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Influence of different biological control agents and compost on total and nitrification-driven microbial communities at rhizosphere and soil level in a lettuce - Fusarium oxysporum f. sp. lactucae pathosystem

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1	Influence of different biological control agents and compost on the rhizosphere and soil total
2	and nitrification driving microbial communities in a lettuce – Fusarium oxysporum f.sp.
3	lactucae pathosystem
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28 Abstract

29 The response of rhizosphere and bulk soil indigenous total microbial communities focusing on 30 nitrifying prokaryotes was evaluated after the application of different biological control agents 31 (BCAs) (Bacillus subtilis, Trichoderma asperellum + Trichoderma gamsii, Trichoderma W2, 32 Pseudomonas putida) and compost in controlling lettuce Fusarium wilt. Experiments were conducted "in situ" over two lettuce cropping seasons. Total fungal, bacterial and archaeal 33 34 microbial populations, as well as the ammonia oxidizing groups (i.e., AOB, AOA) and the pathogen, Fusarium oxysporum f.sp. lactucae (FOL) have been screen out using quantitative 35 polymerase chain reaction (qPCR) method. The BCAs - like communities and different antifungal 36 37 functional genes (fungal chitinase chiA, and bacterial 2,4-diacetylphloroglucinol - phlD and HCN 38 synthase - hcnAB abundances have been also assessed. Quantitative data were corroborated with the 39 disease index, the potential nitrification activity and chemical parameters of the soil.

40 The application of BCAs and compost resulted in a significantly lower disease severity than the 41 untreated control, and Fusarium wilt was reduced by as much as 60%. This result was confirmed by 42 significant negative correlations between abundance of BCA-like microbial population and disease 43 severity. The BCAs and compost treatments did not negatively affect the microbial communities at 44 the rhizosphere or bulk soil level. The presence of FOL in the untreated control resulted in a niche 45 differentiation of the nitrifiers. Overall, the observed decreased FOL abundance, the increased 46 abundance of inoculum-like populations and the active response of nitrifiers community pointed out the sustainability of the used treatments which achieved an important level of disease suppression. 47

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50 Keywords: BCAs; disease suppression; nitrifying prokaryotes; *chiA* gene, *phlD* gene; *hcnAB* gene
51 1. Introduction

52 Lettuce is cultivated and consumed by the whole world and its production exceeded 24 million tons 53 in 2014 (FAO, 2014). Lettuce crops are grown intensively in Mediterranean regions. Spain, Italy 54 and France are the three most important producing countries, with 35, 21, and 13%, respectively, of 55 lettuce production in Europe in 2010 (Eurostat, 2012). Fusarium oxysporum f. sp. lactucae (FOL) is 56 a soil-borne, host-specific lettuce pathogen that has been observed in Italy since 2002 (Garibaldi et 57 al., 2002), which causes serious economic losses (Matheron & Gullino, 2012). Its management has 58 been investigated intensively through different methods, such as solarization (Matheron and 59 Porchas, 2010) or the application of different organic amendments (Franceschini et al., 2016; 60 Gilardi et al., 2016). However, the seed transmission of this pathogen makes soil disinfection only 61 partially effective (Garibaldi et al., 2004; Gilardi et al., 2007). For these reasons, the use of 62 biological control agents (BCAs), based on antagonistic microorganisms or different composts (e.g., 63 compost with inoculated BCA, green compost), is considered an effective and sustainable strategy 64 (Barrière et al., 2014; Pugliese et al., 2015). Species of the Trichoderma genus (T. harzianum, T. 65 *viridi*) have been shown to be efficient "biofungicides", as they are involved in the suppression of 66 different pathogens, including Fusarium oxysporum f. sp. lactucae (Gilardi et al., 2007, 2013; 67 Innocenti et., 2015). Pseudomonas spp. is known to be responsible for the "natural" suppressiveness 68 of some soils against soil-borne pathogens (e.g., Pythium ultimum, Rhizoctonia solani) (Weller et 69 al., 2002; 2007), and the same phenomenon has also been observed for various *Bacillus* species 70 (e.g., B. subtilis, B. thuringiensis, B. amyloliquefaciens) (Cawoy et al., 2011; Kloepper et al., 2004). 71 Many studies have been conducted on the effects of BCAs and compost on disease control 72 (Bonanomi et al. 2007; Berg, 2009; Pane et al. 2013). The presence of BCAs candidates in compost, 73 belonging to the Trichoderma, Pseudomonas and Bacillus genera, has also been demonstrated 74 (Pugliese et al., 2008).

BCAs are known to be aggressive colonizers that indirectly affect the functional guilds in the rhizosphere environment (Gupta et al., 2012). However, their introduction into a new soil system is a complex process which may disturb the indigenous soil microorganisms that play crucial roles in 78 the essential nutrient cycles, especially nitrogen (N) cycle. A critical component of the microbially 79 driven N cycle is the nitrification process, encoded by amoA as key genes that controls the 80 availability of soil nitrates (Nicol et al., 2008; Zhang et al., 2013). N supply and form affect lettuce plant defense and FOL pathogenicity (Orr and Nelson, 2018). Nevertheless, little information are 81 82 available regarding the impact of BCAs and compost may have on ammonia-oxidizing bacteria or 83 on the archaea that harbor the amoA gene (AOB and AOA) (Musyoki et al., 2014; Zimmermann et 84 al., 2016), and in consequence on the soil N form which may further control FOL suppression. 85 Hence, there is a critical need to study the potential side - effects of the applied treatments on both 86 AOB and AOA.

87 The primary objective of this study was to explore the direct impact of BCAs and compost 88 treatments "in situ" on the abundance of total fungi, bacteria and archaeal communities, as well as 89 on the ammonia oxidizing groups (i.e., AOB, AOA) in the FOL affected lettuce rhizosphere and 90 bulk soil. Moreover, fungal and bacterial recruitment upon pathogen infection, has been assessed by 91 studying three anti-pathogen genes that are usually correlated with disease suppression: the 92 Trichoderma based chitinase chiA gene, and two well-described bacterial antifungal genes: phlD 93 (coding for 2,4-diacetylphloroglucinol - 2,4-DAPG) and hcnAB (coding for hydrogen cyanide -94 HCN).

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- 102
- 103 2. Materials and Methods

104 2.1. Layout of the experiments

The experiments were carried out in tunnels (6 x 60 m) on a commercial farm in Moretta (Cuneo, Northern Italy), which had a history of several lettuce cycles prior to the beginning of this study. The soil was characterized as a silty loam soil (silt : sand : clay 60 : 30 : 10 %, with neutral pH and 1.5 % organic matter) naturally infested with the FOL pathogen.

109 The lettuce cultivar 'Novelsky', which belongs to the batavia type and is moderately susceptible to 110 Fusarium wilt was used (Gilardi et al., 2017). The experimental trials were carried out over two 111 years, 2016 and 2017, in order to test the efficacy of pre-planting treatments with different 112 commercially available Bacillus subtilis: Serenade Max - SM, QST713, Bayer Crop Science, Italy 113 and Trichoderma gamsii + Trichoderma asperellum: Remedier - RM ISAGRO, Italy) and 114 experimental BCAs (Trichoderma, TW2, AgriNewTech, Italy and Pseudomonas putida - Pp, Agroinnova, Italy) (Table 1) against lettuce wilting. In addition, two composts (Ant's Compost V 115 116 and Ant's Compost M; AgriNewTech, Italy), produced from green wastes, in a dynamic industrial 117 treatment system, were used (Table 1): one green compost - CV and the same compost inoculated 118 with Trichoderma TW2 - CM. An untreated control was used to monitor Fusarium wilt 119 development. Summarized details of these treatments can be found in Table 1.

Lettuce seeds were sown in 100-plug trays (3.4 cm diameter pots, 4 l of soil, 53 x 42 cm surface)
filled with a peat mixture substrate.

122 Four substrate (i.e., peat) treatments with BCAs were carried out on the plug tray between sowing 123 and transplanting (at the first true leaf stage) in a commercial nursery; incubation until transplanting 124 was conducted under greenhouse conditions at a temperature of 22-24°C (Table 1). The BCAs were 125 applied by spraying them onto the peat surface in a high volume of water (500 ml/tray) using a 1 l 126 capacity hand sprayer. The compost products were distributed in a thin layer of 8 g/plug at sowing 127 (T0) over the surface of 53 x 42 cm plug trays, then mixed with the soil immediately before 128 transplanting (T20). Azoxystrobin (Ortiva, 23.2% a. i., Syngenta Crop Protection, Italy) was used as 129 the reference chemical treatment and was applied once as a soil drench at transplanting.

Twenty day old lettuce plants were transplanted (at T20) at a density of 24 plants/m², drip irrigated 130 131 and grown according to the cultural practices adopted by commercial growers in the region. Each 132 treatment had four biological replicates arranged in a completely randomized block design. The 133 plants were monitored weekly for symptoms development, and the data were recorded starting from 134 the appearance of the first symptoms (yellow leaves and reduced growth). The final disease rating 135 was carried out 48-53 days after transplanting on 16 plants/treatment by dissecting each plant. The 136 disease severity (DS) scale was as follows: 0 = healthy plant, 25% = initial leaf chlorosis, 50% =137 severe leaf chlorosis and initial symptoms of wilting during the hottest hours of the day, 75% =138 severe wilting and severe symptoms of leaf chlorosis; 100% = plant totally wilted, leaves 139 completely necrotic. Lettuce yield was measured at the end of the trials by collecting 12 plants per 140 treatment, and determining the fresh weight of the plants.

