Research Article

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Effects of organic materials on soil bacterial community structure in long-term continuous cropping of tomato in greenhouse

https://doi.org/10.1515/biol-2022-0048
received August 15, 2021; accepted January 10, 2022

Abstract: Long-term fertilization will affect the above-ground vegetation, but we have little understanding of soil bacterial community structure and diversity so far. This study aims to study the effect of organic fertilization on the soil bacterial community structure and diversity of protected long-term continuous tomato cropping by using high-throughput sequencing technology. Results show that (1) fertilization application (chemical fertilizer [CF] and vermicompost [VM]) significantly changed the soil physico-chemistry properties, such as soil pH decreased compared with control treatment and increased the soil organic carbon (SOC), total nitrogen (TN), total phosphorus (TP), and total potassium (TK) contents; (2) VM increased the Shannon index of soil bacteria but decreased the soil Chao1 index; and (3) soil Proteobacteria and Actinomycetes were dominant taxa and the relative abundance of Actinobacteria increased by 36.40% and 25.80–29.35%, with CF and VM, respectively, compared with the control. Pearson correlation analysis showed that soil pH, SOC, TN, TP, and TK were the main environmental factors that affected the diversity and richness of soil bacterial communities. Redundancy analysis (RDA) showed that the SOC and TN play important roles in the composition of soil bacterial communities. In summary, the effect of VM on the soil bacterial community structure of continuous tomato cropping is better than that of CF, which should be used in the sustainable production of facility tomatoes.

Keywords: soil organic carbon, tomato, soil, bacterial community structure

1 Introduction

Microorganisms are a key component of the soil ecological chain. They play important roles in soil nutrient cycling, organic matter (OM) formation and decomposition, soil-borne disease occurrence and prevention, and crop growth and development. Additionally, soil microorganisms are crucial for maintaining soil ecological balance and fertility [1,2]. Therefore, maintaining high soil microbial activity and diversity is fundamental for current soil quality management practices.

Continuous protected tomato cropping is common in southern China. The continuous cropping of a single species often leads to an imbalance in the soil microbial flora, which manifests as a decrease in microbial diversity, an increase in pathogenic bacteria, and a decrease in the number of beneficial bacteria [3,4]. Furthermore, tomato production is often accompanied by excessive application of chemical fertilizers (CFs). This practice affects crop yield, soil physical and chemical characteristics, and quality and directly changes the structure of the soil microbial community [5,6]. Studies have shown that the long-term application of CFs can significantly reduce soil pH and microorganism number and diversity [6].

The use of organic materials (including livestock and poultry manure and straw) to fertilize continuous cropping soils in facilities has a high application value; it can also replace CFs and, therefore, reduce the volume used in facility agricultural production. The input of organic materials can effectively improve the structure of continuous cropping soil aggregates, increase the OM content, and reduce the salt content. Ding et al. [7] performed field experiments showing that long-term combined application of organic and inorganic fertilizers increased the
diversity of soil bacteria and fungi and promoted the growth of beneficial bacteria; conversely, single application of CFs promoted fungi growth. Long-term use of organic manure and straw returning in the field can also increase the abundance of flora such as growth-promoting bacteria (plant growth-promoting rhizobacteria, which promote plant growth and many microbial products that stimulate plant growth) and arbuscular mycorrhizal fungi, which promote soil health [8,9]. However, it is still unknown whether the input of organic materials can significantly enhance the structure of the bacterial community in continuous cropping soil.

China produces billions of tons of organic solid waste every year, including agricultural waste (e.g., straw), which has surpassed 800 million tons produced yearly. A large amount of straw is burned or discarded yearly, polluting the environment and wasting resources [10]. As of 2010, the total amount of livestock and poultry manure emissions in China has reached 1.9 billion tons, 227 million of which have not been properly treated to prevent environmental pollution [11]. This solid waste is rich in nutrients and OM needed for crop growth and development [12,13]. After being composted or adequately processed, organic solid waste can be used for fertilization in agricultural production, promoting sustainable development and reducing environmental pollution [13,14]. Therefore, in this study, we used high-throughput sequencing technology to analyze the effects of composting or earthworm treatment of livestock and poultry manure (vermicompost, VM) and CF on the composition and diversity of soil bacterial communities in facility tomato continuous cropping. We analyzed the feasibility of applying VM and CF in the sustainable production of facility agriculture and provided a theoretical basis for the rational fertilization of facility agriculture.

