1	Soil priming effects and involved microbial community along salt gradients
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19 Abstract

20 Soil salinity mediates microorganisms and soil process, like soil organic carbon (SOC) cycling. Yet, how soil salinity affects SOC mineralization via shaping bacterial communities diversity and 21 22 composition remains elusive. Therefore, soils were sampled along a salt gradient (salinity at 0.25%, 23 0.58%, 0.75%, 1.00% and 2.64%) and incubated for 90 days to investigate i) SOC mineralization (i.e., 24 soil priming effects induced by cottonseed meal, as substrate) and ii) responsible bacteria community, by using high throughput sequencing and natural abundance ¹³C isotopes (to partition cottonseed meal 25 26 derived CO₂ and soil derived CO₂). We observed negative priming effect during first 28 days of 27 incubation and turned to positive priming effect after day 56. Negative priming at the early stage might 28 be due to the preferential utilization of cottonseed meal. The followed positive priming decreased with 29 the increase of salinity, which might be caused by the decreased alpha diversity of microbial 30 community in soil with high salinity. Specifically, soil pH and EC along salinity gradient were the 31 dominant variables modulating the structure of microbial community and consequently SOC priming 32 (estimated by distance-based multivariate analysis and path analysis). By adopting O2PLS, priming 33 effects were linked with specific microbial taxa, e.g., Proteobacteria (Luteimonas, Hoeflea and 34 Stenotrophomonas) were the core microbial genus that attributed to the substrate induced priming effects. Here, we highlight that the increase of salinity reduced the diversity of microbial community 35 36 and shifted dominant microorganisms(Actinobacteria and Proteobacteria (Luteimonas, Hoeflea and 37 Stenotrophomonas)) that determined SOC priming effects, which provides a theoretical basis for understanding of SOC dynamics and microbial drivers under salinity gradient. 38

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40 Keywords: Salt gradient, priming effects, bacterial community, core microorganisms

41 **1. Introduction**

42 Soil salinization is an increasing environmental problem caused by natural and 43 human activities in the arid and semi-arid area (Wichern et al., 2006). Salinization is 44 often a major threat to crop productivity in agricultural land. Soil microorganisms suffer 45 from osmotic stress. Soil salinity often cause microbial death or dormant. It was widely 46 reported that the increased salinity decrease microbial biomass, enzymatic activity, and 47 alpha diversity of microbial community (Laura, 1974; Pathak et al., 1998; Rietz et al., 48 2003). Soil salinity is reported to the major determinants of composition, activity of 49 microbial community (Kamble et al., 2014). Although salinity is reported to be a vital 50 factor in influencing microorganisms in the arid and semi-arid area, limited studies 51 investigated C processes (e.g., priming effect) driven by microbial community in 52 salinity soils (Sardinha et al., 2003).

53 Soil organic carbon (SOC) is the largest pool (1500 Pg C) in the terrestrial carbon 54 (C) cycle, and contains twice as much C as the atmosphere (Filley et al., 2006; 55 Wiesmeier et al., 2019). The input of substrate C can influence the output (i.e., CO₂ 56 release) through a phenomenon called priming effect, which was firstly discovered by 57 LÖhnis (1926). Substrate additions accelerate or decrease soil organic C mineralization, 58 referred to positive or negative priming effects (Kuzyakov et al., 2000). The intensity 59 of the priming effect affects the turnover of SOC and thus storage pool (Sullivan et al., 60 2013). Soil priming effects are affected by many biotic and abiotic factors (Lavelle, et 61 al.;; Martin et al., 2019), to investigate abiotic and biotic mechanisms underlying SOC 62 priming enhance strong understanding of the SOC cycling.

Soil priming effects is affected by soil fauna animals (Scheu et al., 1994), activities,
diversity and composition of microbial community (; Fontaine et al., 2011). The
microbial decomposers are the major player in the decomposition process of added C
sources. The addition of substrate, such as composts (Xun et al., 2016), animal sludges
(Hartmann et al., 2015), sewage sludges (Su et al., 2017; Wagner et al., 2011) and plant
residues (Dai et al., 2017), generally increases soil microbial biomass C and stimulates

the microbial activities thus enhanced the loss of SOC (positive priming effects)
(Fontaine et al., 2003; Bird et al., 2011; Li et al., 2018; Ali et al., 2019).

