1	Soil bacterial community triggered by organic matter inputs
2	associates with a high-yielding pear production
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23	Abstract The roles of microorganisms in enhancing crop production have been demonstrated for a range
24	of cropping systems. Most studies to date, however, have been confined to a limited number of locations,
25	making it difficult to identify general soil biotic and abiotic characteristics underpinning the yield-
26	promotion across various locations. This knowledge gap limits our capacity to harness soil microbiome
27	to improve crop production. Here we used high-throughput amplicon sequencing to investigate the
28	common features of bacterial community composition, ecological networks and physicochemical
29	properties in six yield-invigorating and adjacent yield-debilitating orchards. We found that yield-

30 invigorating soils exhibited higher contents of organic matter than yield-debilitating soils and harboured 31 unique bacterial communities. Greater alpha diversity and higher relative abundances of Planctomycetota 32 and Chloroflexota were observed in yield-debilitating soils. Co-occurrence network analysis revealed 33 that yield-invigorating soils displayed a greater number of functionally interrelated modules (meta-34 modules) and a higher proportion of negative links to positive links. Chloroflexota was recognized as a 35 keystone taxon in manipulating the interaction of bacterial communities in yield-invigorating soils. 36 Structural equation modelling showed that soil organic matter, beta diversity of bacterial community, and 37 network connector (Chloroflexota) were identified as potential key factors in explaining the high-yield 38 pear production. Altogether, we provide evidence that yield-invigorating soils across a range of locations appear to share common features, including accumulation of soil organic matter, higher microbial 39 40 diversity, enrichment of key taxa like Chloroflexota, and maintaining a competitive network. These 41 findings have implications for science-based guidance for sustainable food production.

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Keywords: Soil organic matter, Microbial diversity, Random forest prediction, Co-occurrence network,
Keystone taxa

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46 1 Introduction

47 Soils are essential to human wellbeing due to their great contributions to the production of food, fiber, 48 feed, and medicine (Raaijmakers and Mazzola, 2016). Soil organisms play critical roles in maintaining 49 these ecosystem services, such as driving nutrient cycling, maintaining soil fertility, improving plant 50 productivity and suppressing plant diseases (Bender et al., 2016; Barrios, 2007). Microorganisms 51 participate in nearly all soil biological processes, and the microbial abundance, community composition 52 and activity primarily determine the sustainable productivity of agricultural lands (Philippot et al., 2013). 53 Fungi participate in decomposition of organic matter and deliver nutrients for plant growth (Frac et al., 54 2018), however, considering that bacteria are the most diverse and abundant group of microorganisms in 55 soil, bacterial communities and their functions can be pivotal indicators for crop production in 56 agroecosystems (van der Heijden et al., 2008).

57 In general, an increase in microbial diversity is linked to a high-yielding crop production mainly 58 through improving the host resilience to physical or chemical disturbances, modifying plant competition, 59 and facilitating plant access to nutrients (Chaer et al., 2009; Kennedy and Smith, 1995). Since individual 60 organisms do not live in isolation but rather form a complex system of inter-species interactions in soil, 61 interactions among community members were found to be related to crop production in the potato 62 monoculture system (Lu et al., 2013). Enrichments of key functional microbes in soil were deemed to 63 serve specific soil system functions, such as suppressing soil-borne pathogens and maintaining 64 sustainable crop production (Banerjee et al., 2018). However, the relative contributions of microbial 65 diversity, interactions among community members, or enrichment of key taxa to crop production remain 66 largely unknown. Therefore, it is highly desirable to identify pivotal indicators of bacterial community 67 composition in response to high-yielding crop production.

68 Changes in composition of soil bacterial communities across space are often strongly correlated 69 with soil pH (Fierer and Jackson, 2006). Soil pH has been recognized as a key driver in determining the 70 assembly of bacterial community in arable soils by field or microcosm experiments (Rousk et al., 2010). 71 However, recent studies have demonstrated that compositions of soil bacterial communities were driven 72 by a myriad of soil abiotic traits, such as organic matter contents, forms and contents of soil nutrient 73 (Tian et al., 2018; Wang et al., 2018). For example, soil bacterial community composition, which 74 determines the ability of soil to suppress soil-borne pathogens, was found to be strongly correlated with 75 soil organic matter (Shen et al., 2018). An imbalanced ratio of soil nutrients, *i.e.*, ratio of nitrogen to 76 phosphorus or potassium could be a driving force altering the bacterial community composition in long-77 term fertilized soils (Eo and Park, 2016). Key soil chemical properties identified in controlling the 78 distribution and abundance of bacterial community is largely depending on the geographical distributions 79 of soils. As a consequence, a better understanding of the relationship between soil edaphic properties and 80 bacterial community composition is critical to develop targeted manipulation options to increase soil 81 service provisions.

82 'Sucui No. 1' pear, is an early-maturing variety bred by the Jiangsu Academy of Agricultural 83 Sciences, China, and has been popularly cultivated in Eastern and Central China, due to the advantages 84 including easy to produce, adaptable to the environment, and has good quality and high economic 85 benefits (Lin et al., 2013). With the increasing demand in China, sustainable production of high-quality 86 pear is becoming increasingly important. Manipulation of soil microbiomes has shown to be an effective 87 way to increase soil productivity (Chaparro et al., 2012). Considering that large-scale surveys could 88 exhibit the diversity of soil microbial communities exceeds what is found in host-associated communities 89 (Toju et al., 2018), it is necessary to explore the general microbial characteristics of multiple yield90 invigorating soils and identify key environmental drivers in assembling bacterial community.

