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1	NITROGEN FERTILISERS SHAPE THE COMPOSITION AND PREDICTED FUNCTIONS
2	OF THE MICROBIOTA OF FIELD-GROWN TOMATO PLANTS
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22 ABSTRACT

The microbial communities thriving at the root-soil interface have the potential to improve 23 plant growth and sustainable crop production. Yet, how agricultural practices, such as the 24 25 application of either mineral or organic nitrogen fertilisers, impact on the composition and functions of these communities remains to be fully elucidated. By deploying a two-pronged 26 16S rRNA gene sequencing and predictive metagenomics approach we demonstrated that 27 the bacterial microbiota of field-grown tomato (Solanum lycopersicum) plants is the product 28 of a selective process that progressively differentiates between rhizosphere and root 29 microhabitats. This process initiates as early as plants are in a nursery stage and it is then 30 more marked at late developmental stages, in particular at harvest. This selection acts on 31 both the bacterial relative abundances and phylogenetic assignments, with a bias for the 32 enrichment of members of the phylum Actinobacteria in the root compartment. Digestate-33 based and mineral-based nitrogen fertilisers trigger a distinct bacterial enrichment in both 34 rhizosphere and root microhabitats. This compositional diversification mirrors a predicted 35 functional diversification of the root-inhabiting communities, manifested predominantly by 36 the differential enrichment of genes associated to ABC transporters and the two-component 37 system. Together, our data suggest that the microbiota thriving at the tomato root-soil 38 interface is modulated by and in responses to the type of nitrogen fertiliser applied to the 39 field. 40

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Additional keywords: *Solanum lycopersicum*, Rhizosphere, Root, Microbiota, Nitrogen,
 Fertilisers, Digestate.

Page 3 of 43

44 INTRODUCTION

Limiting the negative impact of agricultural practices on the environment while preserving sustainable crop yield is one of the key challenges facing agriculture in the years to come.

As an essential element for plant nutrition, nitrogen represents a paradigmatic example of 47 such a challenge. Moreover, due to the combined effect of elevated solubility and little 48 retention in soils, the lack of this element is and will be one of the major yield-limiting factors 49 worldwide (Tilman et al., 2011). At the same time, the application of synthetic nitrogen 50 fertilisers is, in many agricultural systems, a low efficiency approach which has been linked 51 with the degradation of natural resources (Elser and Bennett, 2011). One of the strategies 52 adopted to limit the economic and environmental footprint of crop production while 53 maintaining sustainable yield is the "recycling" of, mineral-rich, biodegradable products of 54 the livestock and agricultural sectors. 55

One example of this approach is the application of the digestate, a by-product of the 56 anaerobic digestion of organic waste for the production of biogas (Möller and Müller, 2012) 57 as renewable soil amendment for crop production. The digestate is a mixture of partially-58 degraded organic matter, microbial biomass and inorganic compounds (Alburguergue et al., 59 2012). We recently demonstrated how the digestate can be efficiently used as innovative 60 fertiliser and plant growing media (Ronga et al., 2018b; Ronga et al., 2018a; Ronga et al., 61 2019), yet the impact of digestate applications on the agroecosystem remains to be fully 62 elucidated. 63

For instance, the digestate can be a source of phytoavailable nitrogen, in particular ammonium, capable of impacting on organic matter mineralisation and emission of carbon dioxide from the soil profile (Grigatti et al., 2011). Therefore, it is legitimate to hypothesize that such treatments impact on the composition of the microbial communities thriving at the root-soil interface, collectively referred to as the rhizosphere and root microbiota, which play

a critical role in mobilisation of mineral elements for plant uptake (Alegria Terrazas et al.,
2016). Congruently, several studies indicate that the application of biogas by-product
enhances soil microbial activity (Möller, 2015) and the availability of phytohormones (Scaglia
et al., 2015). However, the intertwined relationship among microbiota composition, soil
characteristics and amendments as well as host plant species-specificity (Bulgarelli et al.,
2013) makes it difficult to infer first principles.

This is particularly true for field-grown crops such as tomato (Solanum lycopersicum L.), one 75 of the most cultivated horticultural crops globally with plantations occupying an area of 4.8 76 million of hectares with a production of 182 million tonnes in 2017 (FAO 2017). Notably, this 77 species is also an excellent experimental model for basic science investigations: tomato was 78 one of the first crops whose genome was sequenced (Consortium, 2012) and provided a 79 superb platform to test the significance of genome editing for evolutionary studies and plant 80 breeding (Zsögön et al., 2018). Perhaps not surprisingly, tomato is gaining momentum as 81 an experimental system to study host-microbiota interactions in crop plants. Recent 82 investigations revealed novel insights into the assembly cues of the microbiota associated 83 to this plant (Bergna et al., 2018; Toju et al., 2019) and the contribution of microbes thriving 84 85 at the tomato root-soil interface to pathogen protection (Chialva et al., 2018; Kwak et al., 2018). However, the composition and functional potential of the tomato microbiota and their 86 interdependency from nitrogen fertilisers remain to be elucidated. 87

Here we report the metagenomics characterisation of the microbiota thriving at the root-soil interface of field-grown tomato plants. We hypothesize that nitrogen treatments shape and modulate the contribution of the tomato microbiota for crop yield. To test this hypothesis, we focused on processing tomato exposed to different nitrogen fertilisers, either digestatebased or containing a mineral fraction. By using a 16S rRNA amplicon sequencing survey we deciphered how the microhabitat (i.e., either rhizosphere or root) sculpts the tomato microbiota which, in turn, is fine-tuned by the type of fertiliser applied. Finally, by using a
predictive metagenomics survey, we inferred the functional diversification imposed by the
nature of the fertilisers on the root microbiota.

