO<sub>2</sub>. Soil micro-food web parameters, including microfauna (protists and nematodes) and  
43 soil microbes (bacteria and fungi from phospholipid fatty acid (PLFA) analysis), as well as  
44 soil C and N variables, were determined at the heading and ripening stages of rice. Results  
45 showed that eCO<sub>2</sub> effects on soil micro-food webs depended strongly on N fertilization, rice  
46 cultivar and growth stage. eCO<sub>2</sub> stimulated the fungal energy channel at the ripening stage, as  
47 evidenced by increases in fungal biomass (32%), fungi:bacteria ratio (18%) and the  
48 abundance of fungivorous nematodes (64%), mainly due to an enhanced carbon input. The eN  
49 fueled the bacterial energy channel by increasing the abundance of flagellates and  
50 bacterivorous nematodes, likely through alleviating the N-limitation of plants and rhizosphere  
51 under eCO<sub>2</sub>. While eCO<sub>2</sub> decreased the abundance of herbivorous nematodes under the  
52 weak-responsive cultivar by 59% and 47% with eN at the heading and ripening stage,  
53 respectively, the numbers of herbivorous nematodes almost tripled ( $\times 2.9$ ; heading) and  
54 doubled ( $\times 1.6$ ; ripening) under the strong-responsive cultivar with eCO<sub>2</sub> at eN due to higher

55 root quantity and quality. Structural equation model (SEM) showed that lower trophic-level  
56 organisms were affected by bottom-up forces of altered soil resources induced by eCO<sub>2</sub> and  
57 eN, and effects on higher trophic level organisms were driven by bottom-up cascades with  
58 69% of the variation being explained. Taken together, strategies to adapt climate change by  
59 growing high-yielding crop cultivars under eCO<sub>2</sub> may face a trade-off by negative soil  
60 feedbacks through the accumulation of root-feeding crop pest species.

61 **Key-words:** Global change; Crop cultivar; Rhizosphere; Soil food webs; Root microbiome;  
62 Soil fauna

## 63 **1. Introduction**

64 Increasing evidence indicates that soil biota can modify ecosystem functions in response to  
65 climate change (Bardgett & van der Putten, 2014; Garcia-Palacios et al., 2015). The rising  
66 atmospheric CO<sub>2</sub> concentration (eCO<sub>2</sub>) often increases plant photosynthetic rate and enhances  
67 carbon allocation belowground by promoting root exudation and root turnover (Ainsworth &  
68 Long, 2005; Hu et al., 1999; Zak et al., 2000). Also, eCO<sub>2</sub> enhances plant water use efficiency  
69 through reducing plant stomatal conductance and density (Jackson et al., 1994). Alterations in  
70 C supply and water resources can lead to changes in the structure and activities of the soil  
71 microbial community and the soil food web (Blankinship et al., 2011; Li et al., 2009; Yeates et  
72 al., 2009; Drigo et al., 2008; Drigo et al., 2010). These alterations in soil biota and their  
73 interactions may in return affect plant productivity through modifying plant-microbe  
74 interactions and nutrient availability (Bardgett & van der Putten, 2014), highlighting the need  
75 to adopt appropriate management practices in agroecosystems for adaption to future climate  
76 change.

77 Some crop cultivars have been found to respond with increased yield to eCO<sub>2</sub>, and  
78 breeding such 'positively responsive' cultivars recently received major attention as a strategy  
79 to optimize crop production under future climate regimes (Brummer et al., 2011; Ziska et al.,  
80 2012; Korres et al., 2016). Yield of rice is highly responsive to eCO<sub>2</sub> (Liu et al., 2008a;  
81 Shimono & Bunce, 2009) with indica varieties being more responsive to eCO<sub>2</sub> than japonica  
82 varieties (Hasegawa et al., 2013; Zhu et al., 2015). Therefore, active selection and breeding  
83 for CO<sub>2</sub> responsiveness among rice varieties was suggested as an effective strategy to increase  
84 global yields and maintain food security under future global climate change scenarios (Ziska

85 et al., 2012). Meanwhile, to satisfy the nutrient demand of high-yielding cultivars, more N  
86 fertilizer is needed (applied as exogenous N or eN). While low N availability in natural  
87 ecosystems may limit plant responses to eCO<sub>2</sub>, high N inputs can mitigate N limitation on  
88 crop growth in agroecosystems (Reich et al., 2006; Feng et al., 2015). N fertilization may also  
89 alleviate N constraints on soil microbes under eCO<sub>2</sub> (Hu et al., 2001; Luo et al., 2004). Yet,  
90 the net-effect of different responsive cultivars to eCO<sub>2</sub> in N-enriched soils on soil food web  
91 and potential feedbacks to plant production have rarely been examined (van der Putten et al.,  
92 2016).

93       The structure complexity of the soil food web, as an integrated indicator of soil biological  
94 interactions that drive soil functioning, can significantly affect crop production in two major  
95 aspects (Bardgett & Wardle, 2010; Neher, 2010). First, soil microbes and soil food web  
96 interactions mediate nutrient cycling. For example, bacterivorous and fungivorous (i.e.  
97 microbivorous) protists and nematodes directly affect the turnover of microbial biomass and  
98 the availability of plant nutrients (Bardgett & Wardle, 2010; Eisenhauer et al., 2012). The  
99 early syntheses found that these microbivores contributed almost 30% of N mineralization in  
100 agroecosystem (Griffiths, 1994; Trap et al., 2016), indicating the pivotal significance of  
101 microbial grazers in N cycling. Similar to nematodes, protists have diverse feeding strategies  
102 including bacterivores, fungivores, and omnivores (Geisen et al., 2016). Previous studies in  
103 grasslands and forests have found eCO<sub>2</sub> tended to favor soil fungi while eN stimulated  
104 bacterial growth (Drigo et al., 2008; Garcia-Palacios et al., 2015). Second, root parasites  
105 among soil microfauna, such as herbivorous nematodes, can directly affect plant growth  
106 (Neher, 2010). Due to this direct trophic link, changes to roots in response to eCO<sub>2</sub> and eN

107 may affect root herbivore performance by altering the quality and quantity of food resources  
108 (Robinson et al., 2012). eCO<sub>2</sub> often leads to a higher C:N ratio in plant tissue, and studies of  
109 aboveground insects found that herbivores must compensate for the differences in elemental  
110 ratios between the food and their requirements through over-grazing (Molles, 2013). However,  
111 it is unclear what factors and/or processes will drive the structure and functioning of  
112 belowground systems in rice paddy soils under future climate change scenarios (Okada et al.,  
113 2014).