91 In this study, compared to local average yield, orchard showing higher pear yield production was 92 recognized as yield-invigorating (YI) orchard while orchard displaying lower pear yield production was 93 regard as yield-debilitating orchard (YD). After field surveys accomplished in 2019, six yield-94 invigorating and adjacent yield-debilitating pear orchards in total were selected for further analysis of 95 soil chemical properties and microbiome. We hypothesized that high input of organic fertilizer could 96 improve soil structure and modify chemical properties, which leading to YI soils harbor unique bacterial 97 communities that maintains the high-yielding pear production. To address this, soil bacterial communities 98 and edaphic properties of the study sites were compared to (1) decipher the differences of taxonomic 99 diversity, and composition of the bacterial community, and (2) determine the contributions of 100 environmental variables to the changes in the structure of bacterial communities.

101 2 Methods

102 2.1 Study sites and experimental design

103 From July - August 2019, a field production survey of orchards cultivated with 'Suci No. 1' pear was 104 performed after pear fruits harvest to compare the differences of soil nutrients and microbiota between 105 yield-invigorating with yield-debilitating orchards. The locations, planting density, cropping years, soil 106 type and total yield were recorded. To minimize the effects of microclimate at each site, only pair-located 107 pear orchards with invigorating and debilitating yield and at similar growth stage were selected for this 108 research. In total, six pair-located yield-invigorating and -debilitating pear orchards distributed in four 109 cities of Jiangsu province, China, were selected in the main pear production areas (Fig. 1, Table S1). The 110 yield per tree was obtained by dividing the total yield per hectare by plant density.

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Fig. 1 here

112 Paired yield-invigorating and yield-debilitating orchards from Fengxian (FX), Suining (SN) and 113 Tongshan (TS) were maintained in the Xuzhou city under the warm temperate sub-humid monsoon 114 climate. This site has a mean annual temperature (MAT) of 14.5 °C and mean annual precipitation (MAP) 115 of 847 mm. Orchards from Taixing (TX) were located in the Taizhou city under the humid southern 116 subtropical climate with a MAT of 15.3 °C and MAP of 1055 mm. Orchards from Gaochun (GC) were 117 located in the Nanjing city under the humid subtropical monsoon climate with a MAT of 15.4 °C and 118 MAP of 1106 mm. Orchards from Zhangjiagang (ZJ) were located in the Suzhou city under the humid 119 subtropical monsoon climate with a MAT of 15.7 °C and MAP of 1094 mm. For paired yield-invigorating and-debilitating orchards, the irrigation and pesticide management practices were similar according to farm records. However, yield-invigorating orchard was usually amended with more organic fertilizer under integrated nutrients management whereas the co-located yield-debilitating orchard received more chemical fertilizer under intensive management. Detailed information about fertilization regimes for each orchard is shown in Table S2.

125 2.2 Soil sample collection and chemical properties determination

126 Along with the field survey, soil sampling campaigns were performed from July - August 2019 after pear 127 fruits harvest. For each yield-invigorating or -debilitating orchard, four subplots with three pear trees in 128 each subplot were randomly selected for soil sampling. Subsequently three soil cores (0-20 cm) under 129 the trunk base for each tree were collected using a 25 mm soil auger. In total, nine soil cores for each 130 subplot were pooled as a composite sample and finally four composite soil samples for each orchard 131 were collected and promptly transported on ice to the laboratory. After sifting through a 2 mm sieve and 132 thoroughly mixing, one portion of each soil sample was air-dried for chemical property analyses while 133 the remainder was stored at -70 °C for DNA extraction.

134 Soil chemical properties including soil pH, content of organic matter (OM), total nitrogen (TN), 135 available phosphorus (AP), available potassium (AK), alkali-hydrolyzale nitrogen (N), exchangeable 136 calcium (Ca), effective magnesium (Mg), effective iron (Fe), effective manganese (Mn), effective copper 137 (Cu) and effective zinc (Zn), were measured according to methods described by Shen et al. (2018) and 138 Huang et al. (2019). Briefly, soil pH was determined using a glass electrode meter in a suspension with 139 a 1:5 soil/water ratio (w/v). Soil OM was determined using the potassium dichromate external heating method. TN was determined using a dry combustion method on an Element Analyzer (Vario EL, 140 141 Germany). AP and AK were determined using the molybdenum blue method after soil was extracted with 142 sodium bicarbonate and flame photometry after soil was extracted with ammonium acetate, respectively. 143 Soil alkaline hydrolysable nitrogen (N) was measured by the alkaline hydrolysable diffusion method. 144 Contents of soil Ca, Mg, Fe, Mn, Cu and Zn were determined by the atomic absorption spectroscopy 145 method using ICE 3300 AAS Atomic Absorption Spectrometer (ThermoScientific, USA) after acid 146 hydrolysis.

