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Increased exopolysaccharide production and microbial activity affect soil water retention and field performance of tomato under water deficit

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3 Summary

4 According to the literature, biological processes in the rhizosphere could play a role in the 5 adaptation of plants to droughts under a changing climate. A previous study has identified significantly different productivity level and fruit quality for two tomato varieties under wa-6 ter deficit conditions. We conducted a field study, with and without water deficit, with these 7 two varieties to examine whether microbiological activity and exopolysaccharides concen-8 tration could affect soil hydrophysical properties. The rhizosphere soil had indeed distinct 9 bio-chemical and hydrophysical properties between the two cultivars and between the two 10 11 water-related conditions. The quantity of soil exopolysaccharide and/or nitrogenous sub-12 stances, and the activity of microorganisms (fungi in particular) explains part of the soil water retention measurements. In addition, these mechanisms are significantly accentuated for 13 the cultivar with the best productive capability under water-limited condition-i.e. with 14 15 commercial yield, fruit dry matter and water use efficiency which are respectively 35%, 28%, 16 and 31% higher for the productive cultivar.

17 Key words

18 Rhizosphere, Water deficit, Soil water retention, Exopolysaccharide, Microbial activity

19 Introduction

The current increase in temperature, evapotranspiration, and in the seasonal variation of water regimes linked to climate change lead to a higher pressure on the water resources, which affects the agricultural production (Jia et al., 2019, Boeck et al., 2011). This calls for the development of more water-efficient agroecological systems, able to withstand more frequent water deficit periods (Lipiec et al., 2013). The water retention capacity of soils has a strong influence on their capacity to satisfy crops' water requirements and could thus mitigate the effects of extreme climatic events (Łabędzki, 2016).

27 The rhizosphere is the area of soil around the roots influenced by plant roots and where the plant, the soil, and the associated microorganisms interact (Bargett et al., 2014; Hinsinger et 28 al., 2009; Angers et al., 1998). The rhizosheath is the portion of soil that is physically in con-29 30 tact with the root system. The rhizosheath is part of the rhizosphere, but rhizosphere may extend beyond the rhizosheath (Marasco et al., 2018; Pang et al., 2017). The rhizosphere is 31 32 both a "hot spot" of biological activity and metabolic processes in the soil (Zhang et al., 2020), and the zone where root water and nutrient uptake takes place. Moreover, the rhizo-33 sphere has specific physicochemical properties: soil aggregates are more stable; bulk densi-34 ty, porosity, water and nutrients transfer as well as pH are modified with respect to the bulk 35 36 soil (Hinsinger et al., 2009). These properties are linked to the specific biochemical activities 37 of roots, such as rhizodeposition and exudation of organic compounds (Personeni et al., 38 2017), which can then consumed by the associated microbial communities (van Veelen et al. 39 2018). In addition, these microbial communities, stimulated by the rhizosphere environment, can also excrete similar organic molecules, including Exo-polymeric Substances (EPS), 40 such as Exo-PolySaccharides (EPSac) (Redmile-Gordon et al., 2014). EPSac may also consti-41

42 tute the majority of root mucilage (Nazari, 2021). These polymeric substances contribute to the stability of soil aggregates (Crouzet et al., 2019), improve soil water retention (Czarnes et 43 al., 2000), and present either hydrophilic or hydrophobic characteristics depending on their 44 state of hydration (Carminati and Vetterlein, 2013). Until now, few studies have focused on 45 the influence of both the physicochemical and biological properties of the rhizosphere on 46 47 plant water use efficiency (WUE) (Benard et al. 2019). However, since all the water transpired by a plant must cross the rhizosphere soil layer, understanding its functioning and its 48 properties is critical (Bengough, 2012). Thus, acting on the functioning of the rhizosphere 49 could be a lever to maintain the productivity of crops suffering from water deficit (Ahmed et 50 51 al., 2018, Doussan et al., 2015).

52 Tomato (Solanum lycopersicum) is widely grown in the Mediterranean region and its produc-53 tion is one of the most important worldwide in terms of market value and quantity (FAO-STAT, n.d.). Understanding how this crop adapts to droughts is therefore of prime im-54 portance. Under glasshouse conditions, water deficit negatively affects plant growth, leaf 55 water potential and gas exchange intensity during the early stages of tomato development 56 57 (Duan et al., 2016) as well as fruit mass and composition at the latter stages (Ripoll et al. 2016). The impact of water deficit was also observed in the field for industrial tomato culti-58 59 vars (Arbex de Castro Vilas Boas et al., 2017). In particular, under water deficit, the cultivar Terradou, compared to H1015, showed a higher dry yield (on average 76 kg.ha⁻¹ and 56.kg 60 ha⁻¹ respectively) and a higher Water Use Efficiency (WUE) (41 and 33 kg.m⁻³ respectively). 61 Terradou fruits had also a higher sugar content (Glucose, Fructose, higher SSC index) while 62 H1015 fruits had more acids (Malic, Citric). The fruit puree viscosity increased in general 63 when the plants underwent water deficit, but more markedly for H1015 than for Terradou. 64

As water deficit is one of the main factors limiting yield in tomatoes (Costa et al., 2007;
Patanè and Consentino, 2010), depending on its intensity, duration, and timing in the crop
cycle (Rinaldi et al., 2007), the response of these cultivars to water deficit deserves further
testing.