114 Taking advantage of the long-term free-air CO<sub>2</sub> enrichment (FACE) platform to assess the  
115 responses of different rice cultivars to CO<sub>2</sub> concentration and N fertilization (Zhu et al., 2016),  
116 we examined the responses of soil micro-food webs (bacteria, fungi, protists and nematodes)  
117 at different growth stages of two rice cultivars with contrasting performance under eCO<sub>2</sub>. We  
118 hypothesized that (1) the bottom-up control plays a primary role in mediating responses and  
119 feedbacks of both the decomposer and herbivorous food webs to eCO<sub>2</sub> and eN, and (2) the  
120 responsiveness of rice cultivars to eCO<sub>2</sub> and eN is a major factor affecting food web structure  
121 via changes in resource input to soil from plant root.

122

## 123 **2. Materials and methods**

### 124 *2.1. Study site and experimental design*

125 An experimental platform of free-air CO<sub>2</sub> enrichment (FACE) was established in 2004, with a  
126 rice-wheat rotation system in Zongcun Village (119°42'0" E, 32°35'5" N), Yangzhou City,  
127 Jiangsu Province. From 2010, the rice-wheat rotation system was changed to a rice-fallow  
128 system. The region has a north subtropical monsoon climate with a mean annual temperature



129 of 16 °C, and mean annual precipitation of 900-1000 mm. The soil at the study site is a  
130 Shajiang-Aquic Cambisol, with 18.4 g·kg<sup>-1</sup> total C, 1.5 g·kg<sup>-1</sup> total N, 57.8% sand, 28.5% silt  
131 and 13.7% clay at 0-15 cm depth.

132 The experiment had a split-plot design with CO<sub>2</sub> as the main factor, and nitrogen  
133 fertilization and rice cultivar as the split plot factors. More details about the FACE system  
134 were described by Zhu et al., (2016). In brief, a randomized complete block design was  
135 established with two levels of target atmospheric CO<sub>2</sub> concentrations. The atmospheric CO<sub>2</sub>  
136 of each FACE ring was enriched by 200 μmol CO<sub>2</sub> mol<sup>-1</sup> over the ambient (Fig. 1). It  
137 consisted of three replicate rings for the eCO<sub>2</sub> and three for the ambient (hereinafter referred  
138 to as aCO<sub>2</sub>). All eCO<sub>2</sub> rings were 12.5 m in diameter, with an area of 80 m<sup>2</sup> that was sampled  
139 after rows on the edge were excluded.

140 We studied treatments with no fertilization (aN) and elevated N fertilization (eN), the  
141 latter receiving urea and compound chemical fertilizer (N: P<sub>2</sub>O<sub>5</sub>: K<sub>2</sub>O = 15:15:15, %) at 22.5  
142 g N m<sup>-2</sup> yr<sup>-1</sup>. Urea was applied as a basal dressing (40% of the total dose) one day prior to rice  
143 transplanting, as a top dressing at early tillering (30%) and at the panicle initiation stage  
144 (30%). Phosphorous (P) and potassium (K) were applied as a compound fertilizer at 9 g P<sub>2</sub>O<sub>5</sub>  
145 m<sup>-2</sup> and 9 g K<sub>2</sub>O m<sup>-2</sup> one day before transplanting. Two contrasting rice cultivars were  
146 planted in aN and eN plots as a split-split-plot in both FACE and ambient rings. Since 2012,  
147 an indica rice IYYou084 and a japonica rice WuYunJing, showing strong (+ 30% yield increase)  
148 and weak (+ 13% yield increase) responses to CO<sub>2</sub> elevation respectively were planted (Zhu et  
149 al., 2015). Compared to the japonica rice, the indica rice was characterized as an increase in  
150 net photosynthetic assimilation, root growth and N uptake capability under eCO<sub>2</sub> (Hasegawa

151 et al., 2013; Zhu et al., 2015) The seedlings were grown under ambient air and were  
152 transplanted by hand into the aCO<sub>2</sub> and eCO<sub>2</sub> plots at a density of three seedlings per hill and  
153 24 hills per m<sup>2</sup> for all six rings on 21<sup>st</sup> June, 2014.

## 154 2.2. Soil sampling and analysis

155 Rhizosphere soil samples were collected with a 3.5 cm diameter corer (0-15 cm depth) no  
156 more than 3.5 cm distant of rice plants (Fig. S1), at rice heading (10<sup>th</sup> Sep) and ripening (27<sup>th</sup>  
157 Oct) stage in 2014. Before sampling, the field was drained for 5 days to facilitate sample  
158 collection. Five soil cores were randomly collected from each treatment in each plot and were  
159 combined to form one composite sample per treatment per plot. Soil samples were stored at 4  
160 °C and analyzed within 7 days after sampling.

161 Dissolved organic carbon (DOC) and nitrogen (DON) was exacted from 10 g fresh soil  
162 using 50 mL ultrapure water by centrifugation (8000 rpm, 10 min). The filtrate that passed  
163 through a 0.45 mm filter membrane was analyzed with a total C analyzer (Elementar,  
164 Germany) and a continuous flow analyzer (Skalar, Holland), respectively. The NH<sub>4</sub><sup>+</sup>-N and  
165 NO<sub>3</sub><sup>-</sup>-N were extracted with 2 M KCl in a 1:5 (soil: water) suspension and the suspension was  
166 filtered through ashless filter paper. The filtrates were determined by a continuous flow  
167 analyzer (Skalar, Holland). Mineral N (MN) was calculated by the sum of NH<sub>4</sub><sup>+</sup>-N and  
168 NO<sub>3</sub><sup>-</sup>-N and total extractable nitrogen (Ext N) was calculated by the sum of DON, NH<sub>4</sub><sup>+</sup>-N  
169 and NO<sub>3</sub><sup>-</sup>-N. Concerning root sampling, representative samples of three individual rice hills  
170 were dug out and pooled from each plot. Roots were carefully washed from the soil, then  
171 oven-dried and weighed. The C and N contents of roots were analyzed with an elemental  
172 analyzer (Elemental, Germany).

173 The soil microbial community was characterized using phospholipid fatty acid (PLFA)  
174 analysis as described by Blight & Dyer (1959) with slight modifications. GC conditions and  
175 nomenclature were as described by Buyer & Sasser (2012). Briefly, 8.0 g freeze-dried soil  
176 was extracted with a chloroform-methanol-citrated buffer mixture (25 mL at a 1:2:0.8 volume  
177 bases). Lipid classes were separated into phospholipid, neutral and glycolipid by solid phase  
178 extraction (SPE) tubes (ANPEL Laboratory Technologies Inc., China) containing 0.5 g  
179 anhydrous sodium sulfate. The phospholipids were trans-esterified by a mild alkaline  
180 methanolysis (Bossio et al., 1998) and the resulting fatty acid methyl esters were extracted in  
181 hexane and dried under N<sub>2</sub>. Samples were re-dissolved in hexane and analyzed in an Agilent  
182 6850 series Gas Chromatograph with MIDI peak identification software (Version 4.5; MIDI  
183 Inc., Newark, DE). The fatty acids i14:0, i15:0, a15:0, i16:0, 16:1 $\omega$ 7c, i17:0, a17:0, 17:0cy,  
184 18:1 $\omega$ 9, 18:1 $\omega$ 7c and 19:0cy were chosen as bacterial markers, and 16:1 $\omega$ 5c and 18:2 $\omega$ 6.9c  
185 were used as fungal markers (Ruess & Chamberlain, 2010). Selected PLFAs biomarker  
186 associated with specific microbial group see Table S1.