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142 2.2. Rhizosphere and bulk soil sampling

143 The influence of soil-treatments with BCAs and compost on indigenous microbial communities was 144 studied at the rhizosphere and bulk soil levels by collecting samples at the end of the trials (the end 145 of September 2016 and 2017). One composite rhizosphere sample per plot consisted of the roots of 146 five randomly selected lettuce plants. The roots were shaken vigorously to remove the soil particles 147 that were not tightly adhering on them. The rhizosphere soil was gently scraped off with sterile 148 forceps and transferred into sterile sampling bags. Each treatment consisted of three biological 149 replicates, which were kept on ice. Soil bulk samples were taken also in three replicates (0-15 cm 150 soil depth) per treatment. Each replicate consisted of five sub-replicates collected from the same 151 plots as the rhizosphere samples. The replicates were sampled in a W - shaped sampling pattern, to 152 obtain a good bulk soil representation and homogeneity. The samples were then placed in plastic 153 bags, kept on ice and transported to the laboratory.

154 The fresh rhizosphere and bulk soil samples were homogenized, passed through a 2 mm sieve and 155 stored at 4°C. All the samples were split into two parts: one part was sent to the Regional Chemistry Laboratory (Turin, Italy) for further geochemical analysis: humidity, pH, total N (TN), inorganic N as nitrate (NO_3^-) and ammonium (NH_4^+) , total organic carbon (TOC), total phosphorus (TP) and available phosphorus (AP). The second part was used for enzymatic measurements (i.e., potential nitrification activity – PNA) and further microbiological and molecular investigations.

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161 2.3. Molecular analyses

162 2.3.1. Rhizosphere and bulk soils DNA extraction

A NucleoSpin® Soil kit (Macherey-Nagel GmbH & Co. KG, Düren, Germany) was used, according to the manufacturer's instructions, for the rhizosphere and bulk soil genomic DNA extraction. Extraction was conducted on fresh samples (250-500 mg of sample material). DNA quantity and purity were measured spectrophotometrically using a NanoDrop ND-1000 (NanoDrop Technologies, Wilmington, DE, USA).

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169 2.3.2 Microbial abundance (quantitative PCR (qPCR) assays)

170 The abundance of the bacterial and archaeal 16S rRNA genes, fungal 18S rRNA genes and of five 171 functional genes (i.e., bacterial ammonia monooxygenase amoA gene - AOB, archaeal amoA gene -172 AOA, fungal chitinase chiA gene, and bacterial 2,4-diacetylphloroglucinol phlD and HCN synthase *hcnAB* genes) was determined by means of qPCR, using a StepOnePlus[™] Real-Time PCR System 173 174 (Applied Biosystems, Foster City, CA, USA) for the rhizosphere and bulk soil DNA samples. In 175 addition, the abundance of FOL and BCAs - like microorganisms (i.e., Bacillus, Trichoderma and 176 Pseudomonas) was also investigated. A description of the primer sets and amplification details are 177 given in Table 2.

The primers for fungal chitinase *chiA* gene quantification were designed as follows: DNA from rhizosphere and bulk soil samples from the treatments with *Trichoderma* sp. was used to amplify the fungal chitinase *chi42* gene, according to the Nguyen HL et al., (2001) protocol, with CHIf/CHIr primers, which generally amplify a 1450 bp sequence. Four new primer pairs targeting

182 fragment of chi42 an internal the gene, were designed using Gen Script (http://www.genscript.com/tools.html#biology), considering relatively high annealing temperatures 183 184 $(\geq 60 \text{ °C})$ and smaller amplicon sizes than 200 bp to reduce generation of artifacts and to achieve 185 appropriate reaction efficiencies during qPCR. The accuracy of the primers was confirmed using 186 BLAST of the National Centre for Biotechnology Information (http://www.ncbi.nlm.nih.gov); 187 preliminary tests (data not shown) were carried out to assess their specificity, and the best primer 188 pair was then chosen. The designed primer pair (chiaxF 189 (ACCCTGCCGATGACACTCAG)::chiaxR (GGCAGCGATGGAGAGAAGGA) generated an 190 of 122 with following amplicon bp, the sequence: 191 5'ACCCTGCCGATGACACTCAGGCCACCAACATGGTTCTTCTGCTCAAGGAGATCCGAT 192 193 GCTGCC-3'.

194 Amplicons from each target gene were generated for the standard preparation, purified (Invisorb 195 Fragment CleanUp, Stratec Molecular GmbH, Berlin, Germany), and ligated in Strata-Clone PCR 196 cloning vector pSC-A (Strataclone PCR Cloning Kit, Agilent Technologies Inc.); the ligation 197 products were then transformed into StrataClone SoloPack competent cells (Agilent Technologies 198 Inc.). The specificity of the clones used as qPCR standards was checked via sequencing, at LGC 199 Genomics GmbH (Berlin, Germany) and through BLAST analysis. Plasmid DNA was isolated 200 (GenEluteTM Plasmid Miniprep Kit, Sigma-Aldrich, St. Louis, MO, USA) from standard clones and 201 quantified as described above.

As assessed in preliminary tests and in a qPCR assay optimization, the final volume of the qPCR reaction cocktails was 25 μ l for the 16S and 18S rRNA genes, the bacterial and archaeal *amoA* gene, and the FOL and BCA - like microorganisms quantification, and it was 20 μ l for the *chiaA*, *phlD* and *hcnAB* genes. All the quantifications were conducted with a 10 ng DNA template, except for the bacterial 16S rRNA gene, for which a 5 ng DNA template was used (Cucu et al., 2017). The reaction mixtures contained 1x Power SYBR green master mix (Applied Biosystems), 0.12 μ M of 208 each oligonucleotide (Table 2) for the total bacteria, archaea, fungi and FOL, 0.15 μ M for the *phlD* 209 and *hcnAB* genes, 0.32 μ M for the AOB, AOA and BCA - microorganisms and 0.4 μ M for the *chiA* 210 gene.

211 All the considered genes were quantified in triplicate across plates, while standards were run in 212 duplicate in 10-fold serial dilutions. The optimal dilution of the DNA extracts was tested to 213 compensate for any inhibitory reactions by the organic compounds (e.g., humic acids) co-extracted 214 during the DNA isolations. In all experiments, negative controls containing no-template DNA were 215 subjected to the same qPCR procedure to detect and exclude any possible contamination. 216 Amplification efficiency ranged from 96%, (archaeal 16S rRNA, fungal 18S rRNA, BCAs and their 217 functional genes) to 103% (bacterial 16S rRNA). For the nitrifiers, the amplification efficiency 218 were 95.3% and 99.1% for AOB and AOA, respectively. The R2 was always ≥ 0.98 . Melting curves 219 of the amplicons were generated to ensure that the fluorescence signals originated from specific 220 amplicons and not from primer dimers or other artifacts. This was confirmed by checking the 221 amplification products on 1% agarose gel. Gene copy numbers were calculated with StepOne[™] 222 software, version 2.2 (Applied Biosystems). The data were normalized and presented as log copies g⁻¹ dry soil. 223

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225 2.4. Chemical properties of rhizosphere and bulk soil samples

In short, the pH values were measured in water suspensions at a solid: liquid ratio of 1:2.5. The total organic carbon (TOC) and total nitrogen (TN) were quantified using a Leco Tru Spect CN automatic analyzer. Ammonium (NH_4^+) and nitrate (NO_3^-) were measured colorimetrically, by means of a continuous flow auto-analyzer (Alliance Evolution II), using standard colorimetric techniques. The total phosphorus (TP) was determined by means of 'ICP Varian mod. Liberty LR', after microwave digestion with hydrogen peroxide, hydrochloric acid and nitric acid, filtration and dilution. The available P (AP) was extracted using sodium bicarbonate and determined by means of the molybdenum blue method (Olsen et al., 1954), modified for continuous flow colorimetricanalysis (Alliance Evolution II).

The potential nitrification activity (PNA) was determined according to the procedures described byHoffmann et al. (2007).

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238 2.5. Statistical analyses

The data on disease index, gene abundance, chemical properties of the soil and potential 239 240 nitrification activity were subjected to a linear mixed model, with year considered as the random 241 variable, using R software (Software R 3.0.1, R foundation for Statistical Computing, Vienna, 242 Austria, http://www.R-project.org). The effects of different BCAs and compost on the abundance of 243 the studied genes, on the chemical properties of the soil and on disease severity were evaluated. All 244 data were subjected to a Levene test to check for the homogeneity of variance and normality was 245 tested on the residuals using the Shapiro-Wilk test; when not normally distributed data from disease 246 severity (DS) were arcsin transformed while, the data from microorganisms abundance were log-247 transformed and normality was checked again. The means were separated by Bonferroni test. The 248 statistical analysis included treatment rhizosphere soil, bulk soil, year, treatment \times year.