97

99 MATERIALS AND METHODS

100 Field site

A field trial was established in a tomato farm near the city of Ravenna (44°25'40.8"N 101 12°05'53.3"E), Emilia Romagna Region, Italy, during the 2017 growing season. During the 102 period from transplant to harvest, the minimum and the maximum average temperatures 103 recorded were 17.1°C and 32.8°C, respectively, and the rainfall was 101.7 mm. The soil had 104 a silty loam texture (14% clay, 51% silt, 35% sand), a pH 8.3 (in H₂O), 1.1 g kg⁻¹ total N 105 (Kjeldahl method), 7 mg kg⁻¹ available P (Olsen method), 129 mg kg⁻¹ exchangeable K 106 (Ammonium acetate), and 9 g kg⁻¹ organic matter (Walkey-Black method). A schematic 107 illustration of the field trial is depicted in Supplementary Figure S1. 108

109 Plant material

We used the tomato cultivar 'Fokker', a processing-type genotype with blocky fruit, late fruit ripening and suitable for tomato puree, for the experimentation. Seedlings were provided by Bronte Soc. Coop. Agr. A.R.L. (Mira, Italy). Processing tomato seedlings were transplanted at the end of May when they were 6-week old corresponding to plants at the fourth true leaf stage. Plant density was 3 plants m⁻². Plants were transplanted into single row, with a spacing of 0.22 m between plants in each row and 1.50 m between rows.

116 **Experimental design**

We established a randomized complete design with three replicates and seven treatments: 117 pelleted digestate (hereafter PE), liquid digestate (LD), slow-acting liquid digestate (SRLD), 118 organo-mineral fertiliser based on digestate (SC), synthetic fertiliser (MF), slow-acting 119 synthetic fertiliser (SRMF), and no fertilization treatment (NT). The composition of the 120 treatments is summarised in Table 1. For each treatment, we applied a total amount of 121 nitrogen in the ratio 150 N kg ha⁻¹ on the basis of soil analysis, crop rotation and crop 122 nutrients required. Nitrogen was supplied at transplanting time with the exception of the 123 synthetic fertiliser treatment. For this latter treatment, the amount of total Nitrogen was 124

equally divided and applied in 3 times (transplanting, full flowering and fruit ripening) using 125 ammonium nitrate in the first treatment and calcium nitrate in the second and in the third 126 ones. During the trial, 600 m³ ha⁻¹ of irrigation water was distributed by drip irrigation to each 127 treatment. The other soil and crop management practices were performed according to the 128 production rules of Emilia Romagna Region, Italy. Briefly, weeds control was performed with 129 a single treatment (on 11th June) using products based on metribuzin and propaguizafop. 130 Sulphur and Copper were used to control phytopatogenic fungi while imidacloprid, 131 abamectin and spinosad were used as insecticide. 132

133 Yield traits

At harvest we determined the marketable yield (t ha⁻¹), as a weight of fully ripe fruits, and the solid soluble content (°Brix t ha⁻¹) as a proxy for fruit quality. The °Brix parameter was determined using the digital refractometer HI 96814 (Hanna, Italy), while the °Brix t ha⁻¹ was calculated by multiplying the hectare marketable yield by °Brix and dividing the result by 100.

139 Root, Rhizosphere and Bulk soil Sampling and DNA Extraction

At transplanting time (May 2017), 5 root specimens per treatment were collected. Upon 140 uprooting, soil particles loosely bound to roots were dislodged by hand shaking and root 141 segments of ~ 6 cm were placed in sterile 50 mL tubes. The samples were stored in a 142 143 portable cooler (~ 4°C), transported to the laboratory and immediately processed. Root specimens were incubated in 30 mL of PBS (Phosphate buffered saline) and placed on a 144 shaker for 20 minutes in order to separate the soil tightly adhering to plant material, which 145 we operationally defined as "rhizosphere", from the roots. The first tubes were centrifuged 146 for 20 minutes at 4,000 x g and the rhizosphere pelleted was collected in liquid nitrogen and 147 stored at -80°C. The roots were moved to a new sterile tube containing 30 mL PBS and 148 sonicated by Ultrasonics Sonomatic Cleaner (Langford Ultrasonics, Birmingham, UK) for 10 149 minutes (intervals of 30 seconds pulse and 30 seconds pause) at 150 W, as previously 150

reported (Schlaeppi et al., 2014) to enrich for endophytic microorganisms. Roots were then 151 washed in the same new buffer and dried on sterile filter paper. After few minutes, the roots 152 were moved to 50 mL tubes and frozen in liquid nitrogen for storage at -80°C. Three 153 independent soil samples were harvested from unplanted soil in different points of the field, 154 frozen in liquid nitrogen and stored at -80°C. At harvest time (September 2017) the whole 155 plants were harvested, 5 roots per treatment and 3 bulk soil samples were collected, 156 prepared and stored like the previous samples. Frozen root samples were pulverized in a 157 sterile mortar using liquid nitrogen prior DNA preparation. DNA was extracted from all the 158 specimens (i.e., bulk soil, rhizosphere and pulverized roots) using the FastDNA® SPIN Kit 159 for Soil (MP Biomedicals, Solon, USA) following the instruction manual provided by 160 manufacturer. DNA samples were diluted using 50 µL DES water and guantified using the 161 Nanodrop 1000 Spectrophotometer (Thermo Scientific, Wilmington, United States). 162

163 16S rRNA Gene Sequencing

The sequencing library generated using primers specific (515F 5'was 164 GTGCCAGCMGCCGCGGTAA-3' and 806R 5'-GGACTACHVGGGTWTCTAAT-3') for 165 hypervariable V4 region of the 16S rRNA gene. The reverse primers included a 12-mer 166 unique "barcode" sequences (Caporaso et al., 2012) to facilitate the multiplexing of the 167 samples into a unique sequencing run. Individual PCR reactions were performed as 168 previously reported (Robertson-Albertyn et al., 2017), with the exception of the concentration 169 of the Bovine Serum Albumin, added at 10 µg/reaction, and the addition of a Peptide Nucleic 170 Acid (PNA) blocker (PNA Bio, Newbury Park, United States) at a concentration of 0.5 171 µM/reaction to inhibit plastidial amplification. For each barcoded primers, three technical 172 replicates and a no-template control (NTC) were organised and processed starting from a 173 174 unique master mix. Five microliters of amplified samples and cognate NTCs were inspected on a 1% (w/v) agarose gel. Two independent sets of triplicated amplicons, displaying the 175 expected amplicon size and lacking detectable contaminations, were combined in a 176

barcode-wise manner and purified using the Agencourt AMPure XP kit (Beckman Coulter,
Brea, United States) with a ratio of 0.7 mL AMPure XP beads per 1 mL of sample. Purified
DNA samples were quantified using Picogreen (Thermo Fisher, United Kingdom) and
combined in an equimolar ratio into an amplicon pool. This latter material was used for the
preparation of a MiSeq run at the Genome Technology facilities of the James Hutton Institute
(Invergowrie, UK) as previously reported (Robertson-Albertyn et al., 2017).