69 While several studies have investigated the influence of microbial activity, and in particular 70 mycorrhizae (Cavagnaro et al., 2006; Hallett et al., 2009; Bowles et al., 2016), on tomato 71 phenology and its yield components, it is worthwhile to investigate the influence of soil hy-72 drophysical properties. Understanding this possible link between soil bio-chemical and hy-73 drophysical parameters might broaden our understanding of the plants' responses to drought. Our objective was therefore to observe in the field the links between different bio-74 75 chemical parameters of the soil (i.e., microbial biomass and activities, EPSac amounts) and 76 the soil water retention (i.e. Field Capacity and Permanent Wilting Point). We took into ac-77 count the influence of (i) the rhizosheath of tomatoes, (ii) the season stage of the plant development, (iii) the tomato cultivars, and (iv) the water conditions (well irrigated or deficit). 78

We carried out a field experiment with the two cultivars Terradou and H1015 on the same 79 80 Mediterranean experimental site than previously used by Arbex de Castro Vilas Boas et al 81 (2017). We increased water deficit by limiting irrigation, and we measured the biophysical parameters mentioned above both in the rhizosheath and in the bulk soil. Our hypothesis 82 83 was that the EPS exudation and microbial activities would be higher for Terradou compared 84 to H1015, and that these processes would allow for a better yield resilience to water deficit conditions for Terradou. We also assumed that these processes would be relatively more 85 important in the rhizosheath and that their dynamics would follow the development stages 86 87 of the plant.

88 Materials and methods

89 Experimental set-up and sampling

90 The experimental site is an industrial tomato field of a SONITO (French interprofessional or-91 ganisation of the tomato sector). It is located under a Mediterranean climate at Piolenc, 92 France (44°11'16.9 "N 4°48'10.8 "E), and is used for a varietal trial. The soil is a Stagnosol (WRB classification; or Redoxisol in French classification) with a loam texture (main soil 93 properties are shown in Table 1). We studied two widely grown commercial cultivars of to-94 95 mato (Solanum lycopersicum L.): Terradou (T) and H1015 (H), previously tested by Arbex de 96 Castro Vilas Boas et al. (2017). The cultivation practices applied on this field correspond to 97 current conventional practices (density of 3.3 plants.m⁻², fertilization and pesticides). During this experimentation, two irrigation treatments were applied: from planting (May 18, 2018) 98 99 to fructification (July 23, 2018), all the plants received the same amount of irrigation water 100 via daily drip irrigation to compensate 87% of evapotranspiration losses, resulting in nearly 101 460 mm of total water input before the fructification (135 mm from rainfalls and 325 mm 102 from irrigation). Then, from fructification to harvest (August 9), half of the plots had water deficit (WD) conditions i.e., 57% of the water input of the fully irrigated plots (well-watered 103 "WW"). Each cultivar was grown on two plots (42 m length each) of two rows (33 cm apart) 104 105 of tomato plants on cultivation ridges distributed on both sides of the main irrigation tube. For each cultivar, one plot was WW and one plot was WD (Fig.S1). 106

107 Soil samples were collected at the beginning of the flowering stage (June 19, 2018) and later 108 at the fructification stage (August 2, 2018) before harvesting. Five sub-plots were randomly 109 chosen within the plots. In those sub-plots, we sampled two types of soil: the 'rhizosheath'

110 soil and the corresponding 'bulk' soil. For the rhizosheath soil sampling, three plants per cultivar in each sub-plot were excavated with their proximal root system and surrounding soil 111 using a spading fork (to a depth of 20 cm maximum). The plants were then shaken vigorously 112 113 by hand, but without breaking roots, until no more soil aggregates could be detached from roots. The soil remaining on root was collected and judged to be the rhizosheath soil. The 114 115 rhizosheath soil of the three selected plants was pooled (Göttlein, 2006). We selected three 116 subsamples of bulk soil located on the outside bare soil of the ridge facing the plants sampled for the rhizosheath soil over 0-20 cm depth (Fig S1). These three subsamples of bulk soil 117 118 were also pooled. All soil samples were air-dried, sieved at 2 mm and stored at 4°C until 119 analysis.

120 Measurements of soil and plant parameters

121 Soil microbial and fungal catabolic measurements

122 Substrate Induced Respiration (SIR) of soil was assessed with the FungiResp, adapted from the MicroResp[™] colorimetric technique (Ben Sassi et al., 2012; Chapman et al., 2007), con-123 124 sisting in a 96-deep-well microplate filled with soil subsamples. The soil water content was preliminary adjusted to 40% of the water holding capacity (about 0.22 g/g gravimetric water 125 126 content). 25 µl of organic substrates (6.7 mg.g⁻¹ dw soil) were added in each deep-well. Each 127 filled soil plate was then sealed face to face with a CO₂ trap microplate including a pH dye 128 indicator, and an intercalated silicone joint to individualize the plate wells. The systems were 129 then incubated in the dark at (23°C±1) for six hours. Absorbance of the gel CO₂-trap was 130 measured at 570 nm, before and after incubation. A calibration curve of absorbance versus 131 head space equilibrium CO_2 concentration (measured by gas chromatography) was fitted to a 132 regression model, which was then used to compute the amounts of released CO_2 in μg C-

133 CO2.g⁻¹ Dry Soil.h⁻¹. We measured the microbial basal respiration by adding 25 μl of demin-134 eralised water instead of organic substrate (M.Bas). In order to address substrate-induced 135 respiration according to fungal communities, we applied a bacterial inhibitor (25 µL for a final concentration of 78 µg Bronopol.g⁻¹ soil, concentration selected after performing a 136 137 dose/response curve on the respiration of this soil, Ben Sassi et al., 2012) to half of the soil 138 sub-samples distributed in the deep well-plate (the other half of the soil sub-samples re-139 ceived 25 µL of demineralised water). Glucose-induced respiration was used as a proxy of active total microbial biomass (MB) and fungal biomass (FB) in µg C.g⁻¹ DrySoil (Anderson 140 and Domsch, 1978). To access the catabolic profiles of fungal communities (F-CLPP) (F.Tre, 141 F.Cell, F.Ala, F.Gly, F.Mal, F.GlcN) and microbial communities (M-CLPP) (M.Tre, M.Cell, M.Ala, 142 143 M.Gly, M.Mal, M.GlcN) we used various organic substrates (trehalose, cellobiose, alanine, 144 glycine, malate, glucosamine, respectively) distributed in the deep well-plate, and at the same amounts of glucose, with and without the addition of bacterial inhibitor (Bérard et al., 145 146 2011).