71 Concerning abiotic factors, the priming effect can be controlled by climate 72 variables (Hagemann, 2008), and soil properties, like pH, EC, TN, etc (Blagodatskaya 73 et al., 2008; Luo et al., 2017). To understand how environmental and edaphic factors 74 affect the processes of SOC mineralization, is important to estimate terrestrial C pool (Lehmann et al., 2015). Although many studies have tested the effects of soil pH, SOC 75 76 content, and other edaphic variables on soil priming effect, few study investigated soil 77 priming effects in salinity soil (Asghar et al., 2012), especially linked with soil 78 microbial community structure and their functions in C decomposition (Soina et al., 79 2018).

80 Thus, we sampled the soils along natural salinity gradients (0.25%, 0.58%, 0.75%), 81 1.00%, 2.64% apart from total water-soluble salt). Based on these soils, we conducted a 90 days of indoor incubation applying C3 substrate of cottonseed meal (δ^{13} C=-82 23.47‰) to C4 soils with salt gradient (δ^{13} C between -14.21‰ and -16.01‰), to 83 84 investigate: 1) mineralization rate of cottonseed meal and induced soil priming effects 85 along salt gradients; 2) diversity of microbial community in the soils with increased 86 salinity, and 3) identify the bacteria taxa associated Soil priming. We hypothesized that 87 i) soil microbial community diversity and composition will be different with the 88 different in soil variables particularly pH and EC along salinity gradients, and ii) Soil 89 C processes like priming effects will be regulated mainly by microbial community and 90 especially the core microbial species. To clarify the priming effects and involved 91 microbial groups would help us better understanding C sequestration potential and 92 underlying mechanisms in saline soils.

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95 2. Materials and methods

96 2.1. Soil sampling and cottonseed meal production.

97 The soil type was gray desert soil, which was collected from farmlands (82.90° E, 98 44.96° N) in Xiao Yinpan town, Bole City, Bortala, northern Xinjiang Uygur 99 Autonomous Region, northwest China. The farmlands soil is naturally formed original 100 saline-salinity soil and with a continuous 30 years planting of maize (C4 crop) and 101 maize straw returning to soil for 7-8 year. In September 2021, we determining the 102 sampling area, and use the five-point sampling method to collecting non-rhizosphere 103 soil. The soil samples were indoor air drying and hand-picked to remove visible other 104 debris, animal and plant residues and then sieved at field moisture (<2mm) and 105 subsequently adjusted to 40% of water holding capacity (WHC). Determination of 106 five salinity gradients at 0.25%, 0.58%, 0.75%, 1.00% and 2.64% through soil salinity 107 measurements. Texture was determined by the pipette method without carbonate in all soil samples. They were then incubated at 25 °C for 7 days before starting the 108 109 experiments, to allow any early sampling and sieving effects to subside.

110 Cottonseed meal is a kind of reddish or yellow granular material obtained by 111 pressing, leaching and other cottonseed. The cottonseed meal was purchased from the 112 market and dried at 105 °C for 24 h indoor, then further pulverized by a ball mill and 113 passed through < 2 mm sieve.

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115 2.2. Soil and substrate analyses

116 EC and pH of soil and cottonseed meal were measured at a soil: water ratio of 1:5 117 (weight/weight) (Bao, 2000). Air-dry soil (5 g, <2 mm) and 25 ml of deionised water 118 were shaken together for 1 min and left to settle for 30 min, which was repeated once 119 more before pH was determined with a pH electrode. Soil water-soluble salt was 120 analyzed by weighted at a soil:water ratio of 1:5 (weight/weight). Air-dry soil (5 g, <1 121 mm) and 25 ml of deionised water were shaken together for 30 min, filtration to obtain 122 clear filtrate, using thermostat water bath to evaporate and weigh(Bao, 2000). Soil total 123 carbon (TC), total nitrogen (TN) are collect soil to be tested was dried and ground

124 through a 0.15mm screen, and a certain amount of treated soil sample was wrapped in 125 tin foil and placed in an element analyzer for determinatio (air-dried, milled <150 µm) 126 were determined by dry combustion (LECO CNS 2000, LECO Corporation, Michigan, 127 USA). Soil microbial biomass C was determined by fumigation extraction (Vance et 128 al., 1987; Wu et al., 1990). The K₂SO₄ extractable organic C was determined using an 129 organic carbon autoanalyser (Shimadzu, Analytical Sciences, Kyoto, Japan). Soil 130 microbial biomass C (Bc) was calculated from: Bc = 2.22 Ec, where Ec = [(organic C131 extracted from fumigated soil) minus (organic C extracted from non-fumigated soil)]. The natural δ^{13} C (‰) abundance of the soils (air-dried, milled <200 µm) was 132 133 determined using an elemental analyser-isotope ratio mass spectrometer (Sercon Ltd, 134 Crewe, UK). All measurements are given on an oven-dry weight basis (o.d., 105 °C, 24 135 h).