## 2 Materials and methods

### 2.1 Experimental design

The experiments were performed in a greenhouse in the Suzhou Polytechnic Institute of Agriculture (120.64 E, 31.43 N), Jiangsu, China, from March to June 2020. The greenhouse had natural light exposure, and the day and night temperature was controlled to be within 15–35°C.

This study used a completely random design. The fertilization treatments tested were VM, CF, and the control (CK) without fertilization. There were three treatments in total, and each was repeated five times. The total amount of N, P2O5, and K2O was equal in all fertilization treatments. To achieve the local general fertilization amount desired, the N, P2O5, and K2O dosages were set to 0.45, 0.20, and 0.40 g kg⁻¹ soil, respectively (approximately 850 kg N hm⁻², 552 kg P2O5 hm⁻², and 850 kg K2O hm⁻²). The soil’s total nitrogen (TN), total phosphor (TP), and total potassium (TK) were detected by a soil elemental analyzer (FlashSmart, ThermoFisher). The soil moisture content was 43%.

The amount of VM applied was 13.10 g kg⁻¹ soil (approximately 30 t hm⁻²), and the amount of RS was 45.52 g kg⁻¹ soil (approximately 95 t hm⁻²) (Table 1). Additional fertilizers (urea, superphosphate, and potassium sulfate) were added to VM and RS to achieve the same N, P2O5, and K2O content as the CF treatment. We added the total contents of urea, superphosphate, and potassium sulfate of 50, 35, and 30 kg, respectively. The nutrient contents of the fertilizers used are shown in Table 1.

All fertilizers were applied as a base fertilizer simultaneously to thoroughly mix the soil and fertilizer. No top dressing was performed during the tomato growth period.

### 2.2 Soil characteristics

The soil used in this study was utilized for 5 years of continuous tomato cropping in the greenhouse. Soil pH (soil/water = 1/5) and electrical conductivity (EC, soil/water = 1/5) were measured with a pH meter (INESA, Shanghai, China) and a DDS-307 conductivity meter (Shanghai Precise Science Instrument Co., China), respectively. The OM content was estimated by the potassium dichromate dilution calorimetry method [15]. Soil’s total carbon (TC) and TN were measured using a Vario EL III element analyzer (Elementar Co., Germany) [16]. Soil TK was detected by a soil elemental analyzer (FlashSmart, ThermoFisher).

| Treatment | pH | SOC (g kg⁻¹) | TN (g kg⁻¹) | TP (g kg⁻¹) | TK (g kg⁻¹) | C/N |
|-----------|----|--------------|-------------|-------------|-------------|-----|
| VM        | 6.38 | 2,035        | 46.8        | 17.85       | 4.65        | 7.4 |
| RS        | 5.92 | 5,830        | 450.5       | 8.35        | 6.53        | 52.69 |

Note: VM: vermicompost; RS: rice straw; SOC: soil organic carbon; TN: total nitrogen; TP: total phosphorus; TK: total potassium; C/N: soil organic carbon/total nitrogen.
The pH value and EC of the tested soil were 6.17 and 490 μS cm⁻¹, respectively, and the available nitrogen, phosphorus, and potassium contents were 206.9, 374.0, and 564.6 mg kg⁻¹, respectively. The soil organic carbon (SOC) and TN content were 46.63 and 3.07 g kg⁻¹, respectively, and the carbon-to-nitrogen ratio was 8.93. The tomato plants were cultivated in polyethylene pots (30 cm × 28 cm). After the air-dried soil was sieved with a 1 cm mesh, each pot was packed with 15 kg of soil with a bulk density of approximately 1.30 g cm⁻³.

The RS used in the experiment was collected from the experimental rice base of the Suzhou Polytechnic Institute of Agriculture, air-dried, and crushed to 2–3 cm before use.