147 Soil water retention measurements

The measurement of water retention was carried out for matric potentials of 0.3 bars (i.e., Field Capacity) and 15 bars (i.e., Permanent Wilting Point) on soil aggregates by using a pressure plate system. The triplicates of initially saturated samples (about 10 g) were placed in porous plate enclosures into which nitrogen gas was injected at a pressure of 0.3 or 15 bars. Samples were left to drain excess water until equilibrium was reached, with no outflow. The gravimetric water content (g water.g⁻¹ dry soil) was then measured by weighing the soil before and after drying at 105°C (24h).

155 Soil exopolysaccharides measurements

156 The ExoPolySaccharides (EPSac) that are principally involved in soil physical properties are 157 those tightly bound to the soil particles (Chen et al., 2014; Crouzet et al. 2019). We used the 158 physico-chemical Cation Exchange Resin (CER) technique (Redmile-Gordon et al., 2014; Bérard et al., 2020) to extract the bound-ExoPolymeric Substance (bound-EPS) from the soil. 159 160 Before bound-EPS extraction, loosely bound EPS were extracted (each 0.5 g of soil sample) 161 using 5 mL CaCl₂ (10⁻² M) under agitation (50 rpm) for one hour at laboratory temperature 162 (Redmile-Gordon et al., 2014; Bérard et al., 2020). The soil/CaCl₂ mixtures were then centri-163 fuged (8000 xg for 15 min at 10°C), and the supernatants were removed. The remaining centrifuged soil pellets were then extracted by the CER extraction technique. The CER (Dowex 164 Marathon C Na+ form, Sigma Aldrich, Steinheim, Germany) was previously washed with a 165 166 phosphate buffer (consisting of 2 mM Na3PO4. 12H2O, 4 mM NaH2PO4. H2O, 9 mM NaCl, 1 167 mM KCl, adjusted to pH 7 with 1M HCl and stored at 4 °C) until the pH of the solution stabi-168 lized to 7. In a centrifuge tube containing the soil pellet samples, the CER was added to ob-169 tain a ratio of 70 g dry CER per 1 g of soil organic matter. Five mL of phosphate buffer pH 7 170 were then added to the soil and CER. After manual stirring, the samples were incubated for 171 sixteen hours overnight under agitation (50 rpm) at room temperature. Bound-EPSac were 172 measured as the total carbohydrate content of the bound-EPS extracted from soil, using the phenol–sulphuric acid Dubois method (Dubois et al., 1956) with glucose as the standard. 173

174 Plant and Fresh Fruit Measurements

175 At harvest, for each cultivar and irrigation modality, five tomato plants were sampled. We 176 measured the commercial yield (CoY, in t.ha⁻¹) corresponding to the tomato marketable 177 fresh yield for the industry (red and yellow ones), and harvest losses due to rotten tomatoes 178 (RoY, in t.ha⁻¹). The dry mass of this commercial yield (DMY in kg.ha⁻¹) was also measured. We calculated the water use efficiency (WUE - kg.m⁻³) as the ratio between the commercial yield (CoY) and the total water volume provided by irrigation and rainfall to the plant, for a given surface. Some chemical analyses were also performed on the tomato fruits, such as the soluble solid content (SSC, in °Brix), the titrable acidity and the pH (as in Arbex de Castro Vilas Boas et al., 2017).

184 Data Analysis:

185 Statistical analysis

Four modalities were investigated in the experimental design: the type of soil-rhizosheath 186 187 or bulk (SOIL), the season—June or August (SEAS), the cultivar—Terradou or H1015 (CULT), 188 and the plots' water status modalities in August -WD or WW. 76 soil samples were analysed. Quantitative data were analysed using R (R core Team, 2017) with R Studio IDE (R Stu-189 190 dio Team, 2020). We first observed the statistical differences between the three first modali-191 ties (SOIL, SEAS, CULT) for each of the soil parameter studied (microbial, hydro-physical, and 192 chemical) at a level of significance of p < 0.05 with an ANOVA F-test (the R stat package and 193 anova() function). Table 2 shows the main effects and their interactions performed over the 194 whole dataset of soil parameters and on rhizosheath soil dataset. Table 2 also presents spe-195 cific analyses for June, August-WD irrigation and August-WW irrigation datasets, allowing us to compare the WW and WD data. Homoscedasticity of variances (Bartlett test) and normali-196 197 ty (Shapiro test) were checked before data analyses. The mean and standard deviation for 198 the groups of data analysed are reported in Table S1.

199 Principal Component Analysis

200 Each substrate-induced respiration measurement was normalized to the sum of all SIR 201 measurements, to exclude the microbial biomass effect on the individual respirations (Ben Sassi et al. 2012). Principal Component Analysis (PCA) was performed on these normalized 202 measurements of microbial and fungal catabolic profiles carried out on the seven organic 203 204 substrates: glucose, trehalose, cellobiose, alanine, glycine, malate, glucosamine. The Fac-205 toMineR R package with the fviz pca ind() function (Lê et al., 2008) was used to perform 206 PCA analysis. A first global PCA analysis was performed on dataset including the 7 microbial 207 or fungal variables and the whole 76 observations (2 cultivars \times 2 seasons \times 2 soil types \times 2 208 irrigation levels x 5 replicates, 4 data missing). Then, we performed PCA analyses on the June 209 data exclusively, as well as for the August-WW and the August-WD.