187 Protists (amoebae and flagellates) were enumerated using a modified most-probable  
188 number method (Darbyshire et al., 1974), Briefly, 3.0 g fresh soil was suspended in 30 mL  
189 sterile Neff's modified amoebae saline (NMAS) (Page, 1976) and gently shaken (180 rpm)  
190 for 30 min on a vertical shaker. Threefold dilution series with tryptic soy broth (TSB) and  
191 NMAS at 1:9 v/v were prepared in 96-well microtiter plates in quadruplicates. The microtiter  
192 plates were incubated at 15 °C in darkness, and the wells were inspected for presence of  
193 protists using an inverted microscope at  $\times$ 100 to  $\times$ 400 magnification after 7, 14 and 21 days.  
194 Abundance of protists was expressed as the number of individuals per gram of dry soil.

195 Nematode populations were extracted from 100 g fresh soil using a sequential extraction  
196 method (Liu et al. 2008b). After the total numbers of nematodes were counted, 100 specimens  
197 per sample were randomly selected and identified to the genus level. If the total number was  
198 less than 100, all nematodes were identified. The nematodes were assigned to the following  
199 trophic guilds: bacterivore, fungivore, herbivore and omnivore-carnivore (Yeates et al., 1993).

### 200 2.3. Statistical analyses

201 Analysis of Variance (ANOVA) and Fisher's LSD posthoc tests were performed using  
202 Statistica 7.1 (StatSoft Inc., Tulsa, OK, USA). Since the measurements were repeated on the  
203 same plot over time, repeated measures ANOVA was used to test the effects of CO<sub>2</sub>  
204 concentration (ambient and elevated), exogenous N (ambient and elevated), or rice cultivar  
205 (weak- and strong-responsive) on soil nutrients, soil microbial biomass and the abundance of  
206 soil microfauna across rice growth stages. PLFA profiles of microbial groups data on the  
207 nematode communities were analysed by principal component analysis (PCA) using the  
208 software package CANOCO 5.0 (ter Braak and Smilauer, Wageningen-UR, The Netherlands).

209 Structural equation models (SEM) were calculated to investigate how elevated CO<sub>2</sub> and N  
210 input impacted soil micro-food webs and bacterial and fungal energy channels in soil (as  
211 indicated by PLFA profiles and the trophic structure of soil nematode communities). The SEM  
212 were separately calculated for the heading and ripening stage. Root parameters could only be  
213 collected at the ripening stage after destructive sampling of rice plants, therefore we presented  
214 only the full model for the ripening stage (see Fig. S6 for the model for the heading stage that  
215 lacks root parameters ).

216 The *a priori* model evaluated relationships among root C/N, root N, soil environment

217 (DOC, Ext N, MN), bacteria, fungi, flagellates, amoebae and nematode communities at  
218 trophic group level. The SEM was performed in Amos version 17.0.2 (Amos Development  
219 Corporation, Chicago, IL, USA) using maximum likelihood estimation procedures. Model fit  
220 was assessed by  $\chi^2$ -text, the comparative fit index (CFI) and the root square mean error of  
221 approximation (RSMEA).

222

### 223 **3. Results**

#### 224 *3.1. Soil resources and environment*

225 Regardless of rice cultivar and N dose the eCO<sub>2</sub> significantly ( $p < 0.05$ ) increased DOC  
226 content by 35% and 38% at heading and ripening stage respectively (Fig. 2a). eCO<sub>2</sub> increased  
227 DON by 83% across all treatments at the heading stage, but at the ripening stage, DON was  
228 reduced by 38% with eCO<sub>2</sub> only under the strong-responsive rice (Fig. 2b). Also, mineral N  
229 was reduced by 27% with eCO<sub>2</sub> at the ripening stage (Fig. 2c), while N fertilization increased  
230 mineral N on average by 41%, irrespective of CO<sub>2</sub> and rice cultivar (Fig. 2c).

#### 231 *3.2. Soil microbial community*

232 PCA of the microbial communities (PLFA) showed that eCO<sub>2</sub> tended to amplify the difference  
233 imposed by N and cultivar at the heading stage. Microbial communities under the  
234 strong-responsive rice were strongly influenced by CO<sub>2</sub> and N (PC1 = 83.5% and PC2 =  
235 8.84%; Fig. S3a). At the heading stage, eCO<sub>2</sub> and eN had no effect on the biomass of bacteria  
236 and fungi or the fungi:bacteria ratio in the rhizosphere of either cultivar (Fig. S1a; Table 1). At  
237 the ripening stage, effects of eCO<sub>2</sub> on microbial community structure dominated, with PC1  
238 clearly separating the communities under aCO<sub>2</sub> and eCO<sub>2</sub> (PC1 = 73.2% and PC2 = 14.0% of

239 the variation; Fig. S3b). eCO<sub>2</sub> tended to increase the overall microbial biomass at the ripening  
240 stage, increasing fungal biomass by up to 32% ( $p < 0.05$ ; Fig. 3; Table 1), resulting in a  
241 significant increase of the fungi:bacteria ratio (Fig. S2b).

### 242 3.3. Soil microfauna

243 Repeated measures ANOVA confirmed a significant CO<sub>2</sub> effect on soil microfauna across rice  
244 growth stages. At the heading stage, flagellates were significantly reduced at eCO<sub>2</sub> under aN  
245 by 61% and 44% for weak- and strong-responsive cultivars, respectively (Fig. 4a). eN  
246 induced an overall increase in flagellate abundance of beyond 35% at the ripening stage, but  
247 the response to eCO<sub>2</sub> depended on an interaction between cultivar and N (Fig. 4a and 4c;  
248 Table 1). At the ripening stage, flagellates under weak-responsive cultivar increased on  
249 average 1.6-fold from aN to eN, irrespective of CO<sub>2</sub> level; also flagellates under  
250 strong-responsive cultivars increased 1.5-fold at eN, but only with eCO<sub>2</sub> (Fig. 4a and 4c). The  
251 amoebae increased by 31% under eCO<sub>2</sub> at the heading stage, but at the ripening stage the  
252 opposite trend was found under eN (Table 1; Fig. 4b and 4d).