183

184 OTU Table Generation and pre-processing

We used QIIME, version 1.9.0 (Caporaso et al., 2010) to process the sequencing output of 185 the MiSeg machine. Briefly, the command join paired ends.py was used to decompress 186 and merged (minimum overlap 5bp) forward and reverse read FASTQ files. Next, we 187 removed in silico low-quality sequencing reads and sequencing reads without the barcode 188 information. Then, the reads were assigned to individual samples. In these analyses, the 189 190 command split libraries fastg.py was used imposing a minimum PHRED score of 20. The resulted high-quality reads were assembled into an Operational Taxonomic Unit (OTU) table 191 at 97% sequence identity. We used a 'closed reference' approach for OTU-picking using the 192 command pick closed reference otus.py. We imposed the Green Genes database version 193 13 5 (DeSantis et al., 2006) as a reference database to identify microbial OTUs and prune 194 195 for chimeric sequences. We used SortMeRNA algorithm for OTU -picking and taxonomy assignment. Finally, OTUs whose representative sequences were classified as either 196 chloroplast or mitochondria, as well as OTUs accruing only one sequencing read over the 197 function entire dataset (i.e., singletons), were depleted in silico using the 198 filter otus from otu table.py. 199

200

201 Data visualisation and statistical analyses

Agronomic traits were analysed by Analysis of Variance (ANOVA) using GenStat 17th (VSN International, Hemel Hempstead, UK). Means were compared using Bonferroni's test at the 5% level.

The OTU table produced in QIIME was analysed in R using a custom script developed from Phyloseq package (McMurdie and Holmes, 2013).

Initially, the data were filtered removing the samples with less than 1,000 reads and the 207 OTUs with less than 10 reads in at least 5% of the samples. For alpha-diversity calculation, 208 sequencing reads were rarefied at an even sequencing deep of 18,467 reads per sample 209 retaining 2,439 unique OTUs. The number of Observed OTUs and Chao1 index were used 210 as richness estimators, while the Shannon index was used for evaluating the evenness. 211 Upon inspecting distribution of the data using a Shapiro-Wilk test, the means of rhizosphere 212 and root samples at harvest time were compared using a non-parametric Wilcoxon rank sum 213 test. Next, we performed a non-parametric Kruskal-Wallis test independently on 214 rhizosphere and root samples to identify significant effect of the individual treatments on the 215 ecological indices. 216

For beta-diversity calculation, the original counts (i.e., not rarefied) were transformed to 217 relative abundances and we imposed an abundance threshold to target PCR-reproducible 218 OTUs. The differences among microbial communities of the samples were computed using 219 Bray-Curtis index and weighted Unifrac index, with this latter index including phylogenetic 220 information in the analysis (Lozupone and Knight, 2005). A Principal Coordinates Analysis 221 (PCoA) was generated to visualize similarities and dissimilarities of microhabitats and 222 223 treatments. In order to assess the effects of microhabitats and the treatments on the bacterial community composition, a Permutational Multivariate Analysis Of Variance 224 (PERMANOVA) on distance matrices was implemented using the function Adonis in p a 225

two-pronged approach. First, we assessed the effect of nursery/harvest stage on microhabitat composition. Next, we used the same test to assess the impact of the treatment on rhizosphere and root specimens at harvest stage. In the two approaches, the computed R^2 therefore reflects the proportion of variance explained by the given factor in the group of samples tested.

Finally, original counts were used to perform a differential analysis to identify individual bacteria differentially enriched in the tested samples using DESeq2 (Love et al., 2014).

The phylogenetic tree was constructed using the representative sequences of the OTUs significantly enriched in rhizosphere and root specimens and annotated with iTOL (Letunic and Bork, 2006).

236 Functional predictions

Tax4Fun (Asshauer et al., 2015) package in R was used as a predictive tool to obtain a 237 functional profile based on 16S rRNA gene data. Metabolic capabilities are calculated by 238 linking the amplicon data phylogenetic and abundance profile to a set of pre-computed 239 metabolic reference profiles, based on the KEGG Ortholog (KO) database (Kanehisa et al., 240 2008). The input for this analysis was an OTU table obtained with the representative 241 sequences of the OTU table previously generated (see above), reclassified using 242 SILVA 115 taxonomic database (Quast et al., 2013). Similar to a previously reported 243 operational protocol (Kavamura et al., 2018), we focused our analysis in prokaryotic 244 functional categories related to amino acid metabolism, carbohydrate metabolism, cell 245 motility, energy metabolism, membrane transport, metabolism of terpenoids and polyketides 246 and two-component system, trimming the rest of predicted functions from the Tax4fun 247 248 output. A statistical comparison between two groups using a Welch's t-test (Bluman, 2009) filtered at a p-value < 0.01 with Storey's correction for false discovery rate (Storey and 249

250 Tibshirani, 2003) was performed in STAMP, Statistical Analysis of Metagenomic Profiles

251 (Parks et al., 2014).

252 Data and scripts availability

The 16S rRNA gene sequences presented in this study are available at the European Nucleotide archive under the study accession number PRJEB32219. The scripts to reproduce the statistical analysis and figures are available at <u>https://github.com/BulgarelliD-</u> <u>Lab/Tomato_nitrogen</u>. Data frames required for scripts reproducibility are included in Supplementary Database 1.