210 Linear regression models

211 Multiple linear regression models (Schielzeth, 2010; Whittingham et al., 2006) with the Im() 212 function in stats package were established to relate 0.3 bars and 15 bars water contents to 213 soil candidate predictor variables: EPSac, microbiological and fungal activity and biomass 214 measurements. A stepwise regression (Murtaugh, 2009) was used to select the best predictor variables using the AIC information criterion (Hegyi and Gramszegi, 2011) with the func-215 tion ols step both aic() from the olssr R-package. The Variable Inflation Factor (VIF) was 216 computed with the function ols vif tol() from the same package. The VIF must be lower 217 218 than 10 for the variable to be selected (Dormann, 2013). This way, we obtained between one and four predictor variables in the linear regression models. 219

220 Results

221 How soil type, season and cultivar influence soil indicators

Significant differences are observed between most of the bio-physicochemical properties of the rhizosheath soil of tomato plants and the corresponding "bulk" soil far from roots (Tab.2 and S1). Microbial and fungal biomass and activity are almost constantly significantly higher in rhizosheath. The same holds for water retention values at the Permanent Wilting Point (W15b) (p-value = 1.26e-04), the field capacity (W0.3b) showing the same trend (p-value = 6e-2). A higher EPSac amount is also denoted in the rhizosheath on this global dataset (mean respectively equal to 780 and 760 μ g.g⁻¹; p-value=7e-2) (Tab.2 and S1).

229 Microbiological activities, fungal activities, and the amount of EPSac are higher in August 230 compared to June (highly significantly) (Tab.2 and S1). Concerning hydraulic parameters, 231 W15b was also higher in August, but W0.3b is higher in June (p-value=5.4e-2 for both). This 232 temporal difference in hydraulic parameters increases when considering only the rhi-233 zosheath soil (respectively p-value=4.5e-3 and 1.2e-7). In June, some physicochemical and 234 microbial parameters show significantly higher values in the rhizosheath soil than in bulk soil, but for August-WW irrigation, most of the microbial indicators exhibit significant higher 235 236 values in the rhizosheath soil compared to bulk soil (Tab.2 and S1).

237 Concerning the tomato cultivars, a number of microbial soil indicators are significantly high-238 er for the T compared to the H cultivar (Tab.2 and S1). In June, we observe differences in 239 microbial indicators between H and T cultivars (T>H) only for the rhizosheath soils. The soil 240 indicators for these two varieties are then differentiated into two distinct behaviours accord-241 ing to the type of irrigation. Moreover, the analyses of variance show interactions between 242 the factors season and soil, for the water retention variable W0.3b and for the microbial 243 variables F.GlcN, F.Ala, M.Cell. The season and cultivar factors also show interactions, for the variables W15b, EPS, M.Mal and in the rhizosheath for the variables W15b, F.Ala, M.Gly,M.Mal (Tab.2).

246 Differences between soil indicators exacerbated under water deficit

247 Moisture content of soil samples from August campaign sampling differed between WD and 248 WW, with higher water content in WW plots compared to less irrigated WD plots (respective averages of 13.7 and 12.4 g.g⁻¹, p-value=0.031). In August-WD, we observe differences in 249 250 both W15b and W0.3b between rhizosheath and bulk soils, while only W15b values present 251 significant differences in August-WW (Tab.2). However, the trend is the same for the two 252 irrigation treatments: W15b values are higher in rhizosheath soils than in bulk soils while it is 253 the contrary for W0.3b. For August-WD, the soil of T cultivar exhibits parameters (W0.3b and 254 a number of microbial indicators) that are significantly higher than for H cultivar (for rhi-255 zosheath soils alone and for rhizosheath and bulk soils together). In August-WW, the differ-256 ences between cultivars are less pronounced than in August-WD and the ranking of cultivars 257 is inverted: Some microbial and fungal catabolic activities present higher values in H soils 258 compared to T soils (Tab.2 and S1).

259 Microbial and fungal catabolic profiles

The microbial and fungal catabolic profiles of the 76 soil samples were analysed with Principal Component Analysis (PCA). Considering the catabolic profiles from the global dataset, no discrimination appears between cultivars (T/H) and soil types (R/B) for microbial (Supp. Fig. 2-b, 2-c) and fungal (Fig. 1-b, 1-c) catabolic profiles. However, PCA shows differences between August and June sampling seasons. In August catabolic activity of microbial communities show more homogeneity between samples, with points concentrated in the center of the graph, compared to June (Supp. Fig.2, % total of variance=79.5). The fungal communities

show a discrimination between June and August catabolism, the later showing a higher difference in catabolism of nitrogen-containing organic substrates (alanine and glycine, glucosamine) (Fig.1-a, % total of variance=73).

When considering only June data, no discrimination in the catabolism of both the microbial and the fungal communities is denoted, either between the cultivars nor between rhizosheath and bulk soils (Fig.2-a; Fig.3-a; Supp. Fig. 2-a; Supp. Fig. 3-a).

273 On the contrary, in August there are differences in catabolic profiles. Microbial catabolic 274 profiles separate H and T cultivars, with higher catabolism of trehalose and glucose in T 275 plots, whereas glycine and malate are preferentially catabolized in H plots (Fig. 2-b, 2-c). The 276 differences in Fungal catabolic profiles observed in August are marked between rhizosheath 277 and bulk soils (Fig.3 b, c), especially at WD irrigation, where fungal communities preferential-278 ly catabolize nitrogen-containing organic substrates (glucosamine, glycine and alanine) 279 (Fig.3-c).