253 Impacts of eCO<sub>2</sub> on the soil nematode depended on N dose, cultivar and growth stage  
254 (Fig. 5 and S4; Table 1). At the heading stage, eN increased bacterivorous nematodes by 41%  
255 (Fig. 5a) and this effect was maintained at the ripening stage but only for the weak-responsive  
256 cultivar under eCO<sub>2</sub> (Table 1; Fig. 5e). At the ripening stage, fungivorous nematodes increased  
257 2.4-fold with eCO<sub>2</sub> at aN under strong-responsive cultivars, and 2-fold at eN under  
258 weak-responsive cultivars compared to aCO<sub>2</sub> (Table 1; Fig. 5f). Also at the ripening stage,  
259 omnivorous-carnivorous nematodes reached significantly higher densities at eCO<sub>2</sub>,  
260 particularly at eN (Table 1; Fig. 5h).

261 The abundance of herbivorous nematodes consistently increased under the  
262 strong-responsive cultivar at eCO<sub>2</sub>, independent of rice growth stages ( $F = 18.91$ ,  $p < 0.01$ ,  
263 Table 1; Fig. 5c and 5g). At the heading stage under eCO<sub>2</sub>, the numbers of herbivores almost  
264 doubled ( $\times 1.8$  under aN) and tripled ( $\times 2.9$  under eN) under strong-responsive rice with eCO<sub>2</sub>,  
265 while the numbers of herbivorous nematodes decreased by 59% under weak-responsive rice at  
266 eN (Fig. 5c; Table 1). At the ripening stage herbivores increased by 47% when eCO<sub>2</sub> plants  
267 received fertilizer, but decreased by 32% under weak-responsive cultivars (Fig. 5g; Table 1).  
268 On average, herbivores under strong-responsive cultivars increased 1.6-fold under eCO<sub>2</sub>  
269 irrespective of N fertilization. Overall, this resulted in a 1.6-fold increase of root herbivores at  
270 eN compared to aN under eCO<sub>2</sub> conditions (Table 1).

271 *3.4. Effects of eCO<sub>2</sub> and eN on the structure and function of micro-food webs in the*  
272 *rhizosphere of rice*

273 At the heading stage, eCO<sub>2</sub> and eN significantly increased the availability of soil resources, in  
274 particular DOC and DON (Fig. 2 and S6). However, these belowground inputs did not affect  
275 bacterial and fungal biomass (Fig. 3 and S6), likely due to enhanced turnovers of bacterial and  
276 fungal biomass, as indicated by significant higher numbers of bacterivores (nematodes, 26%;  
277 amoebae, 14%) and in particular the omnivorous-carnivorous nematodes (44%) at the third  
278 trophic level (Fig. S6).

279 At the ripening stage, eCO<sub>2</sub> and eN led to increased root biomass and this significantly  
280 increased the abundance of herbivorous nematode (Fig. 6). The increased root biomass  
281 correlated with an enhanced biomass of bacteria (covariance coefficient = 0.39,  $p < 0.05$ ), and  
282 in particular of fungi (covariance coefficient = 0.69,  $p < 0.01$ ; Fig. 6). eCO<sub>2</sub> and eN had a

283 direct impact on soil resources such as DOC and Ext N, and increased the total microbial  
284 biomass. Increased bacterial and fungal biomass was positively associated with the increased  
285 abundance of bacterivorous and fungivorous nematodes, respectively (Fig. 6). Interestingly,  
286 amoebae were directly related to root biomass and statistically marginally associated with  
287 fungi (covariance coefficient = 0.33,  $p = 0.07$ ; Fig. 6).  $eCO_2$  thus directly influenced resource  
288 availability for herbivores (root biomass) and microbes (DOC, Ext N) and subsequently  
289 propagated mainly via the fungal energy channel into the microbial primary consumers and  
290 then further up in the trophic chain to secondary consumers (i.e., omnivorous-carnivorous  
291 nematodes). In total, the model explained 69% of the variance in omnivorous-carnivorous  
292 nematodes, while the remaining relationships between variables were not significant but  
293 improved the fit of the model (Fig. 6).

294

## 295 **4. Discussion**

### 296 *4.1. Responses of bacteria- and fungi-based energy channels of rhizosphere micro-food webs* 297 *to $eCO_2$ and $eN$*

298 Our results showed that positive effect of  $eCO_2$  and  $eN$  on soil micro-food webs were mainly  
299 caused by the altered availability of C and N in the plant rhizosphere, confirming bottom-up  
300 control of the rhizosphere micro-food webs.  $eCO_2$  often stimulates C allocation belowground,  
301 leading to increased microbial biomass and/or microbial respiration (Zak et al., 2000; Hu et  
302 al., 2001; Luo et al., 2004).  $CO_2$ -enhancement of belowground C allocation likely occurs  
303 through increasing root growth and root exudation (Matamala et al., 2003; Norby et al., 2004),  
304 and mycorrhizal fungi (Cheng et al., 2012), providing new resources for microbial growth and



305 subsequent grazers (Blankinship et al., 2011; Mueller et al., 2016). At the ripening stage, with  
306 the root data available, it became clear that the main driving force came from roots (Fig. 6),  
307 especially considering the significant interactive effects of CO<sub>2</sub> and cultivars on root traits  
308 (Fig. S5). In addition, the eCO<sub>2</sub> effect will depend on N availability because the relative  
309 availability of C and N can either drive the bacteria- or fungi-based food web (Bååth et al.,  
310 1981; Mikola & Setälä, 1999).

311 The results of the present study showed that the effects of eCO<sub>2</sub> and eN on soil food webs  
312 can occur through altering biomass and/or turnover rates of each trophic level (Table 1). SEM  
313 indicated a clear dominance of the bacterial energy channel at the heading stage when peak  
314 plant growth occurs (Fig. S6). The fact that the growth of flagellates (Fig. 4a) and  
315 bacterivorous nematodes (Fig. 5a) was strongly restricted by N availability with eCO<sub>2</sub>  
316 suggests that bacterial growth was limited by low N availability resulting from the increased  
317 C availability via root exudation (Hoeksema et al., 2000) and competition between  
318 microorganisms and plant roots for N (Hu et al., 2001; Reich et al., 2006). Abundance of  
319 bacterivores is a long-term indicator of bacterial production, and this may explain why several  
320 previous studies in grasslands or forests observed no significant CO<sub>2</sub> effects on bacterial  
321 biomass or abundance in spite of increasing microbial N limitation (Ebersberger et al., 2004;  
322 Chung et al., 2006; Sinsabaugh et al., 2003). The increased bacterivore numbers under eN  
323 further point towards a strong top-down control over bacterial biomass (i.e. the rhizosphere  
324 microbial loop; see Clarholm, 1985; Bonkowski, 2004). Under elevated CO<sub>2</sub> when the  
325 relative availability of C to N was high, nutrient excretion by bacterivores can alleviate  
326 resource limitation of the grazed microbes to such an extent that reproductive rates of bacteria