On March 12, 2020, seedlings (with 3–4 true leaves) that grew sturdily, neatly, moderately, and relatively uniformly were selected and transplanted into pots. The irrigation was maintained during the growth period at 70% of the field water holding capacity and 80% of conventional field management. The test tomato variety used was “Zhongyan No. 988,” cultivated by the Beijing ZhongYanYiNong Seedling Co., Ltd. The entire growth period was 108 days.

2.3 Sample collection and measurement methods

Soil samples were collected during the tomato full fruit period (75 days). The soil samples were collected using a soil auger (8 cm in diameter). The soils were collected from 5 to 10 points along an S-shaped path within each treatment and mixed for a sample to ensure the representativeness of soil samples for each treatment, and immediately sent to the laboratory for analysis. A part of the fresh soil was passed through a 10-mesh sieve, and the root residues were removed and stored at −80°C for soil bacterial community and diversity analysis. Another part of fresh soil was air-dried and passed through 20- and 100-mesh sieves to determine the soil physical and chemical properties.

The total deoxyribonucleic acid (DNA) of each soil sample (0.5 g) was extracted using a Fast DNA SPIN Kit for Soil (MP Bio, USA), and the DNA was pre-checked with 1% agarose gel to assure purity. Next, the concentration and quality were tested by a NanoDrop 2000 UV spectrophotometer (ThermoFisher Scientific, USA); the qualified products were sent to Zoonbio Biotechnology Co., Ltd, to determine the composition of the soil microbial communities. Polymerase chain reaction (PCR) amplification and product purification were performed using diluted genomic DNA.

The HiFi Hotstart ReadMix PCR kit high-fidelity enzyme (KAPA Biosystems, USA) was used for PCR. PCR products were detected with 2% agarose gel electrophoresis, purified by gel extraction, and analyzed with a NanoDrop 2000 UV micro-spectrophotometer and 2% agarose gel electrophoresis to inspect the library quality. Products that passed the quality inspection were quantified with a Qubit (ThermoFisher Scientific, USA) and mixed in the corresponding proportions according to the data volume requirements of each sample.

The V3–V4 region, a highly variable region of ribosomal genes, was used for bacterial 16S rDNA amplification, and the universal primers used were F341 (5′-ACTCCTAGGGGRSGCAGCAG-3′) and R806 (5′-GGACTACVGGGTATCTA-3′). The index and linker sequences suitable for HiSeq2500 PE250 sequencing were added to the 5′-end of the universal primers. Amplified fragments of approximately 425 and 320 bp were obtained. After adding adapters, the products were sequenced using the HiSeq platform Illumina Miseq PE300 to obtain 2 × 300 bp Paired-End data, which was used to analyze the soil bacterial community. The Project accession number for raw data is SUB10527892.

2.4 Data analysis

QIIME (Version 1.7.0) was used to aggregate the 16S sequence data, and the Ribosomal Database Project (RDP) method was used for species clustering (similarity: 97%). Mothur software was used to analyze the bacterial diversity and richness index. Principal coordinate analysis (PCoA) was used in R to indicate the beta diversity of the soil bacterial community. Adonis analysis was used in R to indicate the soil bacterial community similarity. The Adonis analysis was performed by using the R software vegan package at the operational taxonomic unit (OTU) level. The OTU was used to determine significant differences between different fertilization treatments (P < 0.05); the data were used to construct a PCoA by using an R software vegan package at the OTU level. The Canoco 4.5 data package was used for redundancy analysis (RDA).

SPSS 17.0 (SPSS Inc., Chicago, IL, USA) was used for one-way analysis of variance and Pearson’s correlation analysis. The differences between treatments were tested with the Duncan test. The data in the graphs are all averages of five biological replicates.
3 Results and analysis

3.1 Soil physical and chemical properties

The pH of the soil treated with fertilization was significantly lower than that of CK (pH 7.20), with the most significant decrease occurring with CF (pH 4.54) and the minor decrease occurring with VM (pH 5.35) (Table 2). Compared with CK, fertilization treatments significantly increased the soil TN (6.01–26.61 g kg⁻¹), SOC (10.03–15.88 g kg⁻¹), TP (8.28–9.54 g kg⁻¹), and TK (5.25–6.35 g kg⁻¹) contents (Table 2).