147 2.3 Soil DNA extraction and bacterial abundance quantification

148 Genomic DNA from 0.25 g soil for each sample was extracted by using the DNeasy[®] PowerSoil[®] Kit

149 (QIAGEN GmbH, Germany) according to the manufacturer's instructions. The abundances of soil

- bacteria were determined with Eub338F/Eub518R primer using a 7500 Real Time PCR System (Applied
- 151 Biosystems, USA). Standard curves were generated by using 10-fold serial dilutions of a plasmid
- 152 containing a full-length copy of the 16S rRNA gene from *Escherichia coli*. Quantitative PCR analysis
- 153 was performed in 96-well plates with a 20-µl mixture for each reaction using SYBR[®]Premix Ex TaqTM
- 154 (TaKaRa, Japan). Thermal cycling was conducted according to a standard procedure with three replicates,
- and the results were expressed as log copy numbers g^{-1} dry soil.
- 156 2.4 Sequencing library construction and sequencing
- 157 The gene-specific primers 515F/806R with 12 bp barcode were used to amplify the V4 region of bacterial
- 158 16S rRNA gene on the BioRad S1000 (Bio-Rad Laboratory, CA) according to the protocols described
- by Caporaso et al. (2011). All constructed libraries were sequenced using the Illumina NovaSeq 6000 at
- 160 the Guangdong Magigene Biotechnology Co., Ltd. (Guangzhou, China).
- 161 2.5 Sequence processing

162 Quality filtering of the paired-end raw reads was performed to obtain the high-quality clean reads 163 according to the Trimmomatic V0.33 (Bolger et al., 2014) quality control process. Sequences were 164 assigned to each sample based on their unique barcode, after which the barcodes and primers were 165 removed. Paired-end clean reads were merged using FLASH V1.2.11 (Magoč and Salzberg, 2011). Raw 166 tags were processed to generate the final ASV (Amplicon Sequence Variant) table file at 97% pairwise 167 identity according to the QIIME2 pipeline (Bolyen et al., 2019). The nonbacterial and mitochondrial 168 ASVs and extremely low frequency ASVs (relative abundance < 0.01%) were removed. A representative 169 sequence for each ASV was selected and classified using the RDP classifier against the RDP Bacterial 170 16S database (Wang et al., 2007).

171 2.6 Statistical analyses

172 Statistical analyses were performed using the software SPSS 20.0 (SPSS Technologies, Armonk, NY, 173 USA) and R (http://www.R-project.org/). Non-normal data were square-root or log transformed when 174 necessary. The significance of soil properties or microbial taxa in yield-invigorating or-debilitating 175 orchards was determined based on paired Wilcoxon rank sum test, and adjusted P values (< 0.05) were 176 obtained by the FDR method. Mantel tests were used to identify the correlations between microbial 177 community composition and pear yield, and soil chemical properties using the 'vegan' package (Oksanen et al., 2013) in R. The linear regression analyses relating yield to selected microbial taxa or soil chemical 178 179 properties were conducted using the 'basicTrendline' package (Mei et al., 2018) in R.

180 Principal Coordinate Analysis (PCoA) based on the Bray-Curtis distance was performed in 181 MOTHUR V1.38.1 (Schloss et al., 2009) and visualized by the 'ggplot2' package (Wickham and Chang, 182 2015) in R to explore the differences in microbial community composition. Permutational multivariate 183 analysis of variance (PERMANOVA) was performed to evaluate the significant differences of microbial 184 community composition according to sample locations and orchard yield using the 'vegan' package in R. 185 Microbial alpha diversity indexes (Chao, Shannon) were calculated based on randomly resampled ASV 186 abundance matrices at the same depth (23,800 sequences) in MOTHUR. A Venn diagram was generated 187 based on the final ASVs to compare microbial community composition between yield-invigorating and 188 -debilitating orchard soils. The affiliations of unique and shared ASVs in yield-invigorating and -189 debilitating soils were compared to evaluate the differences of the bacterial community composition and 190 plotted using the 'pheatmap' package (https://cran.r-project.org/web/packages/pheatmap) in R. Fold 191 changes (log2 transformed) of shared ASVs across yield-invigorating and -debilitating soils were 192 calculated. The ASVs with fold change ratios > 2 and unique ASVs in yield-invigorating soils were 193 recognized as potential responders to yield promotion. In addition, to better understand the bacterial 194 community composition, relative abundances at the genus level were compared by STAMP software 195 v2.1.3 (Parks et al., 2014).

196 The phylogenetic molecular ecological networks (pMEN) were constructed using the random 197 matrix theory-based approach to explore the organization of bacterial communities in yield-invigorating 198 (YI) or yield-debilitating (YD) soil samples. Potential ecological interactions among bacteria were 199 determined by modeling the microbial community using Molecular Ecological Network Analysis 200 (http://ieg2.ou.edu/MENA) based on pear yield (Deng et al., 2012). Given the large number of rare taxa 201 that are specific to certain locations, ASVs that occurred in less than half of soil samples and lower than 202 0.01% were filtered, which resulting in 591 and 485 ASVs for YI and YD samples respectively, before 203 networks constructed. The microbial network was constructed using random matrix theory-based at 204 0.94 similarity threshold and visualized using Cytoscape 2.8.3 software (Smoot et al., 2011). Module 205 clustering and composition in yield-invigorating and -debilitating networks were compared and plotted 206 in R using the 'pheatmap' and 'ggplot2' packages. And the threshold values of Zi and Pi was 2.5 and 207 0.62, respectively for topology analysis of the network. Redundancy analysis (RDA) was performed in 208 the R 'vegan' package to examine the relationship among frequencies of ASVs, samples and soil 209 variables, which were selected using 'stepAIC' in R. Variance partitioning analysis (VPA) was used to

210 determine the contributions of soil properties, sample location, and yield, as well as interactions among 211 the variation in a microbial community with hellinger-transformed data. The predictors of selected soil 212 properties for explaining the pear yield were identified by random forest regression analysis (Boulesteix, 213 et al., 2012). The significance of each predictor in the response variables was assessed with the 214 'rfPermute' package (Liaw and Wiener, 2002) with 1000 permutations based on 1000 trees. Structural 215 equation modelling was applied to evaluate relative contributions of soil chemical properties and 216 bacterial community to pear yield (Grace et al., 2006; Schermelleh-Engel et al., 2003). The conceptual 217 SEM fitness was examined on the basis of a non-significant chi-square test (P > 0.05) and the goodness-218 of-fit index (GFI). Model was fitted using the 'lavaan' package in R (Rosseel, 2012).