327 keep up with grazing rates, increasing microbial turnover rates without detectable effect on  
328 the microbial biomass (Alpei et al., 1996; Frey et al., 2001; Trap et al., 2016). Also, similar  
329 magnitudes of increase in flagellates and bacterivorous nematodes (Fig. 4 and 5) suggests that  
330 food quality (C:N ratio of substrate and the specific populations of bacteria) rather than  
331 quantity (e.g. increased bacterial biomass) stimulated bacterivores (Schmidt et al., 2000;  
332 Cesarz et al., 2015). In contrast, eCO<sub>2</sub>-induced changes in amoebae were independent of  
333 nitrogen status at the heading stage, indicating different trophic relationships in comparison  
334 with flagellates (Fig. 4b; Geisen, 2016). Also, the SEM indicated that amoebae were  
335 positively associated with root biomass (Fig. 6). Together, these results suggest that amoebae  
336 were related to enhanced root exudation associated with root growth promotion under eCO<sub>2</sub>  
337 and eN (i.e. the modified microbial loop; Bonkowski and Clarholm, 2012).

338 The alteration of bacterivores, especially nematodes and amoebae at the heading stage  
339 explained a large proportion of the variance in omnivorous-carnivorous nematodes (44%) at  
340 the third trophic level (Fig. S6). These results accord with the conceptual framework of the  
341 microbial loop, indicating a strong top-down control of bacterial biomass by bacterivores  
342 (Bonkowski, 2004; Neher et al., 2004; Wolkovich, 2016), with bottom-up root control and  
343 significant transfer of bacteria-derived C and N to higher trophic levels at the heading stage.  
344 Consistent with this notion, a shift in relative dominance occurred at the ripening stage from  
345 the former bacteria-based energy channel to a fungal-based energy channel under eCO<sub>2</sub> (Fig.  
346 S2). Nevertheless, the fungal-bacterial ratios in the paddy rice system were smaller than those  
347 of grasslands (Hungate et al., 2000). The lower proportion of fungal biomass in paddy rice  
348 may belie their importance, as the abundance of fungivorous nematodes under eCO<sub>2</sub> was

349 twice that of aCO<sub>2</sub> (Fig. 5f). Amoebae showed a clear connection to the fungal channel ( $p =$   
350 0.07; Fig. 6), supporting evidence of strong trophic links between amoebae and fungi in soils  
351 (Chakraborty et al., 1985; Geisen et al., 2016). Also, previous studies found that eCO<sub>2</sub>  
352 increased saprotrophic fungi (Drigo et al., 2007) as well as arbuscular mycorrhizal fungi  
353 (Drigo et al., 2013), but our results indicate that the plant's energy shunt to either bacteria or  
354 fungi is dynamic and switches among growth stages (Dunfield & Germida, 2003; Mougel et  
355 al., 2006; Houlden et al., 2008).

356

#### 357 *4.2. Interaction between rice cultivars and eCO<sub>2</sub> and eN determined herbivore load*

358 Most previous studies on eCO<sub>2</sub> and eN effects on soil food web interactions focused on the  
359 decomposer food web and only a few studies included herbivores or parasitic microbes (Ayres  
360 et al., 2008; Cesarz et al., 2015; Chen et al., 2015). Plant parasitic nematodes can cause  
361 significant yield losses cereal production systems, including rice topping systems (Bridge et  
362 al., 2005), but until now they received limited attention (Liu et al., 2008b; Huang et al., 2015),  
363 most probably due to their hidden form of herbivory (Johnson et al., 2016).

364 Elevated CO<sub>2</sub> levels may affect root herbivores in different ways. On the one hand, the  
365 increased biomass and growth rate of rice roots under eCO<sub>2</sub>, particularly for the  
366 strong-responsive cultivar (Yang et al., 2008; Zhu et al., 2013), will increase food supply to  
367 root herbivores. On the other hand, eCO<sub>2</sub> could adversely affect the herbivores via reduced  
368 food quality (i.e. wider C:N ratio) (Norby & Cotrufo, 1998; Reich et al., 2006), even when  
369 high N was supplied (Sinclair et al., 2000). Therefore, improving crop yield by the selection  
370 of cultivars positively responsive to elevated CO<sub>2</sub> levels (Brummer et al., 2011; Ziska et al.,

371 2012) might mitigate against the predicted effects of future climate change (Cramer et al.,  
372 2001; Olesen & Bindi, 2002). However, our data clearly show that the positive response of  
373 rice cultivars to elevated CO<sub>2</sub> might come at a cost of increased herbivore load.

374 The doubling of the abundance of herbivorous nematodes with eN under the  
375 weak-responsive cultivars at the heading stage (Fig. 5c), and their reduction to control levels  
376 under eCO<sub>2</sub> could be explained by this reduced food quality for herbivores. Accordingly, the  
377 weak-responsive rice with its reduced root biomass and lower food quality supported reduced  
378 levels of herbivorous nematodes under eCO<sub>2</sub>. In contrast, the performance of the  
379 strong-responsive cultivar to eCO<sub>2</sub> is highly dependent on N fertilization (Zhu et al., 2015).  
380 This led to improved resource quality (e.g. increased root N content and reduced C:N ratio)  
381 and quantity (e.g. root biomass) for herbivores (Fig. S4). Thus, the strong-responsive cultivars  
382 face a trade-off between N-limitation and herbivore load, especially if N-limitation is  
383 counterbalanced by fertilization.

384 Our results clearly demonstrate that management strategies intended to mitigate negative  
385 climate change effects on crops can lead to conditions conducive to plant parasite outbreaks.  
386 Therefore adopting high-yielding crop cultivars adapted to climate stress without taking into  
387 account root resistance to herbivores may imperil future crop production, and other global  
388 change factors, such as warming, may even exacerbate this effect (DeLucia et al., 2012).

389 Breeding new crop cultivars with improved resource use efficiency to satisfy food  
390 demand, as well as controlling invasive weeds and pathogens, is one of the most promising  
391 practices for agronomists under the pressure of the ongoing climate change (Bender et al.,  
392 2016; Brummer et al., 2011; Hirel et al., 2007). However, our findings clearly show that

393 highly productive cultivars under eCO<sub>2</sub> may raise pest infestation rates, suggesting an  
394 unexpected trade-off that would generate long-term negative soil feed backs. Cultivars with  
395 comprehensive traits should be taken into account in future integrated crop management  
396 (Huang et al., 2012; Tiemann et al., 2015).