Pearson's linear correlation coefficients were calculated for assessing the relations between
 microbial gene abundance, disease severity and PNA, as well as between *Pseudomonas*, *Bacillus*,
 Trichoderma and FOL abundance and soil chemical properties.

Linear regression between total microbial gene abundance and the functional genes of N cycle (bacterial *amoA*, archaeal *amoA*, *chiA* gene) with soil chemical data were calculated to evaluate how much of their abundance is explained by variation in the soil chemical data.

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- 258 **3. Results**

259 3.1. Disease severity and plant biomass

The lettuce plants in the non-treated control plots showed an average disease severity DS of up to 260 261 60% at the end of both trials (Table 3). The BCA treated plots provided significantly lower disease 262 severities than the untreated ones, and statistically similar results to one application of azoxystrobin, 263 reducing Fusarium wilt by 47.2 to 58.5% in the two years, respectively. As a consequence, the fresh 264 weight of the lettuce was significantly higher in all the treatments compared to control, without statistically differences among them (Table 3). No significant difference was observed between the 265 266 disease severity data. A significant time effect for fresh weight of the lettuce between trials was noted (p < 0.001). However, the *Pseudomonas* and *Trichoderma* treatments were less effective in 267 268 the second trial.

269

- 270 3.2. Microbial abundance
- 271 *3.2.1. FOL abundance*

272 In general, FOL abundance assessed by gene copy quantification was significantly higher in the 273 bulk soil than in the rhizosphere soil. Almost all the treatments resulted in a significant reduction of 274 pathogen - associate gene copies compared to the untreated control - C, in both the rhizosphere and 275 bulk soil at the end of both trials (2016 and 2017 respectively) (Table 4). At the end of first trial 276 (2016), Serenade max - SM - Bacillus treatment was not significantly different compared with the 277 untreated control - C at the rhizosphere level, while at the end of the second trial (2017) the 278 Pseudomonas - Pp treatment was not significantly different compared with the untreated control -279 C. The interaction treatment \times year was not significant (Table 4).

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- 284 *3.2.2. Total microbial population abundance*

At the end of both trials (2016 and 2017 respectively) the number of gene copy of bacterial, archaeal and fungal populations showed an important homogeneity, with no significant interaction treatment × year. Therefore the data were presented as average of all values from both trials.

288 Overall, at the end of both trials (2016 and 2017 respectively) the bacterial 16S rRNA abundance 289 was significantly higher at the bulk soil level than at the rhizosphere level (Figure 1a). The effect of 290 BCAs treatments was more evident in the rhizosphere; the Remedier - RM - Trichoderma spp. and 291 Pseudomonas - Pp resulted in a lower bacterial abundance than the other treatments but not 292 significantly different than the untreated control - C, while the Serenade max - SM - Bacillus 293 treatment resulted in a significantly higher bacterial abundance than the untreated control - C. 294 However, the bacterial community abundance was more or less stable at the soil level over all the 295 treatments (Figure 1a). The archaeal 16S rRNA gene copy number in the rhizosphere followed 296 almost the same trend as the overall bacterial counterpart. The green compost - CV and 297 Trichoderma W2 - TW2 treatments resulted in slightly higher archaeal abundance in the bulk soil 298 samples than the untreated control - C (Figure 1b). The fungal 18S rRNA abundance was 299 significantly higher in the rhizosphere than in the bulk soil samples, in all the treatments as well as 300 in the untreated control - C. Remedier - RM - Trichoderma spp. inoculation resulted in a slightly 301 higher fungal abundance than the untreated control - C, while the total fungal community in the soil 302 samples was more abundant in the Serenade max - SM - Bacillus, Pseudomonas - Pp and green 303 compost - CV treatments (Figure 1c).

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305 *3.2.3. Functional gene abundance*

In general, the ammonia-oxidizing bacterial (AOB) gene abundance was significantly influenced by treatments and was significantly higher in the bulk soil than in the rhizosphere samples (Table 5). After the first trial, 2016, the AOB gene in the bulk soil samples was more abundant in the untreated control - C, while the lower abundance was observed after Serenade max - SM - *Bacillus* treatment. At the rhizosphere soil level the Serenade max- SM - *Bacillus*, green compost - CV and 311 *Trichoderma* W2 – TW2 treatments resulted in a highest AOB gene abundance while no significant 312 difference was observed between Remedier - RM-*Trichoderma* spp., *Pseudomonas* – Pp and the 313 untreated control - C. After the second trial, 2017, the same trend of AOB abundance was observed 314 at the bulk soil level, while at the rhizosphere soil level the Remedier - RM-*Trichoderma* spp. and 315 *Pseudomonas* – Pp treatments resulted in the lowest AOB abundance. The interaction treatment × 316 year was not significant. (Table 5).

In general, the ammonia-oxidizing archaeal (AOA) gene abundance was significantly influenced by
treatments and was significantly higher in the bulk soil than in the rhizosphere soil samples (Table
5).

320 All the treatments resulted in higher archaeal *amoA* gene abundance than the untreated control - C, 321 at both rhizosphere and bulk soil levels for both trials, 2016 and 2017 respectively. After the first 322 trial, 2016, the green compost - CV treatment resulted in the highest AOA abundance than the other 323 treatments at the bulk soil level, while the Serenade max - SM - Bacillus and Trichoderma W2 -324 TW2 treatments resulted in the highest abundance than the other treatments at the rhizosphere soil 325 level. After the second trial, 2017, the green compost - CV and the Remedier - RM-Trichoderma 326 spp treatments resulted in the highest AOA abundance at the bulk soil level, while the Serenade 327 max - SM - Bacillus treatment resulted in the highest AOA abundance at the rhizosphere level. The 328 interaction treatment \times year was not significant.

329 In general, the *chiA* gene (*Trichoderma* based chitinase) abundance was significantly influenced by 330 treatments (Table 6). After the first trial, 2016, the abundance of the chiA gene was higher in the rhizosphere soil samples for the Trichoderma TW2 – TW2 and Remedier - RM-Trichoderma spp 331 332 treatments. However the Remedier - RM-Trichoderma spp was not statistically different compared 333 with the inoculated compost - CM. The Pseudomonas - Pp and the green compost - CV treatments 334 resulted in the lowest chiA gene abundance compared with the other treatments, but not 335 significantly different compared with the untreated control - C. At the bulk soil level, the chiA gene 336 was more abundant in the Remedier - RM-Trichoderma spp, Pseudomonas - Pp and green compost - CV treatments comparing with the other treatments and with the untreated control - C (Table 6).
After the second trial, 2017, all the treatments at the rhizosphere soil level resulted in significant
differences between the *chiA* gene abundances. The highest *chiA* gene abundance was observed in
the *Trichoderma* W2 - TW2 treatment. Also at the bulk soil level all the treatments resulted in
higher *chiA* abundance compared with the untreated control - C. The highest *chiA* gene abundance
was observed in the *Pseudomonas* - Pp and green compost - CV treatments.

343 The abundance of the *phlD* gene was significantly higher in the rhizosphere than in the bulk soil 344 samples for all the treatments and for the untreated control - C (Table 6). After the first trial, 2016, 345 Pseudomonas - Pp and the green compost - CV treatments resulted in the highest phlD abundance at 346 the rhizosphere soil level, while the same treatments resulted in the lowest *phlD* abundance at the 347 bulk soil level. After the second trial, 2017, Pseudomonas - Pp treatment resulted in the highest *phlD* abundance at the rhizosphere soil level. At the bulk soil level, the highest abundance was 348 349 observed after Trichoderma W2 - TW2 treatment, while Pseudomonas - Pp and the green compost 350 - CV treatments resulted in a lower *phlD* gene abundance than the untreated control - C.

351 The abundance of the hcnAB gene was significantly influenced by treatments and it was 352 significantly higher in the rhizosphere than in the bulk soil samples for all the treatments and for the 353 untreated control - C (Table 6). After the first trial, 2016, the highest hcnAB gene abundance was observed after Pseudomonas - Pp, inoculated compost - CM and Trichoderma W2 - TW2 354 355 treatments at the rhizosphere level. At the soil level the hcnAB gene abundance was not significantly different compared with the untreated control -C, with the exception of *Pseudomonas* 356 357 - Pp and the green compost - CV treatments which resulted in the lowest *hcnAB* gene abundance. 358 After the second trial, 2017, the hcnAB gene abundance was highest after the inoculated compost -CM treatment and in the untreated control - C was noted the lowest hcnAB gene abundance at the 359 360 rhizosphere level. At the soil level the *hcnAB* gene abundance was higher in the untreated control C 361 and the Trichoderma W2 – TW2 treatment while the lowest hcnAB gene abundance was noted after *Pseudomonas* – Pp and the green compost – CV treatments. 362

364 3.2.4. BCAs - like microorganisms abundance

Overall, at the end of both trials (2016 and 2017 respectively) the BCA applications resulted in an 365 366 increase in the indigenous populations of Bacillus, Trichoderma and Pseudomonas in all the 367 treatments, compared to the untreated control - C (Table 7). No significant interaction treatment \times 368 year was observed. In general, the abundance of *Bacillus* sp. was significantly higher for the Bacillus - SM treatment than the untreated control - C and all the other treatments in both the 369 370 rhizosphere and bulk soil. However, the inoculated compost treatment - CM also resulted in an 371 increased resident Bacillus abundance in rhizosphere and bulk soils after both trials. Nevertheless, 372 the Trichoderma TW2 - TW2 treatment showed similar levels to those of the untreated control - C. 373 Significant interactions bulk \times rhizosphere were observed (Table 7).