258 **RESULTS**

259 Fertiliser treatment impacts on yield and guality of processing tomato

At harvest time the two most important parameters such as marketable yield and fruit quality were measured to evaluate the effect of 7 different fertiliser performances on processing tomato (Figure 1). The fertiliser treatments had a significant effect on fresh biomass of fruits (ANOVA, Bonferroni's test, P < 0.001). Pelleted digestate registered the best performance followed by synthetic fertiliser and slow acting liquid digestate. In addition, the different fertilisers influenced significantly also the quality of processing tomato (Figure 1) (ANOVA, Bonferroni's test, P < 0.001).

267 The assembly dynamics of the bacterial microbiota of field-grown processing tomato

To gain insights into the relationships between yield traits and microbiota composition in

field-grown processing tomato plants, we generated 5,546,303 high quality 16S rRNA gene

sequences for the 86 samples generated in this study.

Upon *in silico* depletion of OTUs classified as Mitochondria and Chloroplast we reduced the number of analysable sequences to 4,645,503 with a retaining proportion of 83.7% of the original sequences (mean per samples = 54,017.48 reads; max = 111,213 reads; min = 272 reads). The data were further filtered removing the samples with less than 1,000 reads as Page 13 of 43

well as the OTUs with less than 10 reads in 5% of samples. This allowed us to retain 2,515
unique OTUs accounting for 4,308,580 high quality reads and 85 samples.

Then, we computed alpha-diversity calculations on a dataset rarefied at 18,467 reads per 277 sample and alpha-diversity was investigated considering two microhabitats (root and 278 rhizosphere) and the seven fertiliser treatments. OTUs richness was assessed by Chao1 279 index and Observed OTUs while the OTUs evenness was assessed by Shannon index. This 280 analysis revealed a significant effect of the microhabitat on the characteristics of the 281 microbiota thriving at the tomato root-soil interface: regardless of the treatment, the root 282 microhabitat emerged as less diverse and even compared to the rhizosphere one (Wilcoxon 283 rank sum, p <0.01, Figure 2). This observation suggests that root microhabitat represents a 284 gated community compared to the surrounding soil environment. Conversely, the treatment 285 impacted only the number of OTUs observed in the rhizosphere compartment (Kruskal-286 Wallis non parametric analysis of variance followed by Dunn's post-hoc test p < 0.05. Figure 287 288 2).

Congruently, beta-diversity analysis computed on the non-rarefied dataset using both 289 weighted Unifrac and Bray-Curtis indicated a microhabitat-dependent microbiota 290 diversification. In particular, the weighted Unifrac matrix visualised using a Principal 291 Coordinates Analysis revealed such a microhabitat effect on samples processed at harvest 292 time along the axis accounting for the major variation. Interestingly, younger nursery 293 samples displayed a similar degree of diversification, although their communities were 294 separated from the harvest samples on the axis accounting for the second source of 295 variation (Figure 3). These data were supported by a PERMANOVA which attributed a R² 296 of 30% to the microhabitat, a R² of 28% to the 'Nursery/Harvest effect' and a R² of 2% to 297 their interactions (Adonis test, 5,000 permutations, p < 0.01). The analysis conducted on 298 rhizosphere and root samples at harvest stage revealed that, congruently with the observed 299

diversification along the axis accounting for the major variation, the microhabitat remained 300 the major driver of the tomato communities (R² 47%, Adonis test, 5,000 permutations, p 301 <0.01) while the individual fertiliser treatments impacted these plant-associated microbial 302 303 assemblages to a lesser, but significant, extent (R²13%, Adonis test, 5,000 permutations, p <0.01). This suggest that, rather than on richness per se, the fertiliser treatment impacts on 304 the abundances and phylogenetic assignments of members of the tomato microbiota. 305 Remarkably, the Bray-Curtis matrix produced a congruent results, although the temporal 306 effect (i.e., nursery vs. harvest time) explained slightly more variation (~ 29%; 307 Supplementary Figure S2) than microhabitat diversification manifested along the second 308 axis of variation (~ 26%; Supplementary Figure S2). Crucially, also in this case the observed 309 diversification was supported by a PERMANOVA which attributed a R² of 23% to the 310 microhabitat, a R² of 29% to the 'Nursery/Harvest effect' and a R² of 3% to their interactions 311 (Adonis test, 5,000 permutations, p < 0.01). 312

313 Differential bacterial enrichments define microhabitat and treatment "signatures" on the 314 field grown tomato microbiota

To gain insights into individual members of the tomato microbiota responsible for the observed diversification we implemented a series of pair-wise comparisons among microhabitats and treatments at harvest stage. We took a two-pronged approach. First, we identified bacteria underpinning the microhabitat effect i.e., the selective enrichment of bacteria in the roots and the rhizosphere microhabitats amended with no fertiliser. Next, we assessed the effect of the fertiliser treatment on roots and rhizosphere bacterial composition by comparison with bacteria enriched in untreated samples.

This allowed us to identify 170 bacterial OTUs whose abundance was significantly enriched in and differentiated between rhizosphere specimens and unplanted soil samples (Wald test, p <0.01, FDR corrected; Supplementary database 1). Similarly, we identified 374 bacterial OTUs whose abundance was significantly enriched in and differentiated between root

specimens and unplanted soil samples (Wald test, p <0.01, FDR corrected; Supplementary 326 database 1). Of these differentially enriched bacteria, 96 OTUs represented a set of tomato-327 competent OTUs capable of colonising both the rhizosphere and root environments. When 328 329 we then looked into the taxonomic affiliations of this tomato-competent microbiota, we discovered that it is dominated by members of Actinobacteria, Bacteroidetes, Alpha-, Beta-330 , Gamma- and Deltaproteobacteria as well as Verrucomicrobia (Figure 4). Strikingly, the 331 taxonomic investigation revealed a bias for Actinobacteria in the root compartment, possibly 332 reflecting an adaptive advantage of members of this phylum in colonising the endophytic 333 environment. 334

Interestingly, each fertiliser treatment had a distinct impact on these tomato-enriched microbiota. The pelleted digestate (PE) and the slow-acting synthetic fertiliser (SRMF) yielded the highest number of uniquely enriched OTUs regardless of the microhabitat investigated, albeit with a distinct pattern: the SRMF had a more pronounced effect on the rhizosphere communities while the PE impacted more on the bacteria thriving in association with root tissues. (Wald test, p <0.01, FDR corrected; Figure 5; Supplementary database 1).