219 3 Results

220 3.1 Overview of sequencing data

221 In total, 1,622,858 16S rRNA sequences were retained after quality control and a total of 9,394 ASVs

222 were obtained for the 16S rRNA gene sequences based on 97% similarity. Among the total 16S rRNA

223 gene sequences, 159 ASVs with 74,372 sequences were classified as Archaea while 9,235 ASVs with

224 1,548,486 sequences were identified as Bacteria. Among Bacteria, Acidobacteriota, Pseudomonadota,

225 Chloroflexota, Planctomycetota and Actinomycetota were the most abundant phyla (Fig. S1).

226 3.2 Soil chemical properties

227 Soil chemical properties differed significantly among the locations and orchard yield types (Table S3).

228 However, when taken all sites together, only a higher relative abundance of soil organic matter (OM) on

229 average was observed in yield-invigorating orchards compared to that in yield-debilitating orchards

based on the Wilcoxon test.

231 3.3 Bacterial abundances and community compositions

Yield-invigorating orchards together displayed significantly higher abundances of total bacteria than that in co-located yield-debilitating orchards based on real time PCR result (Fig. 2A). Meanwhile, bacterial community compositions at the ASV level were significantly correlated to pear yield (r = 0.460, p = 0.001) (Fig. 2B).

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Fig. 2 here

PCoA based on Bray-Curtis distance matrices clearly revealed location-based differences in
 bacterial community compositions (Fig. 3A). Six distinct groups representing samples from different
 locations (FX, GC, SN, TS, TX and ZJ) were obviously separated and confirmed by PERMANOVA test

(F = 14.9, P = 0.001). At each location, soil bacterial community composition in yield-invigorating orchards was significantly separated from that in co-located yield-debilitating orchards, which was also confirmed by PERMANOVA test (F = 3.6, P = 0.001). Although only the Shannon diversity in yieldinvigorating orchards from GC and ZJ was significantly higher than that in co-located yield-debilitating orchards (Fig. S2), the mean alpha diversity indices of Chao and Shannon in all yield-invigorating orchards were significantly higher than those in all yield-debilitating orchards based on the paired Wilcoxon test (Fig. 3B).

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Fig. 3 here

248 The Venn diagram showed that 4,540 ASVs occupying over 90% of total sequences were shared 249 between yield-invigorating and -debilitating orchards (Fig. 3C). Among these shared ASVs, the fold 250 changes larger than 2 of ASVs in yield-invigorating compared to yield-debilitating orchards were 251 recognized as potential responders linking to yield improvement. Surprisingly, none of these ASVs 252 potentially linked to yield improvement were shared among six separated collocated orchards (Fig. S3). 253 A total of 2,546 unique ASVs with 53,222 sequences were found in all yield-invigorating orchards while 254 2,308 unique ASVs with 44,389 sequences were observed in all yield-invigorating orchards. Among 255 these unique ASVs, almost 70% of ASVs were shared between yield-invigorating orchards and -256 debilitating orchards. However, no shared unique ASVs were found among six separately located 257 orchards. The affiliation of unique and shared ASVs at the phylum level exhibited that the 258 Pseudomonadota, Planctomycetota, Chloroflexota, Acidobacteriota and Actinomycetota were the top 259 five phyla (Fig. 3D).

At the phylum level, the relative abundances of bacterial dominant phyla varied across the location and orchard yield condition. Pseudomonadota, Acidobacteriota, Actinomycetota, Chloroflexota and Planctomycetota were the top five abundant phyla (Fig. 4). The mean abundance of Chloroflexota and Planctomycetota was significantly higher while Firmicutes was significantly lower in yield-invigorating orchards compared to yield-debilitating orchards based on Wilcoxon test (Fig. S4).

At a finer resolution, 967 genera were observed for all soil samples, among which 299 genera appeared in more than half of soil samples in yield-invigorating or -debilitating orchards. However, only 34 genera displayed significant differences between yield-invigorating or -debilitating orchard soils based on STAMP analysis (Fig. S5). Interestingly, *Ornatilinea, Ktedonobacter, Longilinea*, belonging to Chloroflexota, were significantly enriched in yield-invigorating orchard soils. *Gimesia* in

- 270 Planctomycetota and Arenimonas in Pseudomonadota showed significantly higher relative abundances
- in yield-invigorating orchard soils than in yield-debilitating orchard soils.
- 272

Fig. 4 here

273 3.4 Co-occurrence patterns of bacterial community

274 The yield-invigorating network contained 302 nodes, 448 edges, and 11 larger modules (> 5 nodes), with 275 an average connectivity (avgK) of 2.967, average path distance of 5.494 and clustering coefficient 276 (avgCC) of 0.152, while the values in the yield-debilitating network were 235, 334, 9, 2.843, 6.232 and 277 0.131, respectively (Fig. 5A, Table S4). The module eigengene network analysis revealed a difference in 278 the higher-order organization between the two networks. Notably, the node composition was substantially 279 different between the two networks as the relative abundances of dominant phyla were obviously 280 different among different modules (Fig. 5B and C). A higher proportion of nodes in the module of yield-281 invigorating network was unique. ASVs affiliated to Acidobacteriota, Chloroflexota, Pseudomonadota, 282 Actinomycetota, and Planctomycetota within the unique modules (M9, M10 and M11) were observed in 283 the yield-invigorating versus yield-debilitating network.