The inoculation with the *Pseudomonas* - Pp significantly increased the overall *Pseudomonas* community in both the rhizosphere and bulk soils, compared to the untreated control - C and the other treatments. Significant interactions bulk \times rhizosphere were observed (Table 7).

377 The inoculation with the Trichoderma - based formulate significantly increased the overall 378 Trichoderma community in both the rhizosphere and bulk soils, compared to the untreated control -379 C. After both trials, the highest abundance of Trichoderma was observed in the rhizosphere after 380 Trichoderma TW2 - TW2 treatment and in the bulk soil after the green compost – CV treatment. 381 After the second trial, 2017, a highest Trichoderma abundance was also observed after the 382 inoculated compost - CM treatment at the bulk soil level. The Serenade max - SM - Bacillus application resulted in somewhat lower Trichoderma abundance than all the other BCA 383 384 applications. Significant interactions bulk \times rhizosphere were observed (Table 7).

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386

387 3.3. Chemical properties

Generally for both trials, 2016 and 2017 respectively, pH was significantly lower (p < 0.05) at the rhizosphere level than at the bulk soil level, with values ranging from between 5.70 and 6.43 for the rhizosphere and between 6.34 and 6.77 for the bulk soil samples. The concentrations of TN, NH₄⁺, NO₃⁻ and TOC were generally significantly higher in the rhizosphere than in the bulk soil samples. The untreated control - C was characterized by higher NH₄⁺, NO₃⁻ and TOC concentrations than all the treatments. On the other hand, the highest TP and AP contents were observed in the bulk soil samples (Table S1). No significant treatment × year interaction was found.

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396 3.4. Potential Nitrification Activity (PNA)

397 The PNA measurements generally exhibited significantly higher values at the bulk soil level than in 398 the rhizosphere level. No significant treatment \times year interaction was observed, therefore the data 399 were presented as the average of all data from both trials, 2016 and 2017 respectively. The PNA in 400 the bulk soil was significantly (p < 0.05) higher in the untreated control - C, followed by the 401 inoculated compost - CM and the Pseudomonas - Pp treatments. The lowest value was observed 402 after both, Serenade max - SM - Bacillus and Remedier - RM-Trichoderma spp treatments. The 403 PNA in the rhizosphere was significantly lower (p < 0.05) in the untreated control - C than in the 404 other treatments. (Figure 2).

405

3.5. Correlations between the levels of total microbial communities (16S bacteria, 16S archaea, 18S
fungi), the functional genes (fungal *chiA* gene, bacterial and archaeal *amoA* genes, bacterial *phID*and *hcnAB* genes), the *Bacillus, Pseudomonas, Trichoderma* population levels and FOL
abundances, DS and PNA.

In general the correlations were negative, with r coefficients ranging from -0.37 to -0.5 (p < 0.001) at the rhizosphere level and from -0.3 to -0.6 (p < 0.001) at the bulk soil level (Table 8). Very strong positive correlations were observed between FOL and DS, as well as between the nitrification rate - PNA and DS. Strong negative correlations were recorded at the rhizosphere and bulk soil levels especially between AOB and DS at the soil level, and between AOA and PNA at thesoil level (Table 8).

416

417 3.6. Correlations between *Bacillus, Pseudomonas, Trichoderma* population levels and chemical
418 properties in rhizosphere (rhizo) and soil samples

419 In general the correlations were significant only with respect to AP, TP, NO_3^- and TOC, with r 420 values ranging from -0.5 to 0.8 (Table 9).

421

422 3.7. Regressions between indigenous microbial communities and soil chemical properties

423 Variance in the total bacterial abundance was explained by alterations of NO₃⁻ and TP at the 424 rhizosphere level (Table S2). Changes in the total archaeal abundance were explained by alterations 425 of NO_3^{-1} and TP at the rhizosphere and soil level respectively. Changes in the total fungal 426 community were explained by alterations of TP and TOC at the rhizosphere level and TP at the soil 427 level. The variance in *chiA* gene abundance was explained by alteration of TN including NO₃⁻ and NH_4^+ at both rhizosphere and soil level. AOA abundance changes were explained by TN, NO_3^- and 428 429 TOC at the rhizosphere level and by NO_3^- at the soil level. However, a greater percentage of AOB 430 was explained also by NO_3^- (Table S2).

431

432 **4. Discussion**

In the present study, rhizosphere and soil microbial community changes have been investigated monitoring total bacterial, archaeal and fungal highly distinctive genes abundance as indicators of the general alterations that may take place as a result of plant treatments and the infection of lettuce by FOL. In this context, the potential impact of various BCA and compost treatments on the total indigenous microbial communities and on the functional genes that controls nitrogen availability was assayed.

439

440 4.1. Disease severity, FOL abundance and plant biomass

441 The obtained results were in line with previous studies showing that the application of BCAs and 442 certain organic matter can effectively suppress soil pathogens (Zhang et al., 2008; Luo et al., 2009; 443 Wu et al., 2009; Ling et al., 2010; Cao et al., 2011; Lang et al., 2011; Yang et al., 2011; Gilardi et 444 al., 2016). The application of different BCAs and compost significantly reduced wilting by 445 suppressing FOL abundance compared to the untreated control. This result was corroborated by the 446 significant positive correlations (p < 0.001) that were observed between disease severity and FOL 447 abundance. These findings were supported by previous studies that showed that Fusarium 448 abundance in the cucumber and banana rhizosphere was reduced after BCA treatments (Qiu et al., 449 2012; Shen et al., 2015; Fu et al., 2017). Furthermore, FOL abundance was very low after the 450 treatment with inoculated compost (CM), thus indicating that an effective disease control could be 451 achieved with organic amendments, as previously pointed out (Bonanomi et al., 2010). Similar 452 results were obtained in a previous study (Pugliese et al., 2011), confirming that inoculation of 453 compost with *Trichoderma* strains can improve the suppressive activity of the substrate. The 454 application of compost may have increased the competition for resources and the interactions 455 between antagonist species of compost and soil, and this may have resulted in a reduction in the 456 FOL abundance and in the potential activity of the pathogen (Bonilla et al., 2012; Larkin et al., 457 2015).

458

459 4.2. Ecological fitness of the BCA - like microorganisms and their effect on the total indigenous
460 prokaryotes

461 Our results have evidenced an excellent feedback of the BCA - *like microorganisms* after the BCAs 462 were introduced into the soil, as inoculated plantlets or inoculated compost. This was highlighted by 463 significant negative relationships (p < 0.001) between the BCA – like populations and Fusarium 464 wilt severity, which suggested not only the ecological fitness of inoculated microorganisms, but 465 also their efficiency against FOL. Most probably, the efficiency of biological control of FOL was 466 based on mechanisms as saprophytic competition for nutrients, parasitic competition for infection 467 sites and induced resistance in host cells (Haas and Defago, 2005; Nel et al., 2006; Srinivasan et al., 468 2009). Even more, the good relationships between the BCAs - like microorganisms and soil 469 nutrients highlighted the role of the antagonists in solubilising certain compounds (nitrogen, 470 phosphorus, potassium, iron) (Gyaneshwar et al., 2002; Richardson et al., 2009) contributing to the 471 disease suppression.

472 The application of BCA - based formulates resulted in a subsequent boost in abundance of the 473 respective microbial populations (i.e., *Pseudomonas*, *Trichoderma* and *Bacillus*). A remarkable 474 result of this study concerns the high abundance that was observed in the rhizosphere samples of the 475 phlD and hcnAB genes known to encode the production of antifungal compounds such as 2,4-476 diacetylphloroglucinol (DAPG) and hydrogen cyanide (HCN) (Dudenhoffer et al., 2016). We 477 speculated that Pseudomonas spp (e.g., P. fluorescens) abundance would have increased in the 478 rhizosphere environment upon pathogen infection, as these bacteria are strong root colonizer 479 (Looper et al., 2012). This may explain the lower observed abundance of the total *Pseudomonas* 480 community compared with other treatments. This was probably due to the action of antifungal 481 metabolites against FOL, but also against other microorganisms that generally colonize the 482 rhizosphere. The chiA gene abundance confirmed the potential antagonistic activity of the inoculated Trichoderma strain TW2, but also of the inoculated compost. In addition, a significant 483 484 relationship of *chiA* gene abundance with the total and especially with ammonium N (p < 0.001) 485 content was observed (Table S2). This result highlighted the involvement of fungal population in 486 remineralization of nutrients during the decomposition of chitin (Gould et al., 1981) with important 487 contribution to nitrification process.