3.2 Relative abundance of soil bacterial groups at different taxonomic levels

The 3153–3649 OTUs of the treated soil samples belonged to 35 phyla, 92 classes, 160 orders, 329 families, and 687 genera. At the level of bacterial phylum classification, the dominant phyla were Proteobacteria, Actinobacteria, Acidobacteria, Firmicutes, Gemmatimonadetes, Bacteroides, Candidatus Saccharibacteria, Chloroflexi, Verrucomicrobia, and Cyanobacteria (relative abundance >1%) (Figure 1). The relative abundance of the dominant bacterial groups, Proteobacteria and Actinobacteria, increased significantly after the application of organic materials (to 36.40–44.27% and 25.80–29.35%) compared with CK (35.88 and 23.10%); the relative abundance of Firmicutes and Bacteroidetes also increased significantly, but the relative abundance of Acidobacteria, Chloroflexi, Verrucomicrobiobio, Cyrobiobiosella, and Cyanobacteria was significantly reduced. CF treatments had the opposite effect on organic materials, leading to lower Proteobacteria and Actinobacteria abundances (23.46 and 19.13%). Proteobacteria had the highest abundance ratio in the CK treatment (44.27%), and the treatment with the highest abundance of Actinobacteria was VM (29.35%). These results show that the application of organic materials can increase the relative content of phylum-level dominant flora in the soil and decrease the abundance of other microbial flora to varying degrees.

Figure 1 shows the relative abundance of the bacterial genus in different treatments, in which the parts with an average abundance level lower than 0.5% are merged. They are insignificant in the CF (2.42%) treatment, compared with CK (3.65%) and VM (3.86%), slightly increasing the relative abundance of these bacteria. Gemmatimonas represented 4.72% of CM and 4.71% of VM; in CF, this dominant flora in the soil significantly reduced its relative abundance (3.29%). Gp6, Gp16, and Gp4 belong to the acid phylum (Acidobacteria), and their relative abundance is the highest in the CF treatments (8.0, 6.87, and 4.10%, respectively), followed by CK (6.33, 3.25, and 4.06%, respectively). Hence, the application of organic materials reduced the relative abundance of Gp6, Gp16, and Gp4 compared to CF and CK. Gaiella had the highest relative abundance in the VM treatment (3.78%) and the lowest in CF (2.45%). There were also several genera that changed significantly differed in abundance between different treatments. For example, Bacillus and Streptomyces had the highest abundance in CF and the lowest in VM. CK had the lowest. Porphyr- obacter was significantly more abundant in VM than in CK and CF, with increases ranging 1.08–1.61 times. The relative abundance of Hyphomicrobiun was the highest in VM and the lowest in CF.

The heatmap based on species showed the bacterial composition of different treatments (Figure A1). From Figure A3, we could infer that the bacterial compositions of VM and CF were more similar compared to that of CK, indicating that the fertilization treatment significantly changed the soil bacterial composition. Based on Figure A1, the g_staphylococcus was dominant in the VM and CF, whereas f_Peptostreptococcaceae and f_Pseudonocardiaceae were dominant in VM and CK, respectively.

Table 2: Soil physico-chemistry properties under different treatments

| Treatment | pH       | SOC (g kg⁻¹) | TN (g kg⁻¹) | TP (g kg⁻¹) | TK (g kg⁻¹) |
|-----------|----------|-------------|-------------|-------------|-------------|
| CK        | 7.20 ± 0.01a | 160.03 ± 6.21 | 20.20 ± 5.35a | 8.28 ± 3.28b | 5.25 ± 2.10c |
| CF        | 5.35 ± 0.02b | 510.50 ± 40.85 | 22.35 ± 5.14a | 15.35 ± 2.54a | 8.78 ± 1.55a |
| VM        | 4.54 ± 0.01c | 230.30 ± 5.85 | 22.61 ± 4.81a | 9.54 ± 3.25b | 6.35 ± 1.55b |

CK: control; CF: chemical fertilizer; VM: vermicompost; SOC: soil organic carbon; TN: total nitrogen; TP: total phosphorus; TK: total potassium; Lowercase letter indicates the Duncan test at the P < 0.05 level. The table content indicates mean ± standard deviations. All values are the mean values from three replicates.
3.3 Changes in soil bacterial community structure

The bacterial communities were analyzed by 16S rDNA sequencing of 18 soil samples obtained from 54,249 to 64,915 high-quality sequences and from 42,661 to 54,118 effective sequences (74–90% of high-quality sequences).