136 The δ^{13} C (‰) abundance of the cottonseed meal (air-dried, milled <200 µm) was 137 determined using an elemental analyser-isotope ratio mass spectrometer (Sercon Ltd, 138 Crewe, UK). The main elemental composition of the substrate was determined using 139 elemental analysis (Vario EL Cube, Hanau, Germany), with the samples combusted at 140 1200 °C. Natural δ^{13} C (‰) abundance ,the total carbon, total nitrogen contents and C/N 141 of the cottonseed meal was presented in Table 1.

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143 2.3. Experimental design

After pre-incubation, five soils with salinity gradient(salinity at 0.25%, 0.58%, 0.75%, 1.00% and 2.64%) were thoroughly mixed with cottonseed meal at 20 mg C g⁻¹ soil (d.w. basis), and incubated over 90 days following moisture adjustment to 40% of water-holding capacity (WHC) to investigate the substrate mineralization and priming effects. Each soil sample (40 g d.w. basis) was incubated in a 100 ml beaker inside a 1 L brown glass jar. Three jars with only water and NaOH were set as blank. All the jars were sealed with a rubber bung and incubated in a randomized block design 151 at 25 °C for the 90 days of incubation. The NaOH vials were changed after 1, 3, 5, 7,

152 14, 28, 56 and 90 days for determination of evolved CO_2 and ${}^{13}C-CO_2$ (‰). Meanwhile,

soil biomass C, NH_4^+ , NO_3^- , pH, EC, TC, TN and DNA extraction were measured at day 28.

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156 2.4. Soil CO₂-C and its isotopic composition

Soil C evolved as CO₂-C in jars was measured by trapping CO₂ in 1 M NaOH 157 158 (20 ml) during soil incubation. After the NaOH (20 ml) trapping CO₂ at different 159 periods of soil incubation, 5 ml 1 M NaOH of each sample was mixed with 10 ml 160 deionised water and titrated with 0.05 M standardised HCl by the TIM840 autotitrator (Radiometer Analytical, Villeurbanne Cedex, France). Meanwhile, the δ^{13} C (‰) of 161 trapped CO₂-C was precipitated, with 8 ml of the 1 M NaOH (20 ml) mixed with 8 ml 162 163 1.5 M BaCl₂ in vials (Aoyama et al., 2000). The BaCO₃ precipitate was trapped on the 164 glass fibre the filter, rinsed with deionised water several times, and dried overnight (80 °C), weighed (0.100-0.200 mg) into tin capsules, and analyzed for δ^{13} C on an 165 166 elemental analyzer-isotope ratio mass spectrometer (Sercon Ltd, Crewe, UK).

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168 2.5. DNA exaction and sequencing

The total soil DNA was extracted from 0.50 g of moist soil using a FastDNA Spin
Kit (MP Biomedicals, Santa Ana, CA, USA) according to the manufacturer's protocol.
The extracted DNA was dissolved in 50 μl of TE buffer, quantified using a
spectrophotometer and stored at -20 °C until sequencing.

173 V3-V4 hypervariable regions of the bacterial 16S rRNA gene were amplified with 174 primers 341F (5'-CCTAYGGRBGCASCAG-3') and 806R(5'-175 GGACTACHVGGGTWTCTAAT-3'). The PCR reactions were conducted with a thermocycler PCR system (GeneAmp 9700, ABI, USA) by using the following 176 programs: 3 min of denaturation at 95 °C; followed by 27 cycles of 30 s at 95 °C, 30 s 177 at 55 °C, and 45 s at 72 °C; and a final extension at 72 °C for 10 min with a thermocycler 178

PCR system (GeneAmp9700, ABI, USA). PCR amplicons pooled from the triplicate
reactions were purified using a QIAquick PCR purification kit (Qiagen, Shenzhen,
China), and quantified using a NanoDrop ND-1000 spectrophotometer (Thermo
Scientific, Waltham, MA, USA). The PCR products were purified, mixed, and sent to
Majorbio, Inc. (Shanghai, China) for sequencing based on the Illumina MiSeq platform.