397

## 398 **5. Conclusions and outlook**

399 Our results showed differential responses of soil microbes and microbivores to eCO<sub>2</sub> and N  
400 inputs at different growing stages of rice. These results illustrated the highly temporal-dynamic  
401 nature of soil micro-food web responses to the changing climate conditions and call for caution  
402 in extrapolating results from single sampling time and/or single trophic level to predict the  
403 long-term impact of climate change factors on the soil micro-food web. Also, the interactive  
404 effect of eCO<sub>2</sub>, eN and cultivars on soil food web indicated that alteration in resource  
405 availability to microbes can cascade up along the food web. eCO<sub>2</sub> in general increases the C:N  
406 ratio of plant materials and thus reduces the quality for herbivores, which are often assumed to  
407 have negative effects on herbivorous nematodes and other pest insects. However,  
408 strong-responsive cultivar was susceptible to root-feeding pests under elevated CO<sub>2</sub> and N  
409 fertilization, thus rendering the long-term advantage of breeding positively CO<sub>2</sub>-responding  
410 cultivars questionable. Regarding to agricultural managements under future climate change  
411 scenarios, this study highlights crop breeding strategies should integrate knowledge about the  
412 architecture and metabolic footprints of soil food web.

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422

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673 **Table 1.** *F* value of repeated measures ANOVA on the effects of CO<sub>2</sub> (ambient [aCO<sub>2</sub>] and elevated [eCO<sub>2</sub>]), N (ambient [aN] and exogenous  
674 [eN]), Cultivar (weak- and strong-responsive) and all possible interactions on the biomass of microorganisms and the abundance of protists,  
675 nematodes and nematode trophic groups across sampling times (T) at heading and ripening growth stage.

	Microbial biomass	Bacterial biomass	Fungal biomass	Flagellates	Amoebae	Total nematode	Bacterivores	Fungivores	Herbivores	Omnivores- carnivores
eCO <sub>2</sub>	1.47	0.63	5.02*	3.08	4.82*	4.98*	0.01	4.56*	1.96	10.89**
eN	2.21	1.75	3.65	35.93**	0.53	51.80**	19.79**	0.46	29.90**	20.89**
Cultivar	0.84	0.85	1.48	0.18	0.09	20.31**	0.16	0.10	24.56**	4.91*
eCO <sub>2</sub> × eN	0.40	0.17	0.15	12.92**	0.05	6.41*	6.35*	0.65	1.27	7.97*
eCO <sub>2</sub> × Cultivar	0.07	0.07	0.00	7.34*	0.43	29.14**	0.03	0.02	58.39**	1.94
eN × Cultivar	0.04	0.00	0.09	5.99*	0.01	9.88**	3.45	8.81**	5.19*	1.61
eCO <sub>2</sub> × eN × Cultivar	0.01	0.12	0.47	5.23*	0.58	1.48	10.70**	4.91**	1.13	2.57
T	51.17**	50.32**	48.22**	8.46*	0.34	45.81**	42.24**	23.87**	4.99*	45.81**
T × eCO <sub>2</sub>	1.85	0.46	4.50*	5.01*	5.87*	17.10**	0.09	13.21**	11.38**	17.10**
T × eN	0.02	0.01	1.33	0.12	0.91	0.30	0.32	1.19	0.24	0.30
T × Cultivar	2.57	3.12	0.81	0.07	0.19	13.09**	0.24	3.78	6.45*	13.09**
T × eCO <sub>2</sub> × eN	0.35	0.41	0.09	2.63	5.45*	9.32**	1.23	2.38	10.97**	9.32**
T × eCO <sub>2</sub> × Cultivar	0.22	0.32	0.08	0.003	0.49	31.91**	7.17*	2.89	18.91**	31.91**
T × eN × Cultivar	0.76	0.67	0.04	0.16	1.65	0.2	1.61	1.76	2.13	0.20
T × eCO <sub>2</sub> × eN × Cultivar	0.42	0.72	0.01	0.56	1.26	38.63**	9.58**	6.96*	19.70**	38.63**

676 \*, \*\* indicates factors effect significant at  $p < 0.05$ ,  $p < 0.01$ , respectively

677 **Figure captions**

678 **Fig. 1** Photograph showing one of the three FACE rings in Zongcun Village (119°42'0" E,  
679 32°35'5" N), Yangzhou City, Jiangsu Province, China.

680 **Fig. 2** The content of soil resources in varying CO<sub>2</sub> (ambient [aCO<sub>2</sub>] and elevated [eCO<sub>2</sub>]), N  
681 (ambient [aN] and exogenous [eN]), and Cultivar (weak- and strong-responsive) treatment (a:  
682 Dissolved organic carbon, n = 12; b: Dissolved organic nitrogen, n = 12 and 6 at the heading  
683 and ripening stage, respectively; c: Mineral nitrogen, n = 12). Only significantly different  
684 results are presented, and see Table S2 and S3 for all data of soil resources. Means with  
685 different letters indicate significant difference among treatments (Fisher's LSD test,  $p < 0.05$ ).  
686 Error bars are standard errors.

687 **Fig. 3** The biomass of the overall microbial community, bacteria and fungi in varying CO<sub>2</sub>  
688 (ambient [aCO<sub>2</sub>] and elevated [eCO<sub>2</sub>]), N (ambient [aN] and exogenous [eN]), and Cultivar  
689 (weak- and strong-responsive) treatment (a-c: Heading stage; d-f: Ripening stage). Means  
690 with different letters indicate significant difference among treatments (Fisher's LSD test,  $p <$   
691 0.05). Error bars are standard errors (n = 3).

692 **Fig. 4** The abundance of flagellates and amoebae in varying CO<sub>2</sub> (ambient [aCO<sub>2</sub>] and  
693 elevated [eCO<sub>2</sub>]), N (ambient [aN] and exogenous [eN]), and Cultivar (weak- and  
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697 **Fig. 5** The abundance of nematode trophic groups in varying CO<sub>2</sub> (ambient [aCO<sub>2</sub>] and  
698 elevated [eCO<sub>2</sub>]), N (ambient [aN] and exogenous [eN]), and Cultivar (weak- and

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701 bars are standard errors ( $n = 3$ ).