280 Relation between hydro-physicals, chemical and microbiological indicators

Figure 4 presents the relationship between EPSac amounts and the microbial or fungal biomasses. EPSac amounts increase with and correlate with microbial biomass (R^2 = 0.20; pvalue = 4.09e-5), as well as with fungal biomass (R^2 = 0.32; p-value = 1.28e-7).

Both for microbial and fungal biomasses, these correlations with EPSac amounts are stronger when considering bulk soil and weaker for rhizosheath soil solely (Tab. 3). When considering the data for June or August, no significant relationship appears (on global, rhizosheath or bulk soils data), with the exception of a weak relationship for fungal biomass in June (R^2 = 0.22; p-value = 1.6e-3) (Tab. 3).

289 The best linear models explaining water retention variables (W0.3b and W15b) with the 290 measured EPSac and microbial/fungal data variables as predictors are shown Table 4 (for 291 global dataset and for June or August data). Whatever the dataset considered, a significant correlation emerges between water retention and EPSac/microbial variables (R² from 0.16 to 292 293 0.74). Most often, the fungal indicators are the significant predictors and EPSac is linked to 294 W0.3b (except in August-WW irrigation). Fungal catabolism of alanine is also linked to W0.3b 295 (except in June). The link between water retention (W0.3b and W15b) and EPSac/microbial variations is stronger in August (R²=0.3 to 0.74) (and particularly in August-WD irrigation) 296 compared to June (R^2 =0.16 to 0.23). 297

298 Aboveground productive characteristics and fruit quality measurement

There is no significant difference between the two cultivars productive characteristics when well irrigated (WW) (Fig.5-a). However, under WD theses characteristics show systematic higher values for the T cultivar compared to H (Fig.5-b), with significant differences for CoY, DMY and WUE (p-value= 1.9e-2; 4.8e-2 and 1.9e-2 respectively) except for RoY, for which it is the opposite (H>T). Titrable acidity and pH of fruits show no significant differences. The soluble solid content (SSC) is slightly higher for H cultivar compared to T cultivar in context of deficit irrigation (mean respectively equal to 6.40 and 6.07, in °Brix ; p-value=3.4e-2).

306 Discussion

307 Influence of the rhizosheath on soil parameters

308 We highlighted a quantitative effect of the rhizosheath on soil parameters, with higher wa-309 ter retention values and higher amounts of exopolysaccharides in the rhizosheath compared 310 to the bulk soil. We also observed that total microbial and fungal biomass, as well as microbial and fungal catabolic activities were significantly higher in the rhizosheath. This differentiation was more pronounced at the fruit maturation stage in August (especially at WD) compared to June (rhizosheath data: Tab. 2 and S1, interactions between season and soil factors and also between season and variety factors). A phenology effect seems to occur here and increases during the season within the rhizosheath (Bardgett et al., 2005). Between June and August, the rhizosheath soil would have received a fresh carbon supply from the roots (especially EPSac) of the developed tomato plants, stimulating microbial biomass and activities and promoting water retention.

319 Qualitatively, these differences were more pronounced for the catabolic profile of fungal 320 communities at the fruit maturation stage and in water-deficit context (August-WD). Fungal 321 communities are linked to a higher catabolism of nitrogen containing organic substrates in 322 rhizosheath soils (Fig.1, Fig.2). We suggest the hypothesis that carbon exudation by roots boosted by water stress (Henry et al., 2007), could induce a "priming effect" in relation with 323 the stimulation of microbial and fungal exo-enzymes involved in the degradation of soil or-324 325 ganic nitrogen (Bardgett et al., 2014). Another possible cause would be that the develop-326 ment of nitrogen-fixing bacteria was favoured by EPSac secreted by the roots (Nazari, 2021) 327 in case of water deficit. The contribution of organic nitrogen by diazotrophic bacteria could 328 then have favoured a higher catabolism of nitrogen containing organic substrates in rhi-329 zosheath soils (Zuluaga et al., 2020). To confirm these hypotheses, additional measurements on the N cycle (N₂ fixing activities, enzymatic activities, nitrogenous forms...) would be nec-330 331 essary.

332 Biochemical activity of roots and their microbial communities. Relationship with333 soil water retention

Several below ground processes, possibly interacting, can allow the plant to compensate for water deficit. The particular hydro-physical properties of the rhizosheath are partly linked to the physical presence of the roots themselves (Aravena et al. 2011, De León-González et al. 2007), and to the production of EPS by the roots and microorganisms (Chenu 1993; Czarnes et al, 2000; Sher et al., 2020), which are also influenced by changes in moisture and nutrient conditions (Alami et al. 2000, Henry et al. 2007, Lynch and Whipps 1990, Redmile-Gordon et al. 2014, Roberson and Firestone 1992).

341 We observe that the amount of soil EPSac follows the same seasonal dynamics as soil micro-342 bial and fungal biomass and their catabolic activity, increasing from spring to summer, espe-343 cially in rhizosheath soils (Tab. 2, Tab.S1). Indeed, although low, the correlations were posi-344 tive between EPSac and microbial and fungal biomasses (the correlation with fungal bio-345 masses was higher - Tab. 3 and Fig. 4). In August, at the fruit maturation stage, water reten-346 tion at 0.3 and 15 bars was mostly linked to physico-chemical and microbial variables, and more significantly in case of water-deficit than with full irrigation (Tab. 4). In particular, the 347 EPSac was partially predictive of the water retention at field capacity (W0.3b) (Tab. 4). 348

The increase in root and microbial biological stimulation in the rhizosheath, notably by a higher EPS production than their consumption (Redmile-Gordon et al. 2014) could explain these results. This stimulation in the rhizosheath could have increased with the root density and the intensification of exudation per unit of root length linked to a phenology effect and a triggered by water deficit in case of deficit irrigation condition (Henry et al., 2007; Nazari et al., 2020).