185 2.6. Calculations

186 2.6.1. CO_2 - $\delta^{13}C$ emission

187 The mineralisation of cottonseed meal was separated from SOC mineralisation 188 according to the change of stable isotopic composition ($\delta^{13}C$) with time. The standard 189 equation for determining $\delta^{13}C$ (‰) is derived from:

190
$$\delta^{13}$$
C (‰) = [(R_{sample}/R_{VPDB}) - 1] × 1000,

191 where R_{sample} is the mass ratio of ¹³C to ¹²C of each sample and R_{VPDB} is the 192 international PDB(Peedee Belemnite) limestone standard. The labeled ¹³C (%) of 193 cottonseed meal was then estimated from:

Eqn. 1

194
$$CO_2^{-13}C(\%) = (\delta_{treatment} - \delta C4) / (\delta C3 - \delta C4),$$
 Eqn. 2

195 where CO_2 -¹³C (%) is the proportion of evolved CO_2 from C3 (cottonseed meal) 196 matter, $\delta_{treatment}$ is the $\delta^{13}C$ (‰) in treatments of soil with cottonseed meal, $\delta C4$ is the 197 $\delta^{13}C$ (‰) in control soil and $\delta C3$ is the $\delta^{13}C$ (‰) from cottonseed meal. Thus, the CO₂-198 C produced from cottonseed meal during the incubation was calculated from:

199
$$CO_2^{-13}C (\mu g g^{-1} \text{ soil}) = CO_2^{-13}C (\%) \times \text{total } CO_2^{-C} (\mu g g^{-1} \text{ soil})/100, \text{ Eqn. 3}$$

200 CO_2 from SOC was CO_2 -¹³C subtracted from total evolved CO_2 -C. The absolute 201 soil priming effect (or primed soil CO_2 -C) with the addition of cottonseed meal was 202 calculated from:

203 Primed soil CO₂-C (
$$\mu$$
g C g⁻¹ soil) = CO₂-C_{treatment} - CO₂-C_{control} Eqn. 4

where CO_2 - $C_{treatment}$ is the non-isotopically labeled CO_2 -C evolved from cottonseed meal amended soil, CO_2 - $C_{control}$ is non-isotopically labeled CO_2 -C evolved from soil without cottonseed meal.

208 2.7. Statistics

The data of ¹⁶S gene sequencing were processed using the Quantitative Insights 209 210 Into Microbial Ecology (QIIME) 1.9.0-dev pipeline (Caporaso et al., 2010). In brief, 211 Reads with less than length 200 bp and ambiguous bases were discarded. The sequences 212 were then binned into operational taxonomic units (OTUs) by UCLUST (Edgar, 2010) 213 based on 97% pairwise identity. Chimeric OTUs identified by USEARCH (Edgar et al., 214 2011) in QIIME were removed. The most abundant sequence from each OTU was 215 selected to represent that OTU. Taxonomy was assigned to 16S OTUs against a subset 216 of the Silva 104 database. The representative OTU sequences were aligned using 217 PyNAST (Caporaso et al., 2010). We obtained between 64,425 and 89,989 clean_reads 218 per sample for all experimental samples.

219 To avoid potential bias caused by sequencing depth, all sample datasets were 220 rarefied for the bacteria α -diversity and β -diversity analyses. Faith's phylogenetic 221 diversity was calculated to provide an integrated index of the phylogenetic breadth 222 across taxonomic levels (Faith, 1992). To compare β -diversity between samples, principal coordinate analyses based on the unweighted and weighted UniFrac 223 224 (Lozupone et al., 2007a) distances were calculated using the function 'pcoa' in the R 225 package 'Ape'. Additionally, permutational multivariate analysis of variance 226 (PERMANOVA) was carried out using the function 'adonis' in the R 'vegan' to 227 measure effect size and significance on β -diversity. The variable influence projection (VIP) value was processed using the way of O2PLS analysis by the SIMCAP 14 228 229 (Version 14.1.0.2047) (Wang et al., 2016). The y-matrix was defined as the 230 environmental factors datasets and the x-matrix was defined as the microbial 231 community on genus level dataset.

Data were logarithmically transformed and analyzed by ANOVA. All analyses were performed using SPSS software (13th edition). Pearson's correlation analyses were performed to assess the linear correlation among soil physio-chemical properties and 235 microbial community. MULTIVARIATE analysis were operated to investigate236 interaction of salinity treatments on bacteria community parameters.