488 Overall, we did not observe any negative side-effects of BCA application on the total microbial 489 communities, thus indicating that the microorganisms introduced into the soil system may have 490 favoured the resident microbial populations through positive interactions. Our results were in line 491 with previous reports (Ghini et al., 2000; Gullino et al., 1995; Mezzalama et al., 1998). Contrary to 492 our expectations, a higher total fungal abundance was observed in the rhizosphere than in the bulk 493 soil. The boost in the fungal community in all the treatments and in the untreated control was 494 probably supported by the elevated level of organic carbon (Table S1). This may be linked to a 495 competitive potential, based on an increased root exudation, which favours rhizosphere colonization 496 by fungi, including FOL. In this context, Strange (2005) showed that fungi mobilize nutrients 497 through the production of growth regulators. This may explain the observed positive correlation 498 between AP and the total fungal community in both the rhizosphere and bulk soil. On the other 499 hand, a lower archaeal population level was observed in the rhizosphere, probably as a result of the 500 apparently lower pH. Moreover, the archaeal community may also have been lower due to a lower 501 growth rate and competitiveness than the bacteria and fungi (Karlsson et al., 2012). In addition, in 502 the present study, the archaeal community was negatively correlated (p < 0.05) with the disease 503 severity, thus suggesting a likely competition for nutrients with FOL. Thus may underline the 504 possible ability of archaea to contribute to the suppression of lettuce Fusarium wilt development. In 505 a previous study, Mendez et al. (2013) described a relationship between archaeal community 506 composition and soil suppressiveness to Rhizoctonia damping-off. However, further investigations are needed to clarify whether archaea play an important role in the effective protection of lettuce 507 508 against FOL.

509

510 4.3. Ammonia oxidizing archaea and bacteria

511 The archaeal nitrifiers (AOA) were more abundant in the soil samples than their bacterial 512 counterparts. In fact, a lower AOA abundance was observed in the rhizosphere than in the bulk soil, 513 which may be a result of a high organic carbon content and a lower pH (Wessén et al., 2010; Bates 514 et al., 2011). In addition, a low ammonia environment represents the key factor that determines a 515 niche separation of AOA and AOB in acidic/neutral soils. A strong positive relationship was 516 observed between the AOB and TN at the rhizosphere level, and this resulted in a dominance of 517 bacterial nitrifiers and their resilience to lower pH compared to AOA (Wessén et al., 2010). The 518 result supports the finding of Valentine (2007), who showed that all archaea, and their nitrifiers 519 components in particular, in general have a higher competitive advantage under reduced organic 520 resource conditions. Accordingly, we observed a higher AOA abundance in the soil samples, which 521 indicated that the archaeal nitrifiers, and in general the entire archaeal community, were probably 522 adapted to the decomposition of recalcitrant organic matter, as described earlier on by Cucu et al., 523 (2017).

524 Overall, the effect of different applied treatments resulted in the increase of nitrifiers abundance 525 with respect to the untreated control thus suggesting that BCAs may not have negative ecological 526 impacts on other groups of microorganisms. Comparing with all treatments,, a higher AOB 527 abundance was observed in the bulk soil of the untreated and FOL infested control, while the 528 opposite was true for the AOA in the rhizosphere, thus suggesting that the sole presence of FOL 529 may has induced a differentiation, in the rhizosphere and bulk soil, among the indigenous 530 prokaryotes involved in the nitrification process. Hence, the AOB abundance was not reduced in the 531 control samples, probably as a result of a niche differentiation among nitrifiers at the rhizosphere 532 and bulk soil levels induced by the FOL presence, rather than the commonly acknowledged direct 533 resource competition with AOA (Nicole et al., 2008; Wessén et al., 2010; Musyoki et al., 2015). 534 Additionally, a strong negative relationship between the nitrification rate and disease severity was observed at the bulk soil level. However, competition for resources may exist between AOA 535 536 abundance and FOL. This hypothesis was corroborated by the low nitrification rate that was 537 observed at the rhizosphere level especially in the untreated control. Our findings indicated that 538 nitrification may play an important role in disease control when environmental conditions for 539 nitrifiers are favourable. Similarly, Fujiwara et al., (2013) showed that, in a multiple parallel 540 mineralization system, rhizosphere microbiota suppressed the Fusarium wilt disease of lettuce by 541 affecting the morphological characteristics of F. oxysporum. However, further research is needed to 542 obtain a better understanding of the promoting effect of nitrifiers after the application of BCAs.

543

544 **Conclusions and outlook**

545 Our results have shown that the application of different biological control agents and compost 546 resulted in a significant reduction in the Fusarium wilt of lettuce crops, without any significant 547 differences in the degree of disease control among treatments. On the basis of the presented results, 548 it is possible to conclude that the applied treatments (based on Bacillus, Pseudomonas and 549 Trichoderma) did not affect negatively the density of resident microbial communities (i.e., bacteria 550 archaea and fungi) or the nitrifying prokaryotes. Nevertheless, the archaeal nitrifiers had a more 551 dynamic response after the treatments. Hence, more investigations are needed to study the influence 552 of BCA and compost treatments on archaeal communities. In addition, our results have shown that lettuce plants could favour bacterial species which are good lettuce rhizosphere colonizers and may 553 554 act as antagonists.

555 Although no negative effects of BCAs were observed in this study on the no-target rhizosphere or 556 on the soil microbial biomass, their application requires further evaluations, including the 557 consideration of a broader range of soils with different physico-chemical properties, as well as 558 additional soil-borne pathogens and different plant species/cultivars, as they may have various 559 effects on the microbial communities and processes in the rhizosphere and bulk soil (Soderberg et 560 al., 2002; Rasche et al., 2006). In addition, the long-term effect on the same crop should be 561 evaluated in order to fully understand the legacy and the results of the interactions of BCAs and 562 compost with microbial biomass and activity. Although this study was focused on the potential 563 functionality of selected microbial populations in terms of abundance it can be assumed that the 564 introduction of different microbial strains or strain combinations could also influence the structure 565 and composition of the overall microbial communities. Therefore, advanced studies on the potential 566 effects of BCAs and organic matrices on the composition and biomass of non-target microbial 567 populations in the crop rhizosphere and bulk soil, using molecular fingerprinting techniques, are 568 still needed.

569

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575	Conflict of interest
576	The authors declare that they have no conflict of interest.
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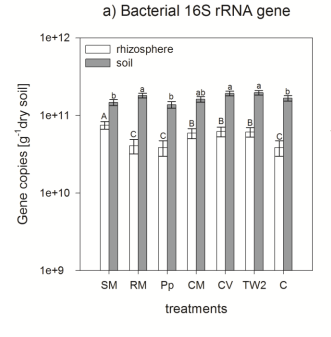
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Figure Captions

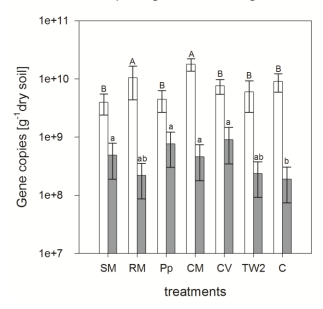
Figure 1 Abundance of bacterial (a), archaeal (b) 16S rRNA genes and fungal (c) 18S rRNA genes in rhizosphere and bulk soil between different treatments* and the untreated control (SM - Serenade Max; RM - Remedier; Pp - Pseudomonas; CM - Green compost plus Trichoderma TW2; CV green compost; TW2 - Trichoderma TW2; C - control) (n = 6, means±standard errors). Different letters above bars indicate significant differences between treatments in rhizosphere (uppercase letters) and bulk soil (lowercase letters) Figure 2 Potential Nitrification Activity (PNA) in the rhizosphere and bulk soil after different treatments* and the untreated control (SM - Serenade Max; RM - Remedier; Pp - Pseudomonas; CM - Green compost plus Trichoderma TW2; CV - green compost; TW2 - Trichoderma TW2; C -control) (n = 6, means \pm with standard errors). Different letters above bars indicate significant differences between treatments in rhizosphere (uppercase letters) and bulk soil (lowercase letters) *Serenade max – SM: Bacillus subtilis; Remedier – RM: Trichoderma asperellum + T. gamsii; Pseudomonas putida – Pp; ANT'S COMPOST M – CM: Green compost + Trichoderma TW2; ANT'S COMPOST V - CV: Green compost; Trichoderma sp. - TW2



1e+9 1e+8 1e+7 1e+7 1e+6 SM RM Pp CM CV TW2 C

treatments

c) Fungal 18S rRNA gene



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Figure1

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b) Archaeal 16S rRNA gene

6 soil rhizosphere ٢ A 5 В N - NO_2^{-} ug / dw⁻¹ В 4 Ç 3 ₽ a L a T 무 D 2 þ þ ¢ T С 1 đ 0 TW2 SM С RM Рр СМ CV treatments

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846 Figure2

Supplementary material

Table S1 Chemical properties of the rhizosphere (rhizo) and bulk soil for all the treatments and the untreated control at the end of trial 1 (2016) and trial 2 (2017). Values are given as average (n = 3). Different letters within a row show significant differences between treatments and the untreated control.