702 **Fig. 6** Structural equation modeling (SEM) analysis of the elevated CO<sub>2</sub> and N fertilization  
703 effects on soil micro-food webs at the ripening stage in a rice field in Jiangsu province, China.  
704 The results of the optimal model fitting [Chi-square ( $\chi^2$ ) = 21.755,  $df = 10$ ,  $p = 0.061$ ,  
705 comparative fit index (CFI) = 0.937, root square mean error of approximation (RMSEA) =  
706 0.221]. Square boxes denote variables include in the models. Values associated with solid and  
707 dashed arrows represent standardized path coefficients. Percentages close to variables indicate  
708 the proportion of variation explained by the model ( $R^2$ ). Solid arrows denote the directions  
709 and effects that were significant ( $p < 0.05$ ) and the thickness represents the magnitude of the  
710 path coefficients. Dashed arrows represent the directions and effects were non-significant ( $p >$   
711 0.05). (ExtN; extractable N, the sum of NH<sub>4</sub><sup>+</sup>-N, NO<sub>3</sub><sup>-</sup>-N and DON; BF: bacterivores; FF:  
712 fungivores; HE: herbivores; OC: omnivores-carnivores; Flag: flagellates; Amoe: Amoebae)

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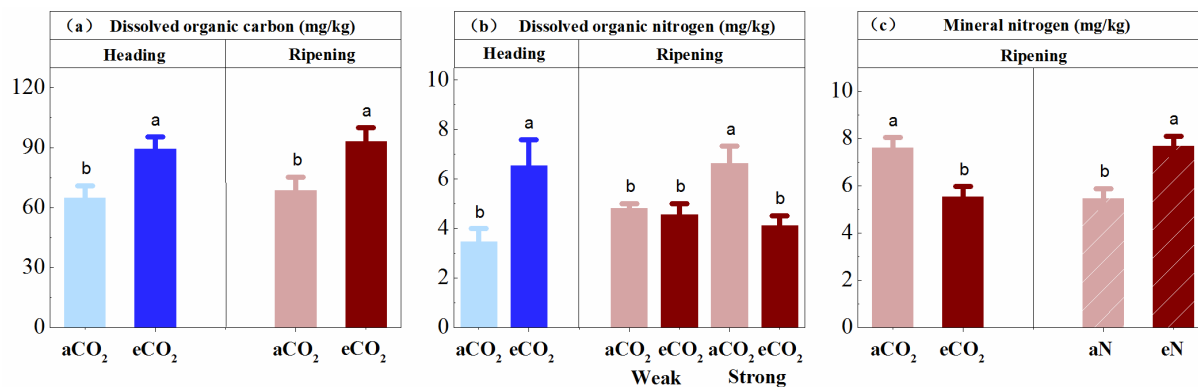
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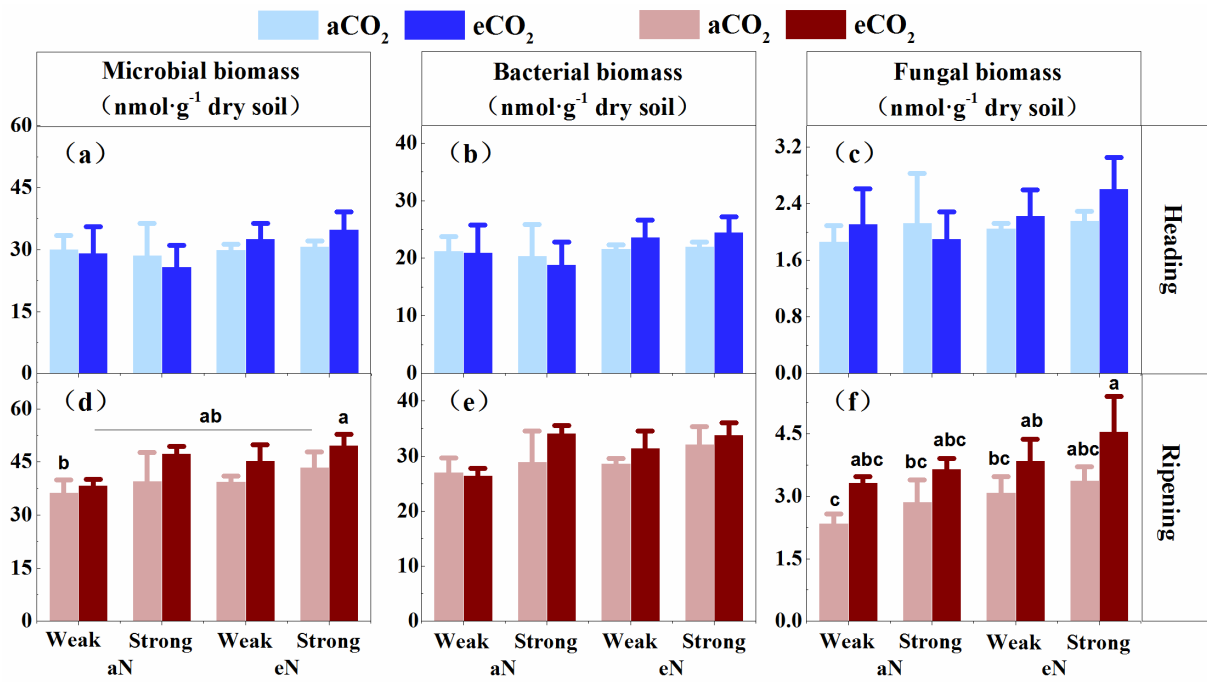