237

238 **3.** Results

239 3.1. Soil physicochemical properties along salt gradients

240 The major soil physicochemical properties along salt gradients were presented 241 (Table 1) and all of soil physicochemical properties has significant difference (P < P242 0.05). The total soluble salinity content in the soils ranged from 0.25% to 2.64% of 243 salinity soils, soil salt gradients increasing gradually from salinity 1 samples to salinity 244 5 samples. The pH and EC in soils ranged from 8.45 to 8.85 and from 1.06 ms cm⁻¹ to 245 7.75 ms cm⁻¹. Soil total C and N were increased with salinity, ranging from 3.16% to 3.57%, and from 0.18% to 0.26%. The δ^{13} C value for soils are between -14.21‰ and -246 16.01‰, which were relatively enriched compared to cottonseed meal (-23.47‰). This 247 allowed separation of soil derived CO₂ from total evolved CO₂, according to the classic 248 249 mixed modeling.

250

251 3.2. Total CO₂ evolution

During the whole 90 days of incubation, the cumulative CO_2 evolved had similar trends, which the amount of CO_2 increased with the incubation times (Fig. S1). The cumulative CO_2 evolved increased more rapidly with the addition of cottonseed meal before 14 days, compared to non-amended soils. At 90 days of incubation. The cumulative CO_2 evolved in the soil with the lowest salinity (Salinity 1) gave the lowest CO_2 emission (597 µg C g⁻¹) in the non-amended soils (Fig. S1, P < 0.001).

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259 3.3. Cottonseed derived ${}^{13}CO_2$ and soil priming effects

260 The total cumulative CO_2 -C was divided three parts based the $\delta^{13}C$ value, 261 including basal soil-derived CO_2 , cottonseed meal-derived CO_2 and primed soil CO_2 262 (Fig.1). The cottonseed meal-derived CO_2 had a significant contribution to the total CO_2 evolved during the early incubation period. The cottonseed meal-derived CO_2 was significantly higher in Salinity 1, Salinity 2 and Salinity 3 than in Salinity 4 and Salinity 5 before 28 days incubation. Meanwhile, the soil priming effects was negative in all amended soil treatments before 28 days incubation and the direction of priming effect in most of soil samples turned into positive after 28 days. During the whole 90 days incubation, there was a negative correlation between cottonseed meal-derived CO_2 and primed soil CO_2 (Fig. 2).

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271 3.4. Bacterial diversity and community structure

272 The number of sequences ranged from 64,425 to 91,261 for per sample (average 273 valve of 80,602). About 27,990 OTUs in total were obtained under different five 274 treatments. Bacterial community diversity was measured by a series of OTU-based 275 analyses of alpha diversity including chaol estimator, and observed species in the 276 QIIME pipeline (Fig. 3). Chao1 diversity estimator and observed_species was 277 significantly different in treatments, being the highest in Salinity 1, followed by Salinity 278 3, Salinity 2, Salinity 4 and Salinity 5 (P < 0.01). In general, bacterial community 279 diversity decreased with increasing salinity (Fig. 3).

The most abundant phylum in the soils and their correlation with salinity were shown in Fig. 4. Among them, *Actinobacteria* was the dominant taxa in all soils, with the abundance ranging from 50.07 % (Salinity 3) to 68.99 % (Salinity 4). The relative abundance of *Bacteroidetes*, *Firmicutes*, and *Deinococcus-Thermus* increased with the salinity, while *Acidobacteria* decreased with salinity degree.

Based on OTUs of five gradient salt treatments, the PCA analysis showed that treatments from Salinity 2 and Salinity 4 clustered together. Meanwhile, soil samples of Salinity 1, Salinity 3 and Salinity 5 distributed in the first, fourth and three quadrant, which indicated that these treatments had large environmental heterogeneity (Fig. S4). In order to visualize the relationship between environmental factors and microbial community, *Canonical Correspondence Analysis* (CCA) was conducted, showing that NO₃⁻-N, EC and TC had a more obvious impact than other factors for microbial community (Fig. 3). Soil EC were positively correlated with pH, NH₄⁺-N, and negatively correlated with TN, TC and MBC. Mantel test and Distance-based multivariate analysis showed the contribution rate of different environmental factors account for 78% of the variability of microbial communities (Table 2). The value of pH (31%) and EC (12%) had a strong influence on microbial community.