The average read length was 414.37 bp. The sequencing coverage rate was 97%, and a total of 3,153–3,649 OTUs were obtained, which shows that the gene sequences in the soil samples reflect the true soil bacterial community status.

The rarefaction curve indicates the sample sequencing depth and can be used to evaluate whether the sequencing volume is sufficient to cover all groups. Figure A2 shows the rarefaction curve for all samples in this test under the condition of similarity of 0.97. As shown in Figure 1, all soil sample dilution curves tended to flatten, indicating that sampling was reasonable, and the confidence in the bacterial community structure in the actual
environment was high, which could reflect the bacterial community of a soil sample in a relatively realistic way (Table 3).

The soil Chao1 and Shannon and Simpson indices differed significantly ($P < 0.05$) between the six treatments. The Chao1 index of the different fertilization modes ranked $\text{CK} > \text{VM} > \text{CF}$ in our analysis; VM and CF treatments led to values significantly lower than CK, whereas there was no significant difference between other treatments and CK. The Shannon and Simpson indices were the highest for VM and the lowest for CF; they were significantly higher for VM than for CF treatment. The Chao1 index was significantly negatively correlated with $\text{NH}_4^+$. The Shannon index was significantly negatively correlated with EC and was significantly positively correlated with pH and TC. The Simpson index was significantly negatively correlated with EC and had a significant positive correlation with SOC (Table 4, $P < 0.05$).

PCoA was used to show the soil bacterial beta diversity (Figure 2). The results showed that among all the treatments, the soil bacterial community compositions of VM and CF were the most similar. The results showed that the first (PCoA1) and the second (PCoA2) principal component axis contributed, respectively, 18.91 and 13.81% to the difference between fertilization treatments. The distances between CF and VM were the closest, indicating that the soil OTU composition of the three was the most similar. This indicates that fertilization changed the composition of the bacterial community in continuous cropping soil, especially for the organic fertilizer treatment. The Adonis results indicate that the soil bacterial structure was significantly changed among CK, CF, and VM (Adonis $P < 0.05$, Figure A3).

A Venn diagram illustrates the number of common and unique OTUs in three treatments, as shown in Figure A4. In total, 2,126 OTUs were detected, of which 1,238 (58.23%) were shared among the CK, CF, and VM soils (Figure A4). There were only 320 OTUs found in CK, accounting for 15.05% of the total. CF soils produced 110 OTUs, accounting for (5.17%), and the OTUs specific for VM were 9, accounting for 0.40%.

### 3.4 Correlation between soil bacterial groups and chemical properties

To clarify the environmental factors that change the structure of the soil microbial community, we analyzed the correlation between the dominant flora at the bacterial phylum and the genus level and the soil chemical properties (Table 5 and Figure 3). At the phylum classification level, Actinomycetes were negatively correlated with EC and significantly positively correlated with $\text{NO}_3^-$, $\text{NH}_4^+$, TC, TN, and OM. Proteobacteria and Blastomonas were significantly negatively correlated with EC and significantly positively correlated with TC. The acid bacterium phylum was significantly positively correlated with EC and significantly negatively correlated with $\text{NH}_4^+$, TC, and OM. Firmicutes were negatively correlated with EC and significantly positively correlated with pH and TC. The Simpson index was significantly negatively correlated with EC and had a significant positive correlation with SOC (Table 4, $P < 0.05$).
with pH, NO$_3^-$, NH$_4^+$, TC, TN, and OM; the correlation with Cyanobacteria/Chloroplast was the opposite. Furthermore, the Bacteroides phylum was negatively correlated with EC and positively correlated with NH$_4^+$, TC, and OM; the verrucomicrobial phylum had the opposite correlations. Candidate Saccharibacteria were significantly negatively correlated with pH and positively correlated with EC, NO$_3^-$, NH$_4^+$, and OM. Finally, the phylum Chloroflexus was positively correlated with EC, TC, and TN (Table 5).