284 Analysis using the threshold values of Zi (within-module connectivity) and Pi (among-module 285 connectivity) showed that majority of nodes from both networks were categorized as peripherals that had 286 only a few links and almost always linked to the nodes within their own modules (Fig. 5D). Although 287 only three nodes affiliated to Acidobacteriota were categorized as module hubs in the vield-invigorating 288 network, seven nodes belonging to Acidobacteriota, Actinomycetota and Pseudomonadota were 289 categorized as module hubs in the yield-debilitating network. Interestingly, four nodes including 290 Longilinea species from Chloroflexota in the yield-invigorating network whereas only one node in the 291 yield-debilitating network was categorized as module connectors (Table S5).

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Fig. 5 here

293 3.5 Relationships between soil chemical properties and microbial community composition

294 Soil chemical properties were significantly correlated to the bacterial community compositions (Mantel:

r = 0.803, p = 0.001). Soil chemical properties, location, and orchard explained 44.9% of the observed

- 296 variation, leaving 55.1% of the variation unexplained for bacterial community composition based on
- 297 VPA result (Fig. 6A). Variation in the community composition was largely explained by soil properties
- 298 (42.3%), and was also influenced by locations and orchard yield types.

299 After forward stepwise selection, the module including soil OM, TN, alkaline N, AP and AK,

available calcium (Ca), copper (Cu) and manganese (Mn) explained the majority of the variation in
bacterial community composition (Fig. 6B). As evidenced by the RDA vectors, OM within the module
was identified as the top important soil property that determines the composition of bacterial community.
Random forest analysis showed that contents of soil Mn, OM and Ca were the top parameters for
predicting the orchard yield (Fig. 6C). Furthermore, soil OM was also significantly correlated with
bacterial communities as revealed by Mantel test (Fig. 6D, Table S6).

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Fig. 6 here

307 3.6 Relationships of soil chemical and microbial indicators with orchard yield

308 Soil OM as potentially key soil chemical properties and bacterial alpha diversity, beta diversity and 309 relative abundance of Chloroflexota and Planctomycetota as potentially key microbial indicators 310 associated with pear yield were used to construct a model to explain yield improvement. Final structural 311 equation modelling (path analysis) (Fig. 7 and S6) showed that the strongest driver explaining yield 312 improvement was beta diversity of bacterial community (PCoA) (r = 0.959, P < 0.001), which was 313 positively affected by content of soil OM (r = 0.843, P < 0.001). Alpha diversity (Chao) of bacterial 314 community also determined yield improvement to a large extent (r = 0.542, P = 0.009). However, alpha 315 diversity was not significantly correlated with content of soil OM.

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Fig. 7 here

317 4 Discussion

318 Although pear is among the most important fruits worldwide, soil microbial communities in pear 319 orchards have been largely under-investigated (Huang et al., 2019). The present study attempts to 320 decipher the bacterial community linked to high-yield production of pear. Our results based on Mantel 321 analysis suggested significant correlations among bacterial community, soil chemical properties and pear 322 yield. Microbial characteristics responding to yield promotion have repeatedly been observed on several 323 crops depending on single experimental site (Zhong et al., 2020; Qiao et al., 2019; Shen et al., 2013). It 324 remained unclear, however, whether these distinctions are ubiquitous at a large-scale. By comparing 325 multiple co-located yield-invigorating and -debilitating orchards, we demonstrate that high-yielding pear 326 production soils exhibited high organic matter contents and harbored bacterial communities with high 327 diversity, significantly enriched indigenous microbes and more interactive network, which was triggered 328 by high-inputs of soil organic fertilizer. Here we discussed these main results and potential mechanisms 329 in detail.

Microbial diversity is critical to soil ecosystems in maintaining the integrity, function and long-term sustainability (Kennedy and Smith, 1995). Higher soil biodiversity is considered to be linked to a more stable system and enhance the combination of vital microbial functions and processes (Cardinale et al., 2006; Bell et al., 2005). In line with a previous report that crop yield was correlated to the soil bacterial diversity (Zhao et al., 2014), greater diversity of bacterial community in yield-invigorating soils was observed in the present study. Hence we infer that higher microbial diversity may result in a more productive agroecosystem, contributing to sustainable pear production.

337 In this study, we found that Pseudomonadota, Acidobacteriota, Actinomycetota, Planctomycetota 338 and Chloroflexota were the top abundant phyla. This result roughly agreed with previous studies showing 339 that Pseudomonadota, Acidobacteriota and Actinomycetota are usually dominant bacterial taxa in 340 agricultural soils (Xun et al., 2019; Dai et al., 2018), while Planctomycetota and Chloroflexota exhibit 341 an unexpectedly high relative abundance in rice cropped soil (Edwards et al., 2015) and sandy loam soil 342 (Pathan et al., 2021). The highest relative abundance of Pseudomonadota was probably explained by the 343 fact that Pseudomonadota are considered as copiotrophic bacteria and flourish in soils with large amounts 344 of available nutrients (Fierer et al., 2007).