Property	Serena	Serenade Max Remedier Pseudomonas		omonas	Compost M		Comp	Compost V		Trichoderma		ntrol		
	Rhizo	Bulk soil	Rhizo	Bulk soil	Rhizo	Bulk soil	Rhizo	Bulk soil	Rhizo	Bulk soil	Rhizo	Bulk soil	Rhizo	Bulk so
2016														
рН	5.76±0.1 ^d	6.34±0.1 ^d	6.01±0.2 ^b	6.67±0.2 ^b	5.98±0.2 ^c	6.73±0.1 ^ª	6.23±0.1 ^ª	6.61 ± 0.1^{b}	5.92±0.2 ^c	6.59±0.2 ^b	6.03±0.2 ^b	6.6±0.1 ^b	5.80±0.1 ^d	6.53±0.1
TN %	0.57±0.2 ^d	0.21±0.05 ^ª	0.50±0.2 ^d	0.19 ± 0.08^{a}	0.27±0.3 ^f	0.21±0.04 ^ª	0.34±0.1 ^e	0.21±0.05 ^ª	0.83 ± 0.1^{b}	0.20±0.03 ^a	0.62±0.3 ^c	0.20±0.08 ^ª	1.02±0.1 ^ª	0.20±0.0
NH₄⁺(mg/kg)	2.9±0.9 ^d	0.23±0.09 ^b	1.6±0.6 ^e	0.39±0.2 ^a	2.07±0.8 ^c	0.27±0.1 ^b	1.13±0.5 ^f	0.29±0.1 ^b	5.29±0.1 ^b	0.33±0.1 ^b	0.6 ± 0.1^{g}	0.36±0.1 ^ª	37.55±1.7 ^ª	0.42±0.1
NO₃ (mg/kg)	18.06±1.2 ^g	5.91±1.9 ^f	22.65±1.8 ^e	8.51±1.6 ^e	39.28±1.9 ^b	22.86±1.7 ^ª	30.0±0.1 ^d	16.83±0.1 ^c	36.0±0.1 ^c	14.30±0.1 ^d	19.15±0.1 ^f	20.58±0.1 ^b	46.43±0.1 ^ª	8.62±0.1
TOC %	19.89±0.1 [°]	1.87±0.1 ^ª	13.82±0.1 ^d	1.90±0.1 ^ª	20.51±0.1 ^c	1.90±0.1 ^ª	7.21±0.1 ^e	1.96±0.1 ^ª	31.22±0.1 ^b	1.90±0.1 ^ª	19.34±0.1 [°]	1.89±0.1 ^ª	43.36±0.1 ^ª	1.71±0.1
TP (mg/kg)	906±1.5 ^f	1262.4±2.1 ^b	1020±1.2 ^d	1337.3±0.1.5 [°]	1296±1.2 ^b	1339.7±1.2 ^ª	1401±1.6 ^ª	1264.9±2.1 ^b	949±1.5 [°]	1254±2.1 ^b	1191±2.2 ^c	1238±2.5 ^b	679±1.5 ^g	1237±2.
AP (mg/kg)	124.4±0.2 ^b	150.2±2.7 ^c	118.5±0.5 ^c	152.7±2.3 ^b	129.3±0.7 ^ª	156.2±2.1 ^ª	125.9±0.1 ^b	158±2.31 ^ª	93.3±0.2 ^d	159.7±1.8 ^ª	113.5±0.1 ^c	156.6±2.6 ^ª	83.5±0.1 ^e	159.5±2.
2017														
рН	5.70±0.1 ^b	6.56±0.1 ^c	6.1±0.23 ^ª	6.88±0.1 ^ª	5.70±0.2 ^b	6.77±0.1 ^b	6.43±0.1 ^ª	6.76±0.1 ^b	5.85±0.1 ^b	6.74±0.2 ^b	5.98±0.2 ^b	6.9±0.2 ^ª	5.7±0.1 ^b	6.55±0.1
TN %	0.67±0.1 ^d	0.12±0.07 ^c	0.47±0.1 ^e	0.21±0.05 ^b	0.33±0.2 ^f	0.32±0.04 ^a	0.35±0.1 ^f	0.19±0.05 ^b	0.97±0.2 ^b	0.12±0.05 ^c	0.79±0.3 ^c	0.18 ± 0.05^{b}	1.20±0.2 ^a	0.18±0.0
$NH_4^+(mg/kg)$	2.59±0.5 [°]	0.25±0.1 ^d	2.6±0.17 ^e	0.41 ± 0.1^{b}	2.9±0.5 ^d	0.25±0.2 ^c	2.27±0.2 ^f	0.3±0.1 ^c	4.43±0.4 ^b	0.27±0.2 ^b	0.5±0.2 ^g	0.41 ± 0.1^{b}	33.76±1.2 ^ª	0.56±0.1
NO ₃ (mg/kg)	18.31±1.2 ^e	6.35±1.5 ^f	25.83±1.2 ^d	7.67±1.5 ^e	45.28±1.6 ^a	25.55±1.5 ^ª	36.0±1.2 ^c	19.91±1.2 [°]	39.0±1.3 ^b	12.47±1.2 ^d	19.21±1.2 ^e	22.47±1.4 ^b	51.29±1.2 ^c	5.37±1.2
TOC %	21.35±2.1 ^d	2.56±1.2 ^b	15.79±2.3 ^e	3.2±1.2 ^ª	22.34±2.1 ^d	2.2±1.2 ^c	13.33±2.2 ^f	2.85±2.4 ^a	27.46±2.2 [°]	2.3±2.3 ^c	32.34±2.3 ^b	1.89±1.2 ^d	56.36±1.3 ^ª	1.71±2.1
TP (mg/kg)	1023±1.5 ^d	1357±2.1 ^ª	980±1.2 ^d	975±2.1 ^d	1136±1.4 ^c	1238±1.2 ^b	1521±2.1 ^ª	1132.9±1.5 ^c	1178±2.1 ^c	1376±1.2 ^ª	1275±2.1 ^b	1231±1.2 ^b	53±2.1 ^e	1129±1-
AP (mg/kg)	112.9±0.1 ^c	176.1±2.1 ^b	131.4±0.5 ^b	164.6±2.2 ^b	131.7±0.5 ^ª	175.4±2.1 ^b	131.2±0.1 ^b	187±2.1 ^ª	112±0.2 ^c	198.5±2.1 ^ª	99.5±0.2 ^c	153±2.1 ^c	76±0.1 ^d	171±2.2
4														

Property	16S bacteria		16S archaea		18S fungi		chiA gene		AOA		AOB	
	rhizo	bulk soil	rhizo	bulk soil	rhizo	bulk soil	rhizo	bulk soil	rhizo	bulk soil	rhizo	bulk soil
рН	ns	ns	ns	ns	ns	ns	ns	ns	ns	ns	ns	ns
TN	ns	ns	ns	ns	ns	ns	0.38*	0.673*	0.77***	ns	0.876	ns
NH_4^+	ns	ns	ns	ns	ns	ns	0.754***	0.85***	ns	ns	ns	ns
NO ₃	0.656***	ns	0.778***	0.697**	ns	ns	0.68***	0.50**	0.79***	0.777***	0.65**	0.856***
Ptot	0.50**	ns	ns	ns	0.679***	0.95***	ns	ns	ns	ns	ns	ns
Pav	ns	ns	ns	ns	ns	ns	ns	ns	ns	ns	ns	ns
TOC	ns	ns	ns	ns	6.975***	ns	ns	ns	0.76***	ns	ns	ns

Table S2 - Linear regressions (R² correlation coefficients, n=21) between microbial abundance and chemical properties in rhizosphere (rhizo) and bulk soil samples

Significance levels: not significant-ns: p>0.05; *p<0.05; **p<0.01; ***p<0.001.

883 Tables

Table 1 – General information on the trials and timing of the operations carried out in 2016 (trial 1)

885 and 2017 (trial 2)

Treatment	Microorganism -	Dosage	Tray	Plot
	active ingredient	10	treatment	Treatment
Serenade max -	Bacillus subtilis QST 713	2.9X10 ¹⁰ cells/L water	T5; T10;	-
SM			T15;T20*	
Remedier - RM	Trichoderma asperellum + T	1.2X10 ⁶ cells/L water	T5; T10;	-
	gamsii		T15;T20	
Pseudomonas	FC7B+ FC8B +FC9B	1x10^7 (cells /ml	T5; T10;	
	FC/BT FC0B TFC9B			-
<i>putida</i> - Pp		water)	T15;T20	
ANT'S COMPOST	Green compost +	8g/seedling;1kg/0.2m ³	Т0	-
M - CM	Trichoderma TW2	of soil		T20
ANT'S COMPOST	Green compost	8 g/seedling;	т0	-
V - CV		1kg/0.1m ³ of soil		T20
Trichoderma sp	Trichoderma strain TW2	1x10^7 (cells /ml)	T5; T10;	
TW2			T15; T20	
Ortiva	Azoxystrobin	0.19 g a.i./Lwater		Т20
Untreated control				
- C	-	-		

886 *Treatments: T0 at sowing; T5: 5 days after sowing; T10: 10 days after sowing; T15: 15 days

after sowing; T 20: 20 days after sowing and immediately before transplanting.