355 Furthermore, fungal biomass and catabolic parameters also seem to be important variables356 explaining water retention: the fungal catabolism of alanine, an amino-acid, was almost sys-

357 tematically found as a predictor variable for water retention at field capacity (W0.3b). In 358 general, fungi are more tolerant to water stress than bacteria (Bérard et al., 2015). Rhizo-359 sphere fungi (and mycorrhizae in particular) are known to act on soil aggregation (and water 360 retention) (Freschet et al., 2021) through the physical action of their hyphae and their pro-361 duction of proteins (e.g. glomalin) and polysaccharides (e.g. scleroglucan), resulting in parti-362 cle sticking actions (Poirier et al., 2018). Studies involving mycorrhizae and tomatoes show 363 that mycorrhizae promote plant resistance to drought by stimulating nutrient cycles (including nitrogen) and root exudation (especially Amino Acids, which can play an osmoregulatory 364 365 role) (Bowles et al. 2016). Moreover, Cavagnaro et al. (2006) showed that mycorrhizae pro-366 mote soil aggregation in tomatoes rhizosheath, especially in the presence of mineral N. One hypothesis would be that tomatoes exude more organic compounds under water deficit 367 368 conditions, which would promote nitrogen catabolism (directly and/or through a "priming 369 effect" or diazotrophic bacteria stimulation as suggested above), fungal development (my-370 corrhization) and its direct or indirect effects on water retention. Thus, these different chem-371 ical and biological phenomena probably interact with the hydro-physical properties of the 372 soil (Hallett et al. 2009). However, our hypothesis is based on proxy measurements of bio-373 mass and fungal catabolism (FungiResp method). It would have been interesting to measure 374 the intensity of root colonization by mycorrhizae to confirm this hypothesis, which we were 375 unable to do during this study and could thus be done in future investigations.

Nevertheless, it seems that the fungal catabolism of alanine is related to the hydraulic properties of the soil without water deficit, while in conditions of water deficit, there could be a shift to other mechanisms explaining water retention, which would be related to EPSacs, among others. Indeed, in August, we found that the fungal catabolism of alanine explained the water retention of the soils with higher R2 values in the WW irrigated plots than in the irrigation-limited plots (WD): in particular for the W0.3b retention at WW irrigation the increase in R2 for F-Alanine was 0.33, while at WD irrigation it dropped to 0.2, with the appearance of the EPSac as an explanatory variable (increase in R2=0.18; Tab.5).

384 Possible relationships between aboveground cultivars characteristics and their385 belowground bio-physico-chemical soil parameters

386 The T cultivar presents a higher production potential compared to the H cultivar in the water 387 deficit situation, and irrigation seemed to mitigate these differences (Fig. 5). Arbex de Casto 388 Vilas Boas et al. (2017) showed in a previous similar experiment that WUE was better for T than H cultivar especially in WD conditions, ranging from 25 kg.m⁻³ for WW irrigation to 41 389 390 kg.m⁻³ for a WD situation in the field. Despite a much higher total water input in this 2018 391 field experiment and, thus, with lower mean WUE values (between 11 and 18 kg.m⁻³), the WUE was still significantly higher for T than for H cultivar in WD situation (Fig. 5), indicating 392 that H seems more sensitive to WD. 393

394 It seems that our biochemical observations on the soils (microbial activities, especially fungal 395 activities and amounts of EPSac, Tab.2) leading to a local modification of soil water retention 396 (Tab.4) are in agreement with the production potential of tomato cultivars T and H, as a 397 function of their water efficiency (Fig. 5). In fact, cultivar H, which seems to have better ex-398 pressed its productive potential under optimal irrigation conditions, presents in its rhi-399 zosheath in August-WW a higher F.Ala catabolism, which contributes to explain the water 400 retentions W0.3b and W15b. On the contrary, cultivar T seems to have better expressed its 401 production potential under water deficit conditions, in correspondence with higher water 402 retention W0.3b, together with higher EPSac amounts and higher indicators of fungal activi-403 ty and biomass in the T soil plots. Of course, if resistance/tolerance is also related to root 404 system characteristics (root density, root depth) (Fang et Xiong, 2015), our results point to 405 the fact that biochemical properties of the rhizosheath may also play a role.

The water retention properties of soils have a strong influence on crop development and could factor in the crop's capacity to withstand droughts. A better understanding of the possible links between soil bio-physicochemical parameters and hydrophysical properties, particularly in the rhizosheath (Bengough, 2012), could help improve crop resistance to water deficit. While creating water deficit conditions by limiting irrigation, we measured biophysical indicators of soil quality and rhizosheath functioning (in comparison to bulk soil). This field study brings new information regarding the links between soil parameters and productivity of two tomato cultivars and two types of irrigation.

414 Conclusion

415 Our study suggests that rhizosheath affects the soil biophysical parameters measured and 416 this influence increased between the end of the vegetative phase and the fruit maturation 417 stage. Secondly, it seems that hydro-physical properties of the soil (e.g., water retention capacities) are partly linked to two interacting phenomena: the presence in the rhizosheath 418 419 environment of specific substances such as exopolysaccharides and/or nitrogenous sub-420 stances from roots and microorganisms, and the presence and activity of fungi (possibly my-421 corrhizae). The importance of fungal activity stood out during the study and it would be in-422 teresting to investigate their effects further by using tools dedicated to fungi and mycorrhi-423 za.