Interestingly, when we inspected the taxonomic composition of the bacteria differentially 341 impacted by the fertiliser treatment we observed an increase of the number of OTUs 342 belonging to phylum of Actinobacteria. In particular, PE had 12 OTUs out of 80 and 14 OTUs 343 out of 105, in root and rhizosphere, respectively, belonging to phylum Actinobacteria. While, 344 MF had 15 OTUs out of 38 and 22 OTUs out of 49 in root and rhizosphere, respectively, 345 belonging to phylum Actinobacteria (Supplementary database S1). Within this phylum we 346 observed the presence of OTUs classified as Streptomyces spp., Agromyces sp., 347 Microbispora sp. and Actinoplanes spp. 348

Together these data suggested that the enrichment of specific bacteria underpins the observed microhabitat effect whose magnitude is fine-tuned by the applied fertiliser.

351 Organic- and synthetic-based fertiliser trigger different metabolic capacities in the tomato 352 root microbiota

To investigate the ecological significance of the observed differential recruitments among 353 fertiliser treatment we employed a predictive metagenomics approach. Briefly, we inferred 354 in silico the functions encoded by the tomato microbiota at harvest stage (Materials and 355 Methods) and we grouped the samples in digestate-based (i.e., PE, LD and SRLD; hereafter 356 'organic') and treatments containing at least a synthetic component (i.e., SC, MF and SRMF; 357 hereafter 'mineral'). We observed that the functions putatively encoded by the communities 358 exposed to either organic or mineral fertilisers can discriminate between treatments in both 359 microhabitats (PERMANOVA: Rhizosphere samples $R^2 = 14\%$, p value <0.01, 5,000 360 permutations; Root samples $R^2 = 16\%$, p value <0.01, 5,000 permutations). Congruently, 361 we identified a set of 14 functions differentially enriched between root communities exposed 362 to either group of treatments (Welch t-test, p < 0.01, FDR corrected; Figure 6). Interestingly, 363 we observed a striking dichotomy between the two groups of treatments: communities 364 exposed to mineral fertilisers are predicted to enrich for genes implicated in the ABC 365 transporter machinery while bacteria exposed to the organic treatments are predicted to 366 enrich for genes implicated in the two-component system. These two set of genes are 367 dominant in communities exposed to both treatments and are also associated to additional 368 distinct enrichment patterns, most notably including nitrogen metabolism (organic 369 communities) and tetracycline biosynthesis (mineral communities). 370

These results suggest that, within tomato roots, the observed taxonomic diversification underpins a functional specialisation of the microbiota which, in turn, may impact on plant growth development and health.

374 **DISCUSSION**

This study revealed that all nitrogen treatments led to an increase of tomato production in comparison with the no fertilization treatment (fold change between 0.8 and 1.73) confirming

that, in the tested conditions, nitrogen limits the yield potential of processing tomato crops 377 as observed in previous studies (Ronga et al., 2015; Ronga et al., 2017). Yet, despite the 378 same amount of nitrogen was applied in each treatment (i.e., 150 kg ha⁻¹), all the treatments 379 380 were statistically different from each other. A prediction of this observation is that, under the tested conditions, the nature of the fertilisers, rather than the amount of nitrogen per se, 381 affect the yield and the fruit quality of tomato plants. These observations and the putative 382 contribution to fertiliser use efficiency of the microbial communities thriving at the root-soil 383 interface (Alegria Terrazas et al., 2016), motivated us to investigate relationships between 384 yield traits and the composition of the tomato rhizosphere and root microbiota under field 385 conditions. 386

387 The tomato rhizosphere and root microbiota are gated communities

First, we characterised the rhizosphere and root microbiota of processing tomato with no 388 treatment. Both alpha and beta diversity discriminated between the communities of 389 seedlings and adult plants. Despite these differences, which could be attributed to both 390 abiotic, e.g., time of residence in soil (Dombrowski et al., 2017), and biotic factors, e.g., 391 developmental-conditioned rhizodeposits (Chaparro et al., 2014), it is striking to note how 392 tomato plants displayed a rhizosphere and root compartmentalisation regardless of the 393 394 developmental stage. This is congruent with the observation that in rice, the assembly and structural diversification of the microbiota is a rapid process which reaches a steady-state 395 level within a few weeks from germination (Edwards et al., 2015). Closer inspection of the 396 rhizosphere and root profiles at harvest stage indicates that these plant-associated 397 communities are phylogenetically related to those of unplanted soil, suggesting that the 398 initial tomato microbiota is further modulated by the growing conditions. 399

Despite this apparent relatedness, the selective enrichment of individual bacterial members of the microbiota discriminates between rhizosphere and root communities for mature plants from unplanted soil profiles (Figure 4). These enrichments displayed a bias for members of 17 the phyla Actinobacteria, Bacteroidetes, Proteobacteria (including the classes Alpha--, Beta, Delta- and Gammaproteobacteria) as well as Verrucomicrobia. Members of these taxa
have routinely been reported in studies focussing on plant-competent bacteria under both
laboratory and field conditions (Bulgarelli et al., 2013; Walters et al., 2018), suggesting that
the experimental approach followed in this study can be considered representative for fieldgrown processing tomato.