RDA results showed that the first and second ordination axes explained 56.0 and 8.6% of the bacterial community changes, respectively, and the two together explained 64.6% of the bacterial community changes (Figure 3). On the first sorting axis, the bacterial communities of organic material treatment (VM) were significantly changed compared with those of CF and CK, and they were distributed on both sides of the first axis. The overall degree of explanation of soil chemical properties for changes in bacterial community structure was 75.2%, of which NH$_4^+$ ($F = 13.237, P = 0.002$) and EC ($F = 15.443, P = 0.002$) were the most relevant, accounting for 32.1 and 24.7% of the explanatory degree, respectively. The effects of pH ($F = 5.944, P = 0.004, 8.0\%$) and TC ($F = 4.462, P = 0.008, 5.3\%$) were the second most important. Comprehensive analysis showed that NH$_4^+$, EC, pH, and TC are soil physical and chemical indicators that are highly correlated with changes in soil bacterial communities.

### 4 Discussion

Soil microbial diversity is an important factor in maintaining soil health [17]. Studies have shown that continuous cropping reduces the microbial diversity around the rhizosphere and increases the abundance of pathogens,

| Phylum         | pH  | SOC  | TN   | TP   | TK   | C/N  |
|----------------|-----|------|------|------|------|------|
| Proteobacteria | −0.214 | 0.803* | 0.215 | 0.072 | 0.305 | 0.264 |
| Acidobacteria  | 0.154 | −0.735* | −0.284 | −0.183 | −0.528* | 0.969* |
| Actinobacteria | −0.228 | 0.864* | 0.891* | 0.314 | 0.872* | −0.454* |
| Chloroflexi    | 0.084 | −0.682* | 0.23 | 0.022 | −0.402* | −0.204 |
| Gemmatimonadetes | 0.286 | 0.430* | 0.136 | 0.801* | 0.785* | 0.343 |
| Bacteroidetes  | −0.028 | 0.785* | 0.24 | 0.892 | 0.234 | 0.787* |

* $P < 0.05$. The table content indicates the correlation coefficient by Pearson’s correlation analysis. * $P < 0.05$.

SOC: soil organic carbon; TN: total nitrogen; TP: total phosphorus; TK: total potassium; C/N: soil organic carbon/total nitrogen.
leading to the occurrence of root diseases [18]. For example, Li et al. [19] analyzed the microbial communities in rhizosphere soil samples of healthy tomatoes and diseased tomatoes suffering from bacterial wilt (*Ralstonia solanacearum*) that were grown continuously for 3 years and showed that the soil microbial diversity was higher in healthy tomato than in diseased tomato. Therefore, improving the diversity of soil microorganisms is key to alleviating soil-borne diseases in continuous cropping soil. This study showed that after fertilization with VM and CM, the soil bacterial alpha diversity increased in continuous tomato cropping and decreased after applying CFs. That might be because the application of VM improved the soil aggregate structure and chemical properties (such as pH and EC value), improving soil microbial diversity [20]. Furthermore, the increase in soil nutrients through the application of VM, especially the increase of OM (Table 2), can promote the growth of soil microorganisms [21]. Considering the chemical properties of the soil, the application of VM can significantly increase the EC value, and there is a risk of secondary soil salinization after long-term application. Hence, fertilization with earthworm manure might be a more appropriate alternative. Similar to our results, Khan et al. [22] compared the effects of long-term application of VM and inorganic fertilizer to soil used for cucumber cultivation and found that the soil microbial diversity treated was the highest with VM treatments. VM is the product of semi-decomposed agricultural or food waste that has been processed by the earthworm intestinal system. Its microbial diversity and aggregate structure are better than traditional compost [23,24]. Soils with high bacterial community diversity have functional redundancy among the communities, and the fluctuation of certain microbial groups has little effect on the overall function of soil microorganisms. Therefore, increasing bacterial community diversity can effectively maintain soil ecological functions and health [17]. The application of VM can increase the diversity of the bacterial community in the continuous cropping soil of facility-grown tomatoes, thereby improving soil disease resistance.