297

298 3.5. Relation between soil microbial community and C dynamics

299 Based on the O2PLS analysis, the variable influence projection (VIP) values of 300 bacterial genus more than 1.00% were showed their contributions to C decomposition 301 of cottonseed meal-derived C, basal soil-derived C, and primed soil C (Table 3). There 302 were many microbial taxa positively correlating to soil primed CO₂, for insatnce, genera 303 Actinomarinales, Nocardioides, of Luteimonas, Hoeflea, Intrasporangium, 304 Nitrolancea, Pseudarthrobacter and Stenotrophomonas had a positive correlation with 305 primed CO₂. In order to further to evaluate the relationship between soil properties, soil 306 bacterial communities and C decomposition, we used the structural equation modeling 307 (SEM) to suggest the direct and indirect impacts of salinity and microbial community 308 on soil C decomposition (Fig. 7). The result showed that soil pH and EC had negative 309 contribution to bacterial diversity, while bacterial diversity had a strong positive 310 influence on the primed soil C (Fig. 5). For instance, salinity properties of EC had a directly negative influence on the bacterial diversity but positive influence on the 311 312 primed soil C. Meanwhile, pH were negatively correlated with bacterial diversity and 313 positively correlated with substrate derived C.

314

315 **4. Discussion**

316 4.1. Soil priming effects along salty gradients

317 Understanding soil C dynamics along salinity gradients is crucial to predict C 318 sequestration in salty soils. In the early stage of the incubation, we observed that the 319 cumulative substrate derived CO_2 in the soils with lower salinity was significantly 320 higher than soils with higher salinity (Fig. 1), which can be possibly explained by that 321 high salinity inhibited microbial activity. Many studies have reported the influence of 322 soil salinity on organic matter decomposition, mostly, the decomposition of organic 323 matter are decreased by salinity (Wichern et al., 2006; Ghollarata et al., 2007; Tripathi 324 et al., 2007; Setia et al., 2012). Yet, the response of microbial community to the 325 increasing levels of salinity and consequent effects on soil priming effects remains 326 largely unknown.

Here, we found soil priming effects was gradually changed from negative to positive priming effect (Fig. 1). The early pattern of the dynamics of the priming effect in this study was similar to other studies showing preferential utilization of labile C substance. The first phase of negative priming effects was likely to be caused by microbial assimilation of substrate. The soil microbes turned to use the new added substrate and thus used less of the original SOC. This was attributed to "preferential substrate utilization" (Perelo et al., 2005).

334 Soil microbial biomass-related growth predominating in the first phase were most 335 likely to utilize SOC, leading to a positive priming effects after substrate was largely 336 vanished. The magnitude of priming effects depends on soil microbial biomass size 337 (Schneckenberger et al., 2008). It was found that the amount of added easily available 338 organic C is beyond 50% of microbial biomass C (Blagodatskaya et al., 2008). Namely, 339 the second phase of positive PEs probably was due to increased biomass size and 340 enhanced demand on SOC. Secondly, C that was assimilated into microbial biomass in 341 the first stage may also be mineralized in the second stage due to the turnover of 342 microbial biomass (Shahbaz et al., 2017; Perelo et al., 2005).

343

344 4.2. Microbial community along salt gradients

345 Previous studies concerning the impact of salinity on soil microbial community346 used different soils with a range of salt levels. In the present study we investigated the

347 influence of soil salinity on microbial communities in soils from the closed area covering a range of salt content. Similarly, Rousk et al. (2011) also used agricultural 348 349 soils from the same area representing a range of soil salinity. Here, we found microbial 350 diversity (alpha diversity) decreased with increasing salinity (Fig. 3). The negative 351 impact on microbial diversity can be explained by that the accumulation of large 352 amounts of salt in the soil raised the extracellular osmotic concentration (Rath et al., 353 2015; Oren, 2011). The high osmotic pressures made it difficult for many 354 microorganisms to adapt to and thus reduce their biological activity. The changes of 355 soil microbial community structure were also explained by salinity (Herlemann et al., 356 2011; Campbell et al., 2013). We found that *Bacteroidetes*, *Firmicutes*, *Acidobacteria* 357 and Deinococcus-Thermus were dominant in these soils (Fig. 4). These results are 358 supported by previous findings that *Firmicutes* possess the high salinity resistance. 359 Other studies also found that *Bacteroidetes* is dominant taxa in alkaline saline soil 360 because of its resistant to salt (Valenzuela-Encinas et al., 2009; Keshri et al., 2013). 361 Other study shows that the dominant phyla are Bacteroidetes and followed by Proteobacteria in the haloalkaline soil (Keshri et al., 2013). These results are consistent 362 363 with the esuarine or marine environments, despite some studies suggest that soil salinity 364 is not found to be a decisive factor for bacterial community and their growth (Rousk et 365 al., 2011).