424 In addition, these two aforementioned relationships could be a part of the adaptation 425 mechanisms of the Terradou cultivar to water deficit, enabling higher agricultural productivi-426 ty via a higher plant WUE. This study confirms the observations made by Arbex de Castro

427 Vilas Boas et al (2017) and provides hypotheses related to the influence of the rhizosheath428 on the responses of these cultivars to water deficit.

429 Our observations—i.e., correlations between the microbial biochemical and hydrophysical 430 parameters of the soil, which are different for the two cultivars—confirm the importance of 431 taking into account soil (and particularly rhizosheath soil) parameters when selecting culti-432 vars for specific agroclimatic contexts (Nazari et al., 2020, Cattivelli et al., 2008). Finally, this 433 study, suggests some links between aboveground and soil parameters in relation with water 434 deficit and plant water use efficiency. This underlines the need of getting a more integrated 435 and quantitative view of the interactions between soil microbial-biochemical parameters 436 and soil hydrophysical properties when investigating agro-ecological solutions to water re-437 lated yield gap.

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448 **References**

Ahmed, M.A., Passioura, J., Carminati, A., 2018. Hydraulic processes in roots and the
rhizosphere pertinent to increasing yield of water-limited grain crops: a critical
review. Journal of Experimental Botany 69, 3255–3265.
https://doi.org/10.1093/jxb/ery183

Alami, Y., Achouak, W., Marol, C., Heulin, T., 2000. Rhizosphere Soil Aggregation and Plant
Growth Promotion of Sunflowers by an Exopolysaccharide-Producing Rhizobiumsp.
Strain Isolated from Sunflower Roots. Appl. Environ. Microbiol. 66, 3393–3398.
https://doi.org/10.1128/AEM.66.8.3393-3398.2000

Anderson, J. P. E., and K. H. Domsch. 1978. "A Physiological Method for the Quantitative
Measurement of Microbial Biomass in Soils." Soil Biology and Biochemistry 10 (3):
215–21. https://doi.org/10.1016 /0038-0717(78)90099-8.

460 Angers, D.A., Caron, J., 1998. Plant-induced changes in soil structure : Processes and 461 feedbacks. Biogeochemistry 42, 55–72. https://doi.org/10.1023/A:1005944025343

462 Aravena, J.E., Berli, M., Ghezzehei, T.A., Tyler, S.W., 2011. Effects of Root-Induced
463 Compaction on Rhizosphere Hydraulic Properties - X-ray Microtomography Imaging
464 and Numerical Simulations. Environ. Sci. Technol. 45, 425–431.
465 https://doi.org/10.1021/es102566j

Arbex de Castro Vilas Boas, A., Page, D., Giovinazzo, R., Bertin, N., Fanciullino, A.-L., 2017.
Combined Effects of Irrigation Regime, Genotype, and Harvest Stage Determine
Tomato Fruit Quality and Aptitude for Processing into Puree. Front. Plant Sci. 8, 1725.
https://doi.org/10.3389/fpls.2017.01725

Bardgett, R.D., Bowman W.D., Kaufmann R. and Schmidt S.K., 2005. A temporal approach to
linking aboveground and belowground ecology. Trends in Ecology & Evolution 20,
634-641. https://doi.org/10.1016/j.tree.2005.08.005

Bardgett, R.D., Mommer, L., De Vries, F.T., 2014. Going underground: root traits as drivers of
ecosystem processes. Trends in Ecology & Evolution 29, 692–699.
https://doi.org/10.1016/j.tree.2014.10.006

476 Bengough, A.G., 2012. Water Dynamics of the Root Zone: Rhizosphere Biophysics and Its

477 Control on Soil Hydrology. Vadose Zone Journal 11, vzj2011.0111.
478 https://doi.org/10.2136/vzj2011.0111

Ben Sassi, M., Dollinger, J., Renault, P., Tlili, A., Bérard, A., 2012. The FungiResp method: An
application of the MicroRespTM method to assess fungi in microbial communities as
soil biological indicators. Ecological Indicators 23, 482–490.
https://doi.org/10.1016/j.ecolind.2012.05.002

Benard, Pascal, Mohsen Zarebanadkouki, Mathilde Brax, Robin Kaltenbach, Iwan Jerjen,
Federica Marone, Estelle Couradeau, Vincent J. M. N. L. Felde, Anders Kaestner, and
Andrea Carminati. 2019. "Microhydrological Niches in Soils: How Mucilage and EPS
Alter the Biophysical Properties of the Rhizosphere and Other Biological Hotspots."
Vadose Zone Journal 18 (1): 180211. https://doi.org/10.3929/ethz-b-000393395

488 Bérard, A., Bouchet, T., Sévenier, G., Pablo, A.L., Gros, R., 2011. Resilience of soil microbial
489 communities impacted by severe drought and high temperature in the context of
490 Mediterranean heat waves. European Journal of Soil Biology 47, 333–342.
491 https://doi.org/10.1016/j.ejsobi.2011.08.004

492 Bérard, A., Ben Sassi, M., Kaisermann, A., Renault, P., 2015. Soil microbial community
493 responses to heat wave components: drought and high temperature. Climate
494 Research 66, 243–264. https://doi.org/10.3354/cr01343

Boeck, H.J.D., Dreesen, F.E., Janssens, I.A., Nijs, I., 2011. Whole-system responses of
experimental plant communities to climate extremes imposed in different seasons.
New Phytologist 189, 806–817. https://doi.org/10.1111/j.1469-8137.2010.03515.x