However, we noticed a differential selective pressure on the bacteria thriving either in the 409 rhizosphere or in the root tissue: this latter environment produced more distinct profiles, i.e. 410 more differentially enriched bacteria compared to unplanted soil, than the ones retrieved 411 from the soil surrounding the roots. This indicates that the diversification of the tomato-412 inhabiting microbial communities from the surrounding soil biota initiates in the rhizosphere 413 and progresses through the root tissue, where it produces a more pronounced microbiota 414 diversification compared to unplanted specimens. This observation is reminiscent of the 415 recruitment patterns of other crops such as barley (Bulgarelli et al., 2015) but it is in striking 416 contrast with studies conducted with both model (Bulgarelli et al., 2012) and field-grown 417 (Rathore et al., 2017) Brassicaceae, whose 'rhizosphere effect' appears negligible. 418

We further noticed that the "root effect" on the microbiota was exerted also at phylogenetic level with a bias for the enrichment Actinobacteria. This observation is in apparent contrast with results gathered from the recent seed-to-seed characterisation of the tomato microbiota which revealed that, albeit averaging 8% of the sequencing reads across microhabitats, members of this phylum did not significantly discriminate root from rhizosphere specimens (Bergna et al., 2018). However, it is worth mentioning that these two studies differed in terms of both soil type and plant genotype used.

Together, our results suggest that both species- and soil-specific traits govern the assemblyof the tomato microbiota in field-grown crops.

428 Nitrogen source impacts on the structural and functional composition of the tomato

429 microbiota

Next, we investigated the impact of the type of nitrogen fertiliser on the tomato microbiota 430 and we demonstrated that each treatment produced "distinct signatures", represented by 431 specific selective enrichment, on both the rhizosphere and root communities. Despite 432 microhabitat-associated variation, the effect of the application of pelleted digestate (PE) 433 resulted in the most distinct microbial profile in the root compartment and the second largest 434 number of specifically enriched OTUs in the rhizosphere Of note, the slow-acting mineral 435 fertiliser (SRMF) follow a "complementary" pattern: its application yielded the greatest and 436 the second greatest number of differentially enriched OTUs compared to untreated samples 437 in the rhizosphere and root profiles, respectively. Remarkably, these two treatments had a 438 discernible effect also on crop yield, with the PE treatment producing the best performance 439 among the various fertilisers. Our data are congruent with studies conducted on wheat which 440 observed a structural diversification of the soil and plant-associated communities exposed 441 to either mineral or organic fertilisers (Kavamura et al., 2018). Yet, the numerical shift in 442 terms of OTUs differentially enriched per se cannot explain the potential impact of these 443 communities on crop yield: owing to the fact that the SMRF treatment, which is associated 444 to a significant reduction in yield traits (compared to PE) is capable of triggering a 445 comparable OTU enrichment. 446

We therefore focused our attention on the taxonomical composition of the rhizosphere and 447 root communities. In particular, we noticed that the proliferation of Actinobacteria in the root 448 compartment was retained in the various treatments. The enriched Actinobacteria included 449 Streptomyces Agromyces sp., Microbispora and Actinoplanes 450 spp., sp. spp. (Supplementary Database 1). Streptomyces spp. are well-known bacteria able to produce a 451 wide diversity of bioactive compounds able to promote plant growth and health (de Jesus 452 Sousa and Olivares, 2016). On the other hand, members of the genus Streptomyces are 453

responsible of economically relevant plant diseases, most notably common scab of potato
caused by *S. scabies* (Loria et al., 2006).

Thus, the taxonomic diversification triggered by both microhabitat and treatment may underpin a functional diversification of the microbiota at the cross-road of mutualism and inter-species competition.

This functional diversification of the root communities is manifested by the differential 459 enrichments of ABC transporter genes (mineral) and the two-component system (organic). 460 Although predictive metagenomics is inherently limited by fact that the individual 461 phylogenetic marker used (i.e., the 16S rRNA gene) may fail to recapitulate the genetic 462 diversity existing among strains of the same phylogenetic lineage (Karasov et al., 2018), 463 ABC transporters have previously been identified as genes underpinning rhizosphere 464 competence in the microbiota of wheat and cucumber (Ofek-Lalzar et al., 2014). Likewise, 465 the two-component system is required for the rhizosphere colonisation of the biocontrol 466 agent Pseudomonas fluorescens WCS365 (De Weert et al., 2006). These observations 467 indirectly support the results gathered from our predictive metagenomics approach. Owing 468 the role played by these classes of genes in uptake of organic compounds (e.g., root 469 exudates, cellular secretion) and stimulus-response mechanisms (e.g., chemotaxis) 470 respectively, it is tempting to hypothesize that the different source of nitrogen define a 471 different metabolic status in and in the vicinity of tomato roots which, in turn, requires a 472 prompt adaptation of the root-inhabiting communities. 473

For instance, experimental data indicate that the abundance of phytoavailable nitrogen, i.e., the scenario of mineral fertiliser treatments, tends to repress the proliferation and activity of members of the microbiota (Ramirez et al., 2012; Terrazas et al., 2019), and this in turn may be reflected in the metabolism of secondary compounds (terpenoid and polyketide metabolism) and membrane transport (ABC transporters).

Page 21 of 43

A "true" comparative metagenomics investigation, whereby the individual communities are subjected to shot-gun sequencing, will be ultimately necessary to test these hypotheses.

We further hypothesize that this adaptation is modulated by mineral nitrogen availability, as manifested by the differential enrichment of functions associated to nitrogen metabolism *per se* and aminoacids. This observation is congruent with results gathered from monocots wheat (Kavamura et al., 2018) and rice (Zhang et al., 2019) and suggests a cross-species pattern whereby plant's adaptation to nitrogen forms and availability is mediated, at least in part, by the associated microbiota.

Finally, it is interesting to note how the production of antibiotics, namely tetracyline, is also among the functions differentially enriched between fertilisers. It is becoming increasingly clear how plant-associated bacteria can act as a reservoir of antimicrobial genes (Cernava et al., 2019) which can be deployed during inter-organismal competition in the plant microbiota. This hypothesis could be tested by leveraging on indexed- and genomeannotated bacterial collection for the tomato microbiota, similar to the approach pursued with bacteria isolated from other plant species (Levy et al., 2018).