366 The difference of microbial community structure is affected by many soil variables, 367 and pH and EC were the most important ones (Fig. 3; Table 2). Our results showed that 368 the value of soil pH and EC would significantly affect the microbial community 369 structure and the combined contribution rate of these two variables to microbial 370 community was 43% (Table 2). At high levels of salt and alkaline arid condition, soil 371 pH has been also shown to have a very powerful influence on the soil bacterial 372 community structures (Bååth et al., 2003; Fierer et al., 2006; Rousk et al., 2010). 373 Meanwhile, it is consequently unlikely that soil pH differences between the studied 374 soils obscured the influence of salt (Rousk et al., 2011). Salinity has been identified as

one of the most potent environmental factors that determine assembly of microbiome.
Salinity has been regarded to play the vital role in shaping microbial community in
different ecosystem. This, despite the clear evidence from aquatic microbial ecology
(Lozupone et al., 2007b), show a potential for salt to affect soil microbial communities
apart from that of pH (Rath et al., 2015).

380

381 4.3. The core microbial taxa regulating C decomposition along salinity gradient

382 The correlation of microbial taxa and SOC decomposition (priming) were found 383 according to the results of O2PLS and SEM (Table 3; Fig. 5). Here we showed that 384 Streptomyces (Actinobacteria), Glycomyces (branch of Actinobacteria), Agromyces 385 (branch of Actinobacteria), and Sphingomonas (branch of Proteobacteria) at the genus 386 level were significantly correlated with the C process particularly primed soil-drived C. 387 Most of these functional taxa belonged to Actinobacteria and Proteobacteria. In a 388 recent study, Ren et al. (2018) found that Actinobacteria had negative impact on SOC 389 mineralization across land-use change (Fierer et al., 2007; Goldfarb et al., 2011) and 390 Proteobacteria drove the positive soil respiration (He et al., 2012; Stevenson et al., 391 2004), indicating the balance of soil C dynamics were largely regulated by these two 392 phyla. We found similar result that Streptomyces (branch of Actinobacteria) had a 393 negative correlation with primed soil CO₂. Actinobacteria are able to grow 394 preferentially on the C-rich refractory materials and relatively easily decompose the 395 cellulose, lignocellulose (Khodadad et al., 2011), indicating these microorganisms 396 preferentially use the C source that is used partially by others.

Although some studies suggest soil salinity may not be a vital factor for C decomposers (Rousk et al., 2011), the composition of microbial community are considered to play a decisive role in determining C dynamic processes in response to salt stress (Ramsey et al., 2005; Schimel et al., 2007; Nottingham et al., 2009). Here, SEM analysis showed that soil pH and EC in salted soils reduced microbial diversity and thus limited the utilization of SOC by microbial community, It was reported that 403 high pH and salinity are the major determinants of soil microbial activity and404 community structure (Kamble et al., 2014).

405

406 **5. Conclusion**

407 Cotton meal is a kind of organic material with high nitrogen content, adding cotton 408 meal in salinised soil can stimulate and promote the release of soil nutrients. The 409 microorganisms mainly use the organic matter in the cotton meal in the pre-culture 410 period, so the soil carbon excitation is negative excitation, Soil priming effect turned 411 from negative to positive at the later stage of incubation (day 28), because 412 microorganisms turned to decompose SOC from the labile substrate. With the increase 413 of salinity, the diversity of microbial community decreased. soil microbial community 414 was mainly controled by soil pH and EC. By O2PLS, we found Actinobacteria and 415 Proteobacteria (Luteimonas, Hoeflea and Stenotrophomonas) dominant in these soils 416 were the core microbial taxa that affecting the process of organic C mineralization, particularly soil primed CO₂. 417

418

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422

423 Data availability

424 The datasets used and analysed during the current study available from the 425 corresponding author on reasonable request.