345 Moreover, a significantly higher abundance of Planctomycetota and Chloroflexota was observed in 346 yield-invigorating orchards, indicating that Planctomycetota and Chloroflexota may be associated with 347 pear yield-improvement. There is no direct evidence showing that Planctomycetota could improve plant 348 growth. However, Planctomycetota has been reported to be involved in many soil biological processes 349 such as ammoxidation, carbohydrate and polysaccharide metabolism (Fuerst, 2017). This implies that 350 Planctomycetota may promote plant production through improving soil fertility. Chloroflexota is a 351 facultative anaerobic phylum including autotrophic, heterotrophic and mixotrophic taxa (Speirs et al., 352 2019). Considering that soil amended with organic fertilizer may enhance the soil water holding capacity, 353 the yield-invigorating soils with more organic material input have a higher soil moisture content, 354 especially after irrigation, probably leading to the enrichment of Chloroflexota in soil.

Network analysis is a systems-level method to explore interactions within an ecosystem that cannot be directly observed through co-occurrence analysis (Fath et al., 2007). Similar to the food web network analyses in macro ecosystems, microorganisms also form complex interactions with other species (Faust and Raes, 2012) and have been widely investigated to explore the linkage of microbial network with soil function, such as nutrient supply (Fan et al., 2021) and disease suppression (Lu et al., 2013). Overall, in

360 line with previous findings (Hu et al., 2020), the topological properties of the constructed networks, 361 including connectivity, average clustering coefficients, average degree distance, and modularity indicate 362 that these networks are scale-free, modular and small world. In short, a scale-free network represents that 363 a network whose connectivity follows a power law, and most of nodes have only a few connections with 364 other nodes. Meanwhile, a small-world network is the network in which most nodes are not neighbors of 365 one another, but most nodes can be reached by a few paths. Modularity is a fundamental characteristic 366 of biological network as a module in the network is a group of nodes that are highly connected within 367 the group, but very few connections outside the group (Deng et al., 2012). Our comparative network 368 analysis indicated that microbial co-occurrence patterns in soils links to different pear production. As a 369 meta-module is usually considered as a group of modules functionally interrelated (Langfelder and 370 Horvath, 2007), a greater number of meta-modules were identified in the network constructed from yield-371 invigorating soils, suggesting that a greater number of network nodes in the yield-invigorating soils were 372 functionally interrelated than those in the yield-debilitating soils. A majority of nodes in the meta-373 modules were not shared between yield-invigorating and -debilitating networks, indicating basal shifts 374 in network architecture during pear production with contrasting yield performance.

375 Furthermore, a higher proportion of negative interactions to positive interactions were identified in 376 the network constructed from yield-invigorating network than the yield-debilitating network. Our results 377 indicated stronger resource competitions in yield-invigorating soils, which means that the soil co-378 occurrence network was more stable to maintain soil ecosystem function (Coyte et al., 2015). In our 379 study, three module connectors and three module hubs were identified as potentially key taxa in the yield-380 invigorating network. Interestingly, among those key species, ASV357 affiliated to Longilinea, 381 belonging to the Chloroflexota, was recognized as a key phylum in improving pear yield. Similarly, 382 Chloroflexota was reported to be key-stone taxa in the constructed network from agricultural soils with 383 40-years fertilization (Fan et al., 2021). Chloroflexota play key roles in connecting network nodes of soil 384 microbiome probably due to that Chloroflexota could participate in degrading plant compounds to create 385 more nutrients via pathways for the degradation of starch, cellulose, and longchain sugars, as it is 386 positively correlated with genes for amino sugars, sugar alcohols and simple carbohydrate metabolic 387 pathways (Hug et al., 2013).

388 In this study, a significantly higher content of soil organic matter was observed in yield-invigorating 389 orchards, demonstrating that soil organic matter could drive the assembly of bacterial community. Consensus is emerging that microbial residues are an important constituent of soil organic matter (Kallenbach et al., 2016), , which participate in almost all soil biological processes (Fierer, 2017). Despite the quality of soil organic matter was not evaluated in this study, the quality of soil organic matter was associated with the diversity of microbial community (Ding et al., 2015), which implies more attentions should be paid to illustrate the relationship between the quality of soil organic matter and microbial community in our future work.

396 Structural equation modelling approach has been widely used to decipher keystone indicators 397 associated with soil function and crop production in agroecosystems (Jiang et al., 2020; Chen et al., 2019). 398 In the present study, we observed that soil organic matter, beta diversity of bacterial community, and 399 network connector were key indicators in supporting high-yield pear production based on the structural 400 equation modelling results. Worth to mention, soil organic matter was not directly linked to the yield in 401 the constructed model, indicating that soil organic matter maintain the high-yielding pear production 402 probably via the indirect ways. Therefore, we proposed that yield-invigorating soils harbour unique 403 bacterial communities that may improve soil biological fertility, which could be driven by soil organic 404 matter and manipulated by keystone species (Chloroflexota) through altering the bacterial interactions.