Our investigation suggests that the bacterial microbiota of field-grown processing tomato is the product of a selective process that progressively differentiates between rhizosphere and root microhabitats. This process initiates as early as plants are in a nursery stage and it is then more marked when plants reached the harvest stage. This selection *a*) acts both on the relative abundances and phylogenetic assignments of members of the tomato microbiota, *b*) is modulated, at least in part, by the nitrogen fertiliser provided which, in turn, *c*) triggers different microbial metabolic specialisations within tomato roots.

It is important to mention that the nitrogen fertiliser may also represent a microbial inoculant *per se*, in particular in the case of organic-based amendments. For instance, a comparative study of 29 different full-scale anaerobic digestion installations revealed that Firmicutes,

followed by Bacteroidetes and Proteobacteria, dominated the resulting microbial 504 communities (De Vrieze et al., 2015). Considering the plant-associated profiles observed in 505 this study, in particular the enrichment of Actinobacteria in the root communities, it is 506 legitimate to hypothesize that the input digestate bacteria may act as in inoculum for a part 507 of the tomato microbiota, which is further fine-tuned by the exposure to soil microbes. Future 508 studies, integrating the microbial profiling of the input fertiliser treatment, will be required to 509 accurately elucidate microbial dynamics associated with the application synthetic (i.e., germ-510 free) and organic fertilisers. 511

512 Towards a lab-in-the-field approach to harness the potential of plant microbiota for climate-513 smart agriculture

Our experiments represent an example of how cultivation-independent approaches can be 514 efficiently deployed to investigate the plant microbiota under field conditions. Although this 515 type of investigation is not novel per se in tomato (Toju et al., 2019), our results revealed 516 fundamentally novel insights into plant's adaptation to nitrogen fertilisers and the implication 517 for crop yield. Similar to what has recently been postulated for tomato pathogen protection 518 (Kwak et al., 2018), our results predicts that the use of field-derived, sequencing data will 519 allow scientists to identify "signatures" of the plant microbiota that can be targeted to 520 enhance plant performance. This approach, which we define as lab-in-the-field, will be key 521 towards the rationalisation of nitrogen (and other treatments) application in agriculture and 522 we anticipate will pave the way for the effective exploitation of the plant microbiota for 523 agricultural purposes (Schlaeppi and Bulgarelli, 2015; Toju et al., 2018). 524

525

526 AUTHORS CONTRIBUTION

527 DR conceived of and designed the field experiment. DR and FC harvested the field data and 528 samples. FC, RAT, EF and DB conceived of and designed the analysis of the microbiota. 529 FC, MC, CVGA, SR-A performed the microbiota experiments. FC, RAT and DB analysed Page 23 of 43

the sequencing data. FC, RAT and DB wrote the initial draft of the manuscript. All the authors

discussed the results and commented on the manuscript.

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551 CONFLICTS OF INTERESTS

552 The authors declare no conflicts of interest.

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TABLES

Table 1. Composition and information on fertilisers used in this study.TOC = Total organic carbon; N = Nitrogen; P = Phosphorus; K = Potassium; H2O = water

content

Treatment	TOC%	(N)%	(P)%	(K)%	H ₂ O%	Additional information
Synthetic fertiliser (MF)		41				Ammonium nitrate (N 26%) and calcium nitrate (N 15%)
Pelleted digestate (PE)	39.70	1.50	2.50	2.00	7.80	(Pulvirenti et al., 2015)
Slow acting liquid digestate (SRLD)	3.74	0.34		0.95		Liquid digestate plus the nitrogen stabilizer Vizura [®] (BASF, 2 L ha ⁻¹),
Liquid digestate (LD)	3.74	0.34		0.95		EC 1.07 dS m-1 and pH 8.3
Organo-mineral fertiliser (SC)	10.50	10.00	5.00	15.00	7.00	Produced by SCAM Spa (Modena, Italy), based on solid digestate for the organic fraction
Slow acting synthetic fertiliser (SRMF)		15.00	15.40	15.00		NPK Original Gold® (Compo Expert)

741 FIGURE LEGENDS

742 Figure 1. Effect of the nitrogen treatments on tomato yield traits

Mean and standard deviation of (**A**) marketable yield and (**B**) Brix of tomato plants exposed to the following treatments: LD (Liquid Digestate), SRLD (Slow acting Liquid Digestate), PE (Pelleted Digestate); SC (Organo-mineral fertiliser); MF (Mineral Fertiliser); SRMF (Slow acting Mineral Fertiliser). Different letters denote statistically significant differences between treatments by Analysis of Variance (ANOVA). Means were compared using Bonferroni's test at the 5% level (P <0.001).

Figure 2. The tomato root microbiota is a gated community

Average (A) number of observed OTUs, (B) Chao 1 index and (C) Shannon index computed 750 on the indicated rhizosphere and root specimens. Abbreviations LD (Liquid Digestate), 751 SRLD (Slow acting Liquid Digestate), PE (Pelleted Digestate); SC (Organo-mineral 752 753 fertiliser); MF (Mineral Fertiliser); SRMF (Slow acting Mineral Fertiliser). Asterisks denote statistically significant differences between microhabitat by non-parametric Wilcoxon rank 754 sum test (P < 0.01). Different blue letters within individual microhabitats denote statistically 755 significant differences between treatment means by Kruskal-Wallis non parametric analysis 756 of variance followed by Dunn's post-hoc test (P < 0.05); ns, no significant differences 757 observed. 758

Figure 3. The tomato rhizosphere and root microbiota host compositionally differentcommunities.

PCoA calculated using a weighted UniFrac matrix calculated on the OTUs clustered at 97%
identity among the indicated microhabitat and treatments. Abbreviations: LD (Liquid
Digestate), SRLD (Slow acting Liquid Digestate), PE (Pelleted Digestate); SC (Organomineral fertiliser); MF (Mineral Fertiliser); SRMF (Slow acting Mineral Fertiliser).

Figure 4. The enrichment of Actinobacteria is a distinctive feature of the tomato rootmicrobiota.