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Target group	Primer (reference)	Amplification details
Fusarium	FLA0001F (Shimazu et al., 2005)	40 cycles
oxysporum f.sp.	FLA0001R (Shimazu et al., 2005)	95°C 15s, 60°C 60s, 72°C 45s
<i>lactucae</i> (FOL)		
All bacteria	Eub338 (Lane 1991)	40 cycles
(16S rRNA gene)	Eub518 (Muyzer et al. 1993)	95°C 30s, 55°C 35s, 72°C 45s
All archaea	340F (Gantner et al., 2011)	40 cycles
(16S rRNA gene)	1000R (Gantner et al., 2011)	95°C 30s, 57°C 30s, 72°C 30s
All fungi	FR1 (Vainio and Hantula, 2000)	45 cycles
(18S rRNA gene)	390FF (Vainio and Hantula, 2000)	95°C 30s, 50°C 30s, 70°C 60s
Ammonia	AmoA-1f (Rotthauwe et al. 1997)	45 cycles
oxidizing	AmoA-2r (Rotthauwe et al. 1997)	, 95°C 30s, 57°C 45s, 72°C 45s, 78°C 20s
bacteria (AOB)	, ,	, , ,
Ammonia	Arch-amoAf (Francis et al., 2005)	45cycles
oxidizing	Arch-amoAr (Francis et al.,2005)	, 95°C 30s, 53°C 45s, 72°C 45s, 78°C 20s
archaea (AOA)		
<i>chiA</i> gene	chiaxf (this study)	35cycles
-	chiaxr (this study)	95°C 15s, 59.9°C 30s, 70°C 30s
phID gene	BPF2 (McSpadden Gardeneret al.,	40 cycles
	2001)	95°C 15s, 60°C 45s, 72°C 45s, 78°C 20s
	BPR4 (McSpadden Gardeneret al.,	
	2001)	
hcnAB gene	PM2 (Svercel, Duffy & Défago 2007)	40 cycles
	PM-26R (Svercel, Duffy & Défago	95°C 15s, 60°C 45s, 72°C 45s, 78°C 20s
	2007)	
Bacillus	Forward <i>B. subtilis</i> (Gao et al., 2011)	40 cycles
	Reverse B. subtilis (Gao et al., 2011)	95°C 30s, 60°C 60s, 80°C 10s
Trichoderma	uTf (Hagn et al., 2007)	35 cycles
	uTr (Hagn et al., 2007)	95 °C 30 s, 55.5 °C 30 s, 72 °C 30 s
Pseudomonas	Pse435F (Bergmark et al., 2012)	40 cycles
	Pse686R (Bergmark et al., 2012)	95°C 30s, 60°C 60s, 80°C 10s

Table 2 Description of the primer sets and amplification details used for the quantitative PCR.

896 Table 3 Effect of the preventative soil treatments with BCAs and compost on disease severity and fresh weight at the end of trials 1 and 2, 2016 and

897 2017, respectively

	Microorganism	D	Disease	severi	ty*	Fresh weight g/12 plants					
Treatment	Microorganism - active ingredient			%							
	active ingredient	20	016	2	017	20	016	2017			
Serenade max - SM	Bacillus subtilis QST 713	26.9	±3.6 ^ª	22.1	±3.3 ^{ab}	3135.6	±128.9 ^a	2727.2	±371.5 ^{ab}		
Remedier - RM	Trichoderma asperellum +	27.7	±3.2 ^ª	29.5	±7.0 ^{ab}	2731.0	±79.0 ^a	2868.4	±348.7 ^{ab}		
	T. gamsii										
<i>Pseudomonas putida</i> - Pp	FC7B+ FC8B +FC9B	28.1	±2.9 ^a	35.2	±6.3 ^{ab}	3090.7	±49.6 ^ª	1528.0	±67.2 ^{bc}		
ANT'S COMPOST M - CM	Green compost +	25.3	±3.3 ^ª	13.0	±4.7 ^a	3256.3	±267.6 ^a	3766.0	±481.6 ^a		
	Trichoderma TW2										
ANT'S COMPOST V - CV	Green compost	32.0	$\pm 4.1^{a}$	29.0	±5.3 ^{ab}	3323.9	±146.7 ^ª	3747.6	±130.1 ^a		
Trichoderma sp TW2	Trichoderma strain TW2	32.2	±2.7 ^a	36.4	±3.3 ^b	2815.7	±78.6 ^ª	1917.6	±134.7 ^{bc}		
Ortiva	Azoxystrobin	24.8	$\pm 4.0^{a}$	19.8	±2.3 ^{ab}	3014.0	±185.3 ^ª	3360.4	±273.0 ^a		
Untreated control - C	-	61.0	±3.5 ^b	61.7	±4.1 ^c	1745.7	±93.0 ^b	770.8	±243.1 ^c		

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900 *Sixteen plants/treatment were evaluated for disease severity using the rating scale: 0 = healthy plant, 25 = initial leaf chlorosis, 50 = severe leaf chlorosis and initial symptoms of

901 wilting during the hottest hours of the day, 75 = severe wilting and severe symptoms of leaf chlorosis; 100 = plant totally wilted, leaves completely necrotic.

Table 4 Abundance of *F. oxysporum* f.sp. *lactucae* (FOL) in the rhizosphere and bulk soil after
different treatments* and in the untreated control at the end of trial 1 (2016) and trial 2 (2017)
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Treatment	Rhizospher	e 2016	Bulk soil	2016	Rhizospher	e 2017	Bulk soil	2017
	FOL (logcopy	DNA -1)						
Serenade Max - SM	6.359	а	6.711	с	6.345	b	6.796	b
Remedier - RM	5.626	С	6.758	с	5.692	С	6.828	b
<i>P. putida -</i> Pp	6.034	b	7.117	b	6.179	b	6.959	а
ANT'S COMPOST M - CM	5.423	d	6.086	d	5.243	d	6.119	с
ANT'S COMPOST V - CV	5.897	с	6.769	с	5.897	с	6.821	b
<i>Trichoderma</i> sp TW2	6.122	b	5.950	d	6.292	b	5.983	с
Untreated control - C	6.445	а	7.298	а	7.635	а	7.073	а
P(F) Treat	0.000							
P (F) Year	0.000							
P(F) Treat*Year	ns							
P(F) Rhizo*Bulk	0.001							

906 Values represent marginal means

907 *Serenade max – SM: Bacillus subtilis; Remedier – RM: Trichoderma asperellum + T. gamsii; Pseudomonas putida – Pp; ANT'S COMPOST M –

908 CM: Green compost + *Trichoderma* TW2; ANT'S COMPOST V – CV: Green compost; Trichoderma sp. – TW2

921	Table 5 Abundance of bacterial (AOB) and archaeal (AOA) amoA genes in the rhizosphere and
922	bulk soil after different treatments* and in the untreated control at the end of trial 1 (2016) and trial
923	2 (2017)

Treatment	Rhizospher	e 2016	Bulk soil	2016	Rhizospher	e 2017	Bulk soil 2	2017
	AOB (logcopy	DNA -1)						
Serenade Max - SM	7.417	а	7.374	С	7.493	а	7.383	С
Remedier - RM	7.139	с	7.653	b	7.192	с	7.741	b
<i>P. putida -</i> Pp	7.045	с	7.628	b	7.086	С	7.660	b
ANT'S COMPOST M - CM	7.236	b	7.680	b	7.320	b	7.715	b
ANT'S COMPOST V - CV	7.407	а	7.656	b	7.446	а	7.693	b
<i>Trichoderma</i> sp TW2	7.396	а	7.703	b	7.410	а	7.698	b
Untreated control - C	7.039	c	7.951	а	7.341	b	7.854	а
P(F) Treat	0.000							
P (F) Year	0.000							
P(F) Treat*Year	ns							
P(F) Rhizo*Bulk	0.007							
	AOA (logcopy	DNA -1)						
Serenade Max - SM	7.057	a	8.829	с	7.100	а	8.834	b
Remedier - RM	6.653	b	8.744	с	6.826	b	8.830	b
<i>P. putida -</i> Pp	6.415	с	8.958	b	6.653	с	8.985	b
ANT'S COMPOST M - CM	6.613	b	8.823	С	6.690	с	8.828	b
ANT'S COMPOST V - CV	6.721	b	9.163	а	6.800	b	9.224	а
Trichoderma sp TW2	6.974	а	9.042	b	6.844	b	9.123	а
Untreated control - C	5.799	d	8.118	d	5.839	d	8.422	с
P(F) Treat	0.000							
P (F) Year	0.000							
P(F) Treat*Year	0.015							
P(F) Rhizo*Bulk	0.006							

926 Values represent marginal means

927 *Serenade max – SM: Bacillus subtilis; Remedier – RM : Trichoderma asperellum + T. gamsii; Pseudomonas putida – Pp; ANT'S COMPOST M –

928 CM: Green compost + Trichoderma TW2; ANT'S COMPOST V – CV: Green compost; Trichoderma sp. – TW2

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Table 6 Abundance of *chiaA*, *phlD*, and *hcnAB* genes in the rhizosphere and bulk soil after different

treatments* and the untreated control at the end of trial 1 (2016) and trial 2 (2017)