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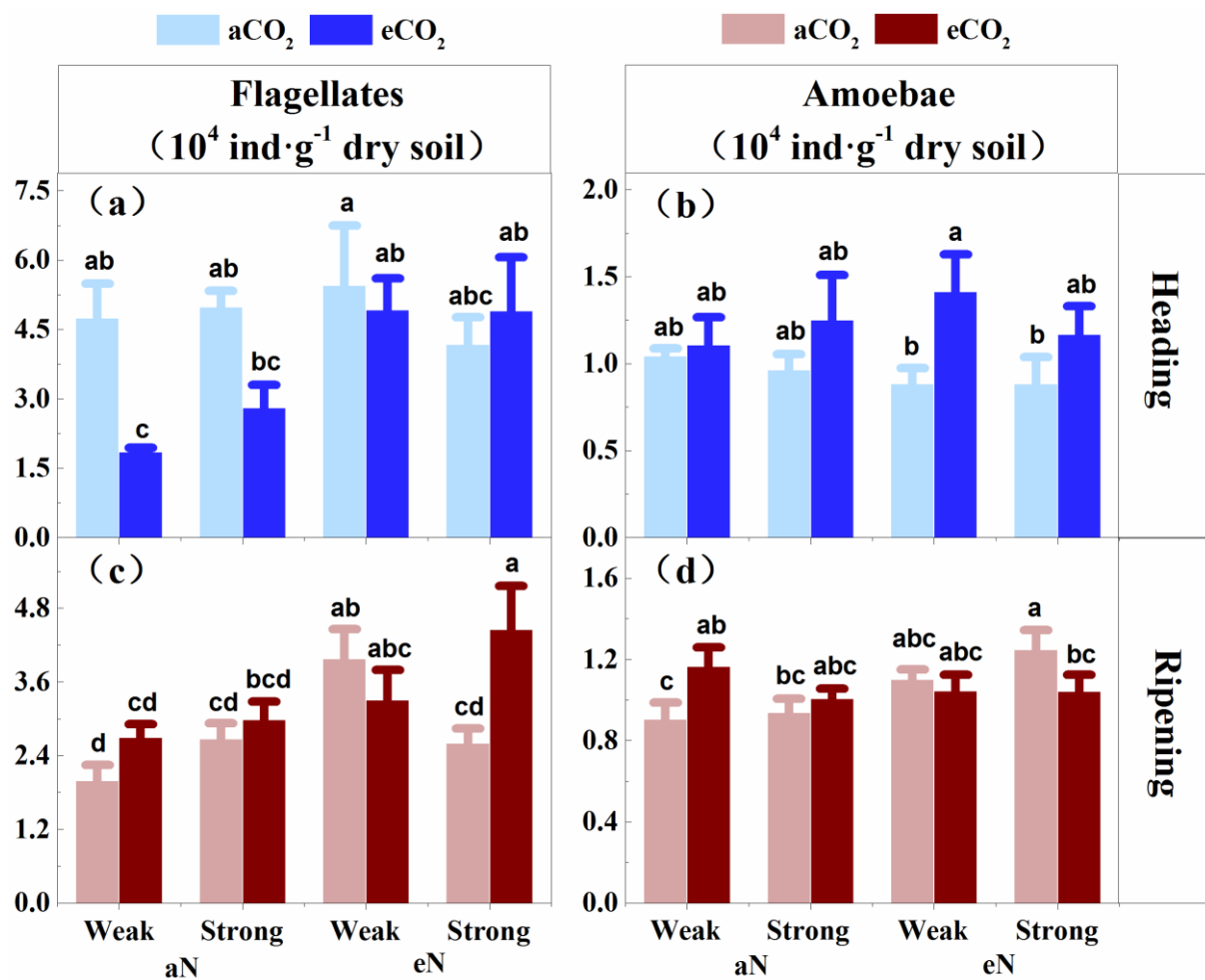


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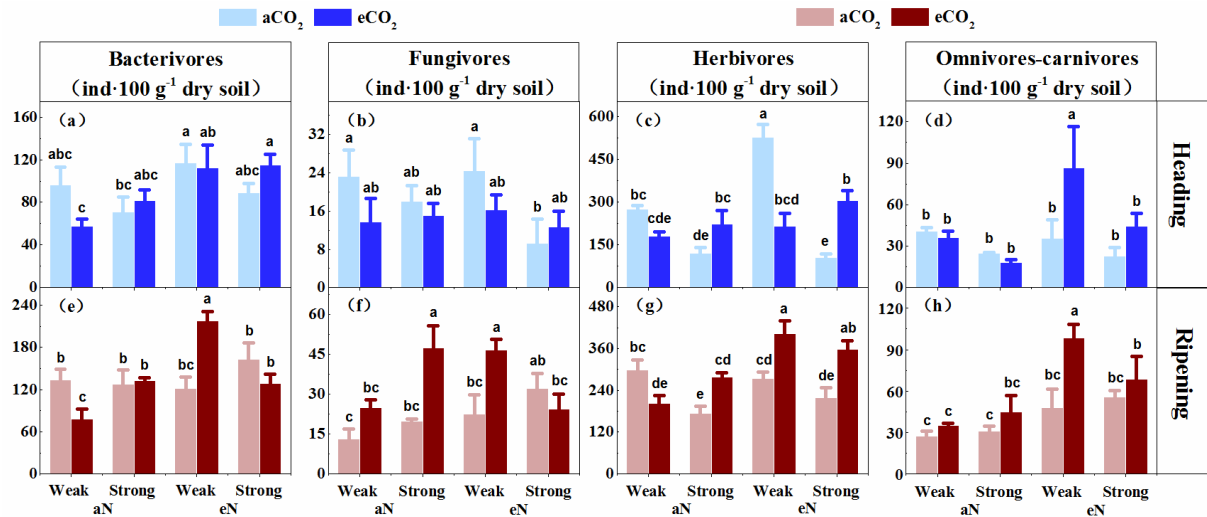
741 letters indicate significant difference among treatments (Fisher's LSD test,  $p < 0.05$ ). Error

742 bars are standard errors ( $n = 3$ ).

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746 **Fig. 5** The abundance of nematode trophic groups in varying CO<sub>2</sub> (ambient [aCO<sub>2</sub>] and

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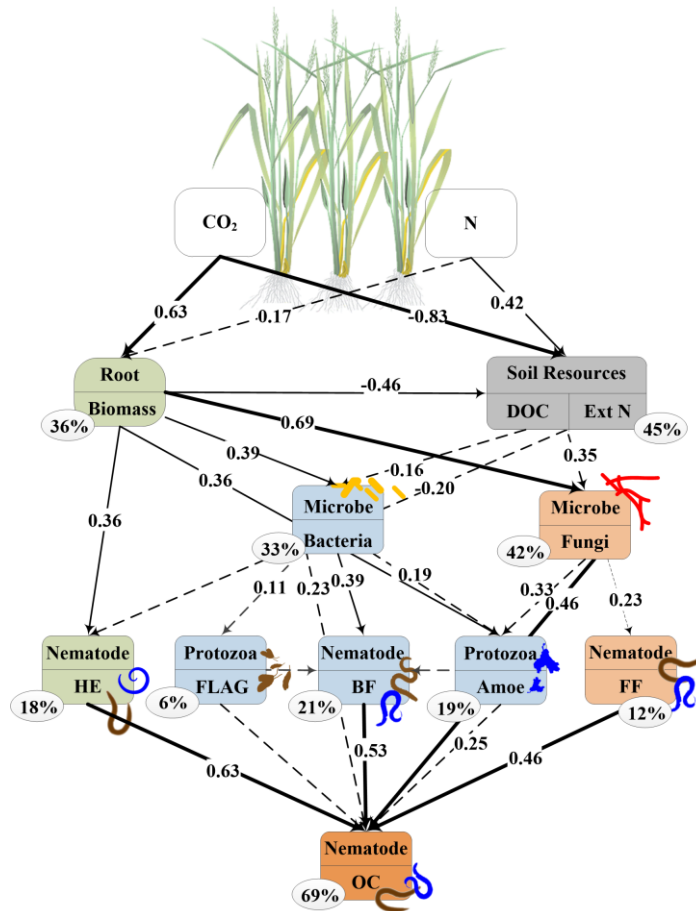
748 strong-responsive) treatment (a-d: Heading stage; e-h: Ripening stage). Means with different

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