426

427 Author contributions

428 K.W. conceptualized and conducted the experiment. H.Z. and D.C. conducted the 429 data analysis and wrote the manuscript, conducted the indoor experiment. C.M. and

Z.Z. assisted in conducting the experiment. All authors reviewed the manuscript.Allauthors contributed to the manuscript and approved the submitted version.

432

433 **Competing interests**

- 434 The authors declare no competing interests.
- 435

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 Table 1. Soil samples and Cottonseed meal properties

	Salinity 1	Salinity 2	Salinity 3	Salinity 4	Salinity 5	Cottonseed meal
Total C (%)	3.38b	3.18c	3.16c	3.57a	3.35b	42.98
Total N (%)	0.18d	0.19d	0.20c	0.22b	0.26a	5.84
C/N ratio	18.32a	16.56b	15.71c	16.54b	12.94d	7.38
δ ¹³ C value (‰)	-14.21a	-14.79c	-14.60b	-14.55b	-16.01d	-23.47
pH (H ₂ O)	8.85a	8.45c	8.58b	8.59b	8.55b	7.63
EC (dS m ⁻¹)	1.06e	1.96c	1.28d	2.64b	7.75a	2.56
Salinity (%)	0.25e	0.58d	0.75c	1.00b	2.64a	ND

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Table 2. Mantel test and Distance-based multivariate analysis relevance and671 contribution rate between soil properties and bacterial community compositions.

	pН	EC	NO3 ⁻ -N	NH4 ⁺⁻ N	MBC	TN	TC
Correlation	0.74**	0.56**	0.36**	0.68**	0.31**	0.11	0.27
Contribution	0.31**	0.12**	0.05	0.04	0.16	0.03	0.07**

672 Note:* P < 0.05, ** P < 0.01

675	Table 3. The variable influence projection (VIP) value and Spearman's correlation
676	between the relative abundances of genera and C dynamic.

		Cottonseed meal	Primed soil	Basal soil
Phylum-Genus			$CO_{1} C(u = 1)$	
		CO_2 - $C(\mu g g^{-1})$	CO_2 - $C(\mu g g^{-1})$	CO_2 - $C(\mu g g^{-1})$
Actinobacteria-Actinomarinales	1.36		0.63**	
Proteobacteria-Luteimonas	1.31		0.80**	
Actinobacteria-Nocardioides	1.30		0.54*	
Proteobacteria-Hoeflea	1.29		0.73**	
Actinobacteria-Streptomyces	1.27		-0.84**	
Actinobacteria-Glycomyces	1.26	0.63**		
Actinobacteria-Marmoricola	1.26	-0.52		
Proteobacteria-Nitrosospira	1.23		0.59	
Actinobacteria-Intrasporangium	1.22		0.60*	
Actinobacteria-Agromyces	1.19			0.58*
Proteobacteria-Sphingomonas	1.18			0.65**
Actinobacteria-Myceligenerans	1.16			
Chloroflexi-Nitrolancea	1.15		0.65**	
Actinobacteria-Pseudarthrobacter	1.06		0.62**	
Proteobacteria-Stenotrophomonas	1.00	-0.50	0.72**	



Fig. 1. Partitioning of CO₂ evolution after addition of cottonseed meal in different five salinity soils. Cumulative CO₂ evolved from salinity soil of 0.25 % (a) , 0.58 % (b) , 0.75 % (c) ,1.00% (d) and 2.64% (e) . Error bars represent standard errors of the means (n = 3).

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Fig. 2. Correlation between primed soil mineralisation and cottonseed meal
mineralisation following different five salinity soils during 90 days incubation



Fig. 3. Microbial community alpha diversity (Chao1) observed_species and beta diversity. Within each panel, boxplot data refer to maximum date (top line), 99%(the second line), mean (the third line), 1% (the fourth line) and minimum date (bottom line) of the different treatments, with statistical significance (P < 0.05).



Fig. 4. The top 10 of phylums and genes in bacterial community in soils with a gradient

of salinity

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 $\chi^2\,{=}\,0.85,\,P\,{=}\,0.65,\,GFI\,{=}\,0.98,\,RMSEA\,{<}\,0.001$

Fig. 5. Path analysis detecting the underlying causal relationships between soil salinity physicochemical factors and microbial community composition of carbon dynamics in the soilt system. Red lines indicate positive relationships, while blue lines indicate negative relationships. The width of arrows indicates the strength of significant standardized path coefficients (P < 0.05). Paths with non-significant coefficients are presented as gray lines. ***P < 0.001; **P < 0.01; *P < 0.05