405 **5** Conclusions

In conclusion, yield-invigorating soils displayed a higher content of organic matter and harboured unique bacterial community with greater diversity than yield-debilitating soils. Further Chloroflexota was significantly enriched and identified as a potential keystone taxon in manipulating the interaction of bacterial community in yield-invigorating soils. These findings indicated that soil organic matter triggered the assembly of soil microbiome, which both participated in maintaining crop production . Such knowledge is a first step toward harnessing soil microbiome in support of sustainable agroecosystems.

412

413 **Data availability.** Raw amplicon sequencing data for each sample used in this study was deposited at 414 the National Center for Biotechnology Information (NCBI) in the FASTQ format and is available under 415 the accession number PRJNA749397. Other data that support the findings of this study are available on 416 request from the corresponding author (Xiaomei Ye).

417

418 Authors' contributions

419 L. Wang: performed all experiments; L. Wang, X. Ye, and Z. Shen: designed the study, and wrote the

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439	References
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437	Supplementary figures and tables to this article can be found in the supplemental material.
436	Appendix A. Supplementary data
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434	42090065).
433	Innovation Fund (CX(19)3094) and National Natural Science Foundation of China (31801842 and
432	Financial support. This research was supported by Jiangsu Agricultural Science and Technology
431	
430	all pear orchards owners for providing access to the soil sampling.
429	Acknowledgements. We sincerely thank all those who have assisted me with any part of this paper and
428	
427	claims in published maps and institutional affiliations.
426	Disclaimer. Publisher's note: Copernicus Publications remains neutral with regard to jurisdictional
425	
424	Competing interests. The authors declare that they have no conflict of interest.
423	
422	The authors read and approved the final manuscript.
421	Lin, and D. Chen: participated in the design of the study, provided comments and edited the manuscript.
420	majority of the manuscript; L. Wang and Z. Shen and C. Tao: analyzed the data; H. Hu, J. Du, Y. Xi, J.

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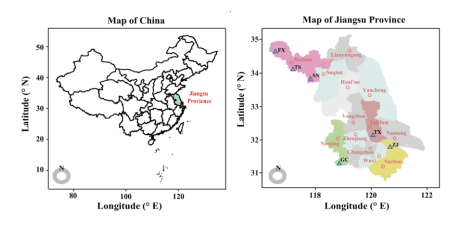
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597 Figure legends

598 Fig. 1 Distribution of studied field sites. Map showing the sites of six pair-located orchards sampled in

599 this study.

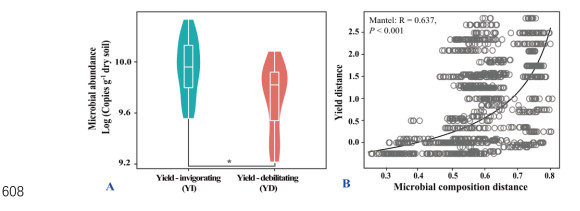
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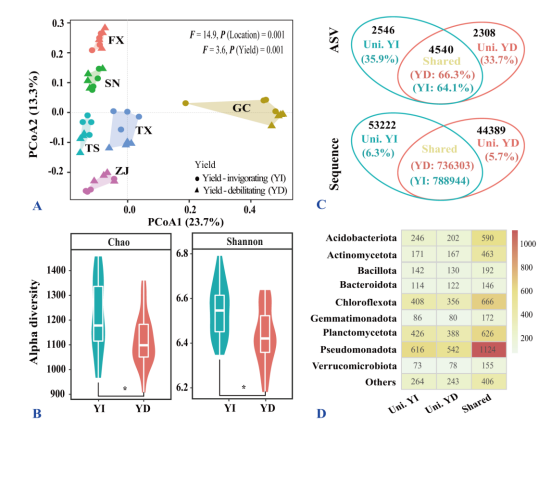
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603 Fig. 2 Quantitation of the abundance of bacteria population, and linkage of microbial composition

to pear yield. (A) Violin plot showing the abundance of total bacteria for all selected orchards. * indicates a significant difference between yield-invigorating (YI) and yield-debilitating (YD) orchards based on Wilcoxon tests (p < 0.05). (B) Correlation plot showing the relationship of microbial composition and yield based on braycurtis distances calculated by Mantel test.



610 Fig. 3 Overview of bacterial composition and alpha diversity. (A) Principal Coordinates Analysis 611 (PCoA) plot displaying the bacterial community composition calculated based on braycurtis distances. 612 (B) Violin plot showing the alpha diversity indices (Chao and Shannon) for all selected orchards. * 613 indicates a significant difference between yield-invigorating (YI) and yield-debilitating (YD) orchards based on Wilcoxon tests (p < 0.05). (C) Venn plot depicting the unique and shared bacterial ASVs 614 615 between yield-invigorating (YI) and yield-debilitating (YD) orchards at ASV and sequence insights. 616 Uni. YI and Uni. YD represent unique ASVs or sequences in the YI or YD soils while Shared represent 617 shared ASVs or sequences between the YI and YD soils. (D) Heatmap displaying the composition of unique and shared ASVs at phylum level in YI and YD soils. Numbers in the cell represent the number 618



619 of ASVs affiliated to that phylum.