Treatment	Rhizosphere	2016	Bulk soil	2016	Rhizospher	e 2017	Bulk soil	2017
	chiA gene (log	copy DN	A ⁻¹)					
Serenade Max - SM	4.528	с	4.406	b	4.572	d	4.539	b
Remedier - RM	4.761	ab	4.632	а	4.815	с	4.513	b
<i>P. putida -</i> Pp	4.317	d	4.612	a	4.482	d	4.635	a
ANT'S COMPOST M	_	u	1.012	u	1.102	u	1.055	u
CM	4.905	b	4.337	b	5.080	b	4.358	С
ANT'S COMPOST V	_							
CV	4.292	d	4.566	а	4.334	e	4.621	а
Trichoderma sp	-							
TW2	5.676	а	4.433	b	5.711	а	4.520	b
Untreated control - C	4.327	d	4.385	b	3.915	f	4.284	d
P(F) Treat	0.000							
P (F) Year	0.000							
P(F) Treat*Year	ns							
P(F) Rhizo*Bulk	ns							
	phID gene (log	conv DN	(A^{-1})					
Serenade Max - SM	4.952	c	2.627	а	4.842	с	2.613	b
Remedier - RM	4.864	c	2.662	a	4.726	c	2.642	b
<i>P. putida -</i> Pp	5.371	a	2.166	b	5.554	a	2.198	c
ANT'S COMPOST M		u		b		a		
CM	4.814	с	2.756	а	4.852	с	2.663	b
ANT'S COMPOST V	-							
CV	5.119	а	2.280	b	5.159	b	2.249	с
Trichoderma sp		,	0.055		5.0.00	,	0.040	
TW2	5.248	b	2.957	а	5.260	b	2.942	а
Untreated control - C	4.950	с	2.786	а	3.959	d	2.621	b
ond eated control - C	4.950	L	2.780	d	3.939	u	2.021	D
P(F) Treat	ns							
P (F) Year	0.000							
P(F) Treat*Year	ns							
P(F) Rhizo*Bulk	0.013							
	hcnAB gene (la	ogcopy D	NA -1)					
Serenade Max - SM	6.319	b	2.613	а	6.482	b	2.556	b
Remedier - RM	3.734	d	2.642	а	3.754	e	2.670	b
<i>P. putida -</i> Pp	6.707	a	2.198	b	6.555	b	2.155	с
ANT'S COMPOST M ·								
СМ	6.910	а	2.663	а	6.940	а	2.643	b
ANT'S COMPOST V	- F (01	4	2.240	h	E COC		2 2 2 7	
CV	5.601	d	2.249	b	5.636	с	2.227	с
Trichoderma sp. ·	- 6.010		2042		6 201	ь	2 002	
TW2	6.810	а	2.942	а	6.391	b	2.892	а
Untreated control - C	4.950	с	2.791	а	4.864	d	2.811	а
- (-) -			I		ı	1		
P(F) Treat	ns							
P (F) Year	0.000							
P(F) Treat*Year	ns							
P(F) Rhizo*Bulk	0.016							

936

937

938 Values represent marginal means

939

940 *Serenade max – SM: Bacillus subtilis; Remedier – RM: Trichoderma asperellum + T. gamsii; Pseudomonas putida – Pp; ANT'S COMPOST M –

941 CM: Green compost + *Trichoderma* TW2; ANT'S COMPOST V – CV: Green compost; Trichoderma sp. – TW2

Table 7 Abundance of *Bacillus*, *Pseudomonas*, and *Trichoderma* genes in the rhizosphere and bulk
soil after different treatments* and the untreated control at the end of trial 1 (2016) and trial 2
(2017)

Treatment	Rhizospher	e 2016	Bulk soil	2016	Rhizospher	e 2017	Bulk soil	2017
	Bacillus (logo	opy DNA	⁻¹)					
Serenade Max - SM	5.039	а	5.394	а	5.087	а	5.477	а
Remedier - RM	4.249	с	4.109	d	4.194	b	4.130	d
P. putida - Pp	3.989	С	4.320	с	3.980	с	4.331	С
ANT'S COMPOST M	-							
СМ	4.702	b	4.790	b	4.308	b	4.814	b
ANT'S COMPOST V	-							
CV	3.982	с	4.428	С	3.990	с	4.228	С
Trichoderma sp.	-							
TW2	3.654	d	3.908	d	3.641	d	3.898	e
Untreated control - C	3.615	d	3.871	d	3.617	d	3.879	е
P(F) Treat	0.000							
P (F) Year	0.000							
P(F) Treat*Year	ns							
P(F) Rhizo*Bulk	0.000							
	Pseudomonas	s (logcopy	, DNA -1)					
Serenade Max - SM	4.295	b	3.928	с	4.337	b	3.902	с
Remedier - RM	4.249	b	4.109	с	4.194	b	4.148	с
P. putida - Pp	4.599	а	4.506	a	4.921	a	4.781	a
ANT'S COMPOST M	-							
СМ	4.781	а	4.301	b	4.837	а	4.515	b
ANT'S COMPOST V	-							
CV	4.292	b	3.566	d	3.880	с	4.554	b
Trichoderma sp.								
TW2	3.654	с	3.908	С	3.735	с	3.681	d
Untreated control - C	3.495	с	3.234	е	3.435	d	3.252	е
			I		1	I		
P(F) Treat	0.000							
P (F) Year	0.000							
P(F) Treat*Year	ns							
P(F) Rhizo*Bulk	0.000							
	Trichoderma	Поасору	DNA^{-1})					
Serenade Max - SM	3.988	d	2.928	с	3.936	d	2.902	d
Remedier - RM	4.249	c	3.109	bc	4.345	d	3.252	c
<i>P. putida</i> - Pp	3.989	cd	3.320	b	3.992	d d	3.507	b
ANT'S COMPOST M		ιu	5.520	D	3.352	u	5.507	U
CM	4.905	b	3.337	b	5.001	b	4.621	а
ANT'S COMPOST V	-				1			a
CV	4.292	с	3.566	а	4.775	с	4.668	а
Trichoderma sp.	-				1			u
TW2	5.676	а	3.433	b	5.658	а	4.418	b
								~
Untreated control - C	3.327	e	2.384	d	4.207	d	3.293	d
D(E) Treat	0.000							
P(F) Treat	0.000							
P (F) Year	0.000							
P(F) Treat*Year	ns							
P(F) Rhizo*Bulk	0.000							

946

947 Values represent marginal means

948 *Serenade max – SM: Bacillus subtilis; Remedier – RM: Trichoderma asperellum + T. gamsii; Pseudomonas putida – Pp; ANT'S COMPOST M –

949 CM: Green compost + Trichoderma TW2; ANT'S COMPOST V – CV: Green compost; Trichoderma sp. – TW2

Table 8 Pearson's correlation coefficient between microbial gene abundance, disease severity (DS)
and potential nitrification activity (PNA) assesses in rhizosphere (rhizo) and bulk soil samples

55					
	Microbial	DS		PNA	
56	abundance				
		rhizo	bulk soil	rhizo	bulk soil
57	FOL	0.73***	0.57***	ns	-0.7***
58	16S Bacteria	ns	0.23		
	16S Archaea	-0.35	-0.24		
i9	18S Fungi	ns	-0.39**		
	AOB	-0.45**	0.75***	-0.31	-0.5***
50	AOA	-0.67***	-0.65***	-0.5***	0.87***
	Bacillus	-0.38**	-0.38*		
51	Pseudomonas	-0.44***	-0.35*		
62	Trichoderma	ns	-0.65***		
	<i>chi</i> A gene	ns	-0.46**		
63	phID gene	-0.7***	-0.5**		
13	hcnAB gene	-0.67***	-0.76**		
54	PNA	0.253	-0.6***		

966 Significance levels: not significant-ns: p>0.05; *p<0.05; **p<0.01; ***p<0.001

- /0/

Table 9 Pearson's correlation coefficient between, *Bacillus, Pseudomonas, Trichoderma*, FOL
population gene abundance and chemical properties assessed in rhizosphere (rhizo) and bulk soil
samples

Property	Bacillus		Pseudomonas		Trichoderma		FOL	
	rhizo	bulk soil	rhizo	bulk soil	rhizo	bulk soil	rhizo	bulk soil
рН	ns	ns	ns	ns	ns	ns	ns	ns
TN	ns	ns	ns	ns	ns	ns	ns	ns
${\sf NH_4}^+$	ns	ns	ns	ns	ns	ns	ns	ns
NO ₃	-0.5	ns	ns	0.60**	-0.57**	ns	ns	ns
PT	-0.80***	-0.50**	0.60***	ns	0.6**	ns	-0.5**	-0.76**
AP	0.50**	-0.60**	0.75***	0.40*	0.55**	0.60**	0.565**	0.438*
TOC	-0.6**	0.4 *	-0.70***	0.60**	-0.75***	0.76***	0.657***	0.543**

980 Significance levels: not significant-ns: p>0.05; *p<0.05; **p<0.01; ***p<0.001