Phylogenetic relationships of the OTUs enriched in rhizosphere and root compartment. Individual external nodes represent one of the OTUs enriched in either (or both) rhizosphere or root samples in no treatment conditions (Wald test, P value < 0.01, FDR corrected) whose colour reflects their taxonomic affiliation at Phylum level. A black bar in the outer rings depicts whether that given OTU was identified in the rhizosphere- or root-enriched subcommunities, respectively. Phylogenetic tree constructed using OTUs 16S rRNA gene representative sequences.

774

Figure 5. Nitrogen fertiliser modulates bacterial enrichment in the tomato rhizosphere and

776 root compartments.

Number of OTUs significantly enriched (Wald test, P value < 0.01, FDR corrected) in the 777 indicated treatment versus untreated controls in (A) rhizosphere and (B) roots. In each 778 panel, blue bars denote the total number of enriched OTUs for a given treatment, the black 779 bars denote the magnitude of the enrichment in either the individual treatment or among two 780 or more treatments highlighted by the interconnected dots underneath the panels. 781 Abbreviations: LD (Liquid Digestate), SRLD (Slow acting Liquid Digestate), PE (Pelleted 782 Digestate); SC (Organo-mineral fertiliser); MF (Mineral Fertiliser); SRMF (Slow acting 783 Mineral Fertiliser). 784

Figure 6. Digestate- and mineral-based fertilisers trigger a functional diversification of the
tomato root microbiota.

Prokaryotic functions discriminating between Digestate-based (indicated as 'R_organic':
 Liquid Digestate; Slow acting Liquid Digestate and Pelleted Digestate) and mineral-based
 fertilisers (indicated as 'R_mineral': Organo-mineral; Mineral Fertiliser and Slow acting

- 790 Mineral Fertiliser) retrieved from Tax4Fun functional profiles (Welch's t-test FDR corrected,
- 791 p<0.01).



Figure 1. Mean and standard deviation of (A) marketable yield and (B) Brix of tomato plants exposed to the following treatments: LD (Liquid Digestate), SRLD (Slow acting Liquid Digestate), PE (Pelleted Digestate); SC (Organo-mineral fertiliser); MF (Mineral Fertiliser); SRMF (Slow acting Mineral Fertiliser). Different letters denote statistically significant differences between treatments by Analysis of Variance (ANOVA). Means were compared using Bonferroni's test at the 5% level (P < 0.001).

159x155mm (150 x 150 DPI)



Figure 2.Average (A) number of observed OTUs, (B) Chao 1 index and (C) Shannon index computed on the indicated rhizosphere and root specimens. Abbreviations LD (Liquid Digestate), SRLD (Slow acting Liquid Digestate), PE (Pelleted Digestate); SC (Organo-mineral fertiliser); MF (Mineral Fertiliser); SRMF (Slow acting Mineral Fertiliser). Asterisks denote statistically significant differences between microhabitat by non-parametric Wilcoxon rank sum test (P < 0.01). Different blue letters within individual microhabitats denote statistically significant differences between treatment means by Kruskal-Wallis non parametric analysis of variance followed by Dunn's post-hoc test (P < 0.05); ns, no significant differences observed.

139x228mm (300 x 300 DPI)



Figure 3. PCoA calculated using a weighted UniFrac matrix calculated on the OTUs clustered at 97% identity among the indicated microhabitat and treatments. Abbreviations: LD (Liquid Digestate), SRLD (Slow acting Liquid Digestate), PE (Pelleted Digestate); SC (Organo-mineral fertiliser); MF (Mineral Fertiliser); SRMF (Slow acting Mineral Fertiliser).

166x193mm (300 x 300 DPI)



Figure 4. The enrichment of Actinobacteria is a distinctive feature of the tomato root microbiota. Phylogenetic relationships of the OTUs enriched in rhzosphere and root compartment. Individual external nodes represent one of the OTUs enriched in either (or both) rhizosphere or root samples in no treatment conditions (Wald test, P value < 0.01, FDR corrected) whose colour reflects their taxonomic affiliation at Phylum level. A black bar in the outer rings depicts whether that given OTUs was identified in the rhizosphere- or root-enriched sub-communities, respectively. Phylogenetic tree constructed using OTUs 16S rRNA gene representative sequences.

163x191mm (300 x 300 DPI)



Figure 5. Nitrogen fertiliser modulates bacterial enrichment in the tomato rhizosphere and root compartments.

Number of OTUs significantly enriched (Wald test, P value < 0.01, FDR corrected) in the indicated treatment versus untreated controls in (A) rhizosphere and (B) roots. In each panel, blue bars denote the total number of enriched OTUs for a given treatment, the black bars denote the magnitude of the enrichment in either the individual treatment or among two or more treatments highlighted by the interconnected dots underneath the panels. Abbreviations LD (Liquid Digestate), SRLD (Slow acting Liquid Digestate), PE (Pelleted Digestate); SC (Organo-mineral fertiliser); MF (Mineral Fertiliser); SRMF (Slow acting Mineral Fertiliser).

167x226mm (300 x 300 DPI)



Figure 6. Prokaryotic functions discriminating between Digestate-based (indicated as 'R_organic': Liquid Digestate; Slow acting Liquid Digestate and Pelleted Digestate) and mineral-based fertilisers (indicated as 'R_mineral': Organo-mineral; Mineral Fertiliser and Slow acting Mineral Fertiliser) retrieved from Tax4Fun functional profiles (Welch's t-test FDR corrected, p<0.01).

91x253mm (300 x 300 DPI)



Supplementary Figure S1. Images of the filed at (A) plant's transplant and details of (B) an individual tomato plant and (C) the plantation at harvest stage.

131x87mm (300 x 300 DPI)



Supplementary Figure S2. PCoA calculated using a Bray-Curtis matrix calculated on the OTUs clustered at 97% identity among the indicated microhabitat and treatments. Abbreviations: LD (Liquid Digestate), SRLD (Slow acting Liquid Digestate), PE (Pelleted Digestate); SC (Organo-mineral fertiliser); MF (Mineral Fertiliser); SRMF (Slow acting Mineral Fertiliser).

224x189mm (300 x 300 DPI)