Ai et al. [25] found that soil disease resistance is positively correlated with the relative abundance of Proteobacteria and Actinomycota in rhizosphere soil. In rhizosphere soils with low plant incidence, the total number of these microorganisms is higher than in rhizosphere soil with high plant incidence. This may be due to the strong disease resistance of some unique microorganisms belonging to these categories [26]. We found that the relative abundances of Actinomycetes, Firmicutes, and Bacteroides were significantly increased after applying organic materials, especially VM (Figure 2), which contrasts with the results obtained by Arjun et al. [27]. This indicates that the application of VM has the potential to improve the resistance to soil pathogens. It is generally believed that the phylum Proteobacteria, which includes trophic bacteria [28], can use complex OM and plant residues as carbon and nitrogen sources [29]. Here, the content of nutrients (NH₄, NO₃, TC, TN, and OM) increased after applying organic materials to the soil (Table 2), which stimulated the growth of Proteobacteria. Our results show a positive correlation between Proteobacteria and TC content, which may be because these bacteria are more sensitive to carbon nutrients [30]. Actinomycete bacteria can produce a variety of secondary metabolites (antibiotics) and extracellular enzymes that play an important role in the defense against plant diseases. Borrero et al. [31] showed that some Actinomycota bacteria have a good inhibitory effect on *Fusarium oxysporum*, the pathogen that causes Fusarium wilt. Hertweck [32] performed whole-genome sequencing on a microbial strain of the genus Ilumato bacter and found that it contained two type I polyketide synthase enzymes that synthesize polyketides, which are highly effective antibiotics. The increase in beneficial microorganisms caused by VM application may be due to the intestinal microorganisms of earthworms. Previous studies have shown that after organic materials are digested in earthworm intestines, the feces produced contain more beneficial microorganisms than the original organic materials [33]. After these beneficial microorganisms enter the soil with VM, they grow and multiply under suitable environmental conditions, promoting the defense of soil ecosystems against soil-borne diseases.

5 Conclusion

VM application has greater advantages than CF, RS, and conventional compost by adjusting soil pH and maintaining low soil salinity. In addition, the rich nutrient content and better soil aggregate structure in the soil VM fertilization can significantly improve the soil microbial environment and diversity, especially by increasing the abundance of some microorganisms that have an anti-pathogenic effect, which is useful to overcome the limitations of continuous cropping.

Acknowledgments: The authors thank Chengzhong Wang from Suzhou Polytechnic Institute of Agriculture, Suzhou, for providing assistance.
Funding information: The work was financially supported by the Basic Research on Suzhou Science and Technology Planning Project (SNG2020062) and SZAI Science and Technology Incubation Project (PY2104).

Author contributions: Jun Chen, Yichun Du and Wei Zhu designed and did the experiment and prepared this manuscript. Xin Pan helped to finish the bioinformatic analysis. Zhen Wang revised this manuscript and language editing. All coauthors contributed to manuscript editing.

Conflict of interest: The authors state no conflict of interest.

Data availability statement: The datasets generated during and/or analyzed during the current study are available from the corresponding author on reasonable request.

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Appendix

Figure A1: Heatmap diagram based on specie level of bacteria from the different soil samples. CK: control; CF: chemical fertilizer; VM: vermicompost.
Figure A3: Distance box plot of soil bacteria from the different soil samples based on Adonis analysis. CK: control; CF: chemical fertilizer; VM: vermicompost.

Figure A2: Rarefraction diagram of soil bacteria from the different soil samples. CK: control; CF: chemical fertilizer; VM: vermicompost.

Figure A4: Venn diagram showing the numbers of shared and exclusive OTUs of bacteria from the different soil samples. CK: control; CF: chemical fertilizer; VM: vermicompost.

Figure A3: Distance box plot of soil bacteria from the different soil samples based on Adonis analysis. CK: control; CF: chemical fertilizer; VM: vermicompost.