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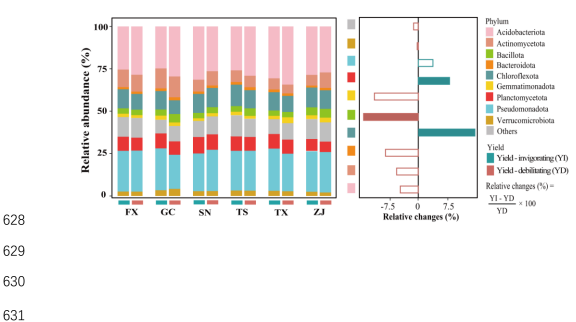
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624 Fig. 4 Key taxonomic groups in distinguishing yield-invigorating (YI) and yield-debilitating (YD)

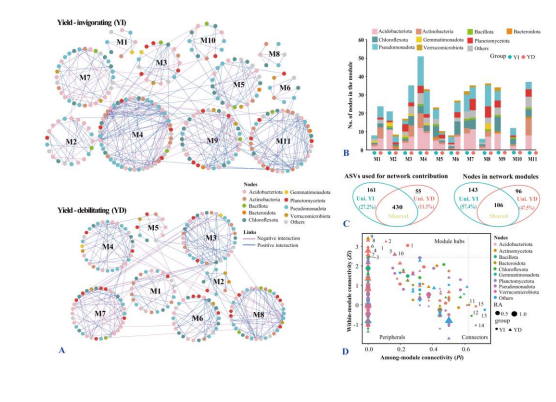
625 orchards. Stacked bar chart (left panel) showing dominant phyla affiliation in YI and YD soils for six

626 pair-located sites while horizontal histogram (right panel) depicting relative changes of dominant phyla

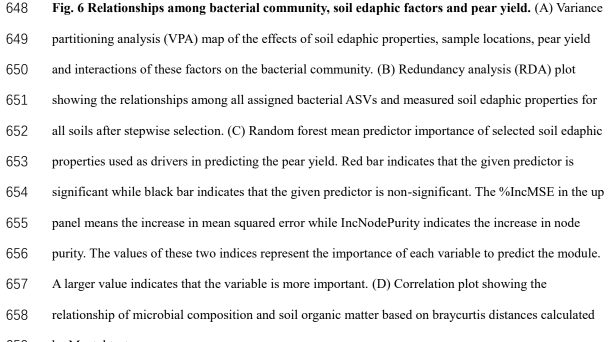


627 in YI soils compared to those in YD soils.

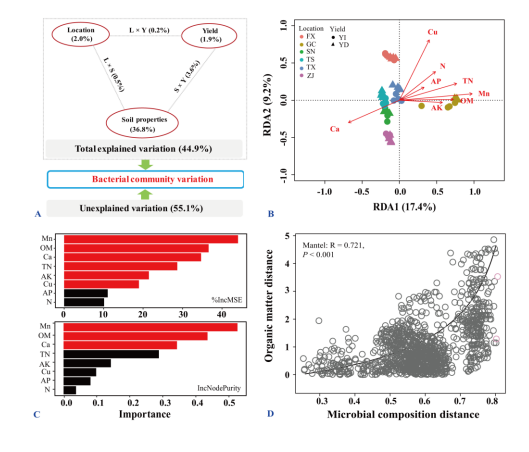
632 Fig. 5 Co-occurrence networks of bacterial community and identified keystone taxa in 633 distinguishing yield-invigorating (YI) and yield-debilitating (YD) orchards. (A) An overview of 634 microbial phylogenetic molecular ecological networks constructed from YI and YD soils. Line with blue 635 color indicates positive correlations whereas lines with red color signifies negative correlations in each 636 network. Modules containing larger than five nodes in the networks are labeled with corresponding letter 637 followed by a number. Circular node colors indicate different bacterial phyla. (B) Stacked figure showing 638 the relative abundance of nodes in each module within each network at the phylum level. (C) Venn plot 639 depicting the unique and shared bacterial ASVs between two networks construed from YI and YD soils. 640 Left panel is plotted based on the original nodes used in building network while right panel is plotted 641 based on the nodes from modules. Uni. YI and Uni. YD represent unique ASVs in the YI or YD networks 642 while Shared represent shared ASVs between the YI and YD networks. (D) Zi-Pi plot showing the 643 distribution of nodes based on their topological roles. The threshold values of Zi and Pi for categorizing 644 OTUs were 2.5 and 0.62 respectively. Node colors indicate different bacterial phyla and node size 645 represent the relative abundance in each network.



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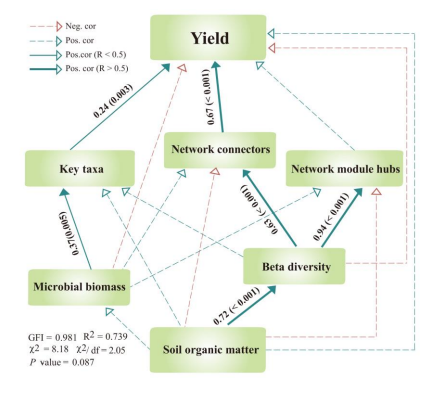
by Mantel test.



663 Fig. 7 Structural equation modeling (SEM) describing the biotic and abiotic factors in affecting

664 the crop production. Structural equation model was built incorporating soil organic matter, microbial

- biomass, beta diversity (PCoA), key taxa, network hubs including module hubs and network
- 666 connectors, and yield. The path analysis numbers adjacent to arrows indicate the relationship's effect
- size and the associated bootstrap *P*-value. Cyan and red arrows indicate positive and negative
- relationships, respectively. Paths with non-significant coefficients are presented as gray lines.



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