Bowles, T.M., Barrios-Masias, F.H., Carlisle, E.A., Cavagnaro, T.R., Jackson, L.E., 2016. Effects
of arbuscular mycorrhizae on tomato yield, nutrient uptake, water relations, and soil
carbon dynamics under deficit irrigation in field conditions. Science of The Total
Environment 566–567, 1223–1234. https://doi.org/10.1016/j.scitotenv.2016.05.178

502 Carminati, A., Vetterlein, D., 2013. Plasticity of rhizosphere hydraulic properties as a key for
 503 efficient utilization of scarce resources. Annals of Botany 112, 277–290.
 504 https://doi.org/10.1093/aob/mcs262

505 Cattivelli, L., Rizza, F., Badeck, F.-W., Mazzucotelli, E., Mastrangelo, A.M., Francia, E., Marè,
 506 C., Tondelli, A., Stanca, A.M., 2008. Drought tolerance improvement in crop plants:

507An integrated view from breeding to genomics. Field Crops Research 105, 1–14.508https://doi.org/10.1016/j.fcr.2007.07.004

Cavagnaro T.R., Jackson L.E., Six J., Ferris H., Goyal S., Asami D., Scow K.M., 2006. Arbuscular
 mycorrhizas, microbial communities, nutrient availability, and soil aggregates in
 organic tomato production. Plant Soil. 282(1):209–225. doi: 10.1007/s11104-005 5847-7

513 Chapman, Stephen J., Colin D. Campbell, and Rebekka R. E. Artz. 2007. "Assessing CLPPs 514 Using MicroRespTM." Journal of Soils and Sediments 7 (6): 406–10. 515 ttps://doi.org/10.1065/jss2007.10.259.

516 Chen, L., Rossi, F., Deng, S., Liu, Y., Wang, G., Adessi, A., De Philippis, R., 2014. 517 Macromolecular and chemical features of the excreted extracellular polysaccharides 518 in induced biological soil crusts of different ages. Soil Biology and Biochemistry 78, 1–

519 9. https://doi.org/10.1016/j.soilbio.2014.07.004

520 Chenu, C., Roberson, E.B., 1996. Diffusion of glucose in microbial extracellular polysaccharide
521 as affected by water potential. Soil Biology and Biochemistry 28, 877–884.
522 https://doi.org/10.1016/0038-0717(96)00070-3

523 Cornu, P., Valceschini Egizio, Mght-Bournay, O., 2018. Histoire de l'INRA, entre science et po-524 litique, Quae. ed. 78026 Versaille Cedex, France

525 Costa, J. Miguel, Maria F. Ortuño, and M. Manuela Chaves. 2007. "Deficit Irrigation as a
526 Strategy to Save Water: Physiology and Potential Application to Horticulture." Journal
527 of Integrative Plant Biology 49 (10): 1421–34. https://doi.org/10.1111/j.1672528 9072.2007.00556.x

529 Crouzet, O., Consentino, L., Pétraud, J.-P., Marrauld, C., Aguer, J.-P., Bureau, S., Le 530 Bourvellec, C., Touloumet, L., Bérard, A., 2019. Soil Photosynthetic Microbial 531 Communities Mediate Aggregate Stability: Influence of Cropping Systems and 532 Herbicide Use in an Agricultural Soil. Front. Microbiol. 10, 1319. 533 https://doi.org/10.3389/fmicb.2019.01319

534 Czarnes S., Hallett P. D., Bengough A. G., Young I.M., 2000. Root- and microbial-derived
535 mucilages affect soil structure and water transport. European Journal of Soil Science
536 51: 435-443.

537 De León-González, F., Gutiérrez-Castorena, M.C., González-Chávez, M.C.A., Castillo-Juárez,
538 H., 2007. Root-aggregation in a pumiceous sandy soil. Geoderma 142, 308–317.
539 https://doi.org/10.1016/j.geoderma.2007.08.023

540 Dormann, C.F., Elith, J., Bacher, S., Buchmann, C., Carl, G., Carré, G., Marquéz, J.R.G., Gruber, 541 B., Lafourcade, B., Leitão, P.J., Münkemüller, T., McClean, C., Osborne, P.E., 542 Reineking, B., Schröder, B., Skidmore, A.K., Zurell, D., Lautenbach, S., 2013. 543 Collinearity: a review of methods to deal with it and a simulation study evaluating 544 https://doi.org/10.1111/j.1600their performance. Ecography 36, 27–46. 545 0587.2012.07348.x

546 Doussan, C., Cousin, I., Berard, A., Chabbi, A., Legendre, L., Czarnes, S., Toussaint, B., Ruy, S.,
547 2015. Crop systems and plant roots can modify the soil water holding capacity 17,
548 9285.

549 Duan, H., Wu, J., Huang, G., Zhou, S., Liu, W., Liao, Y., Yang, X., Xiao, Z., Fan, H., 2016.
550 Individual and interactive effects of drought and heat on leaf physiology of seedlings
551 in an economically important crop. AoB PLANTS plw090.
552 https://doi.org/10.1093/aobpla/plw090

Dubois, M., Gilles, K.A., Hamilton, J.K., Rebers, P.A., Smith, F., 1956. Colorimetric Method for
 Determination of Sugars and Related Substances | Analytical Chemistry [WWW
 Document]. URL https://pubs.acs.org/doi/pdf/10.1021/ac60111a017 (accessed
 8.18.20).

557 FAOSTAT [WWW Document], n.d. URL http://www.fao.org/faostat/en/#data (accessed 558 5.11.20).

Fang, Y., Xiong, L. General mechanisms of drought response and their application in drought
resistance improvement in plants. Cell. Mol. Life Sci. 72, 673–689 (2015).
https://doi.org/10.1007/s00018-014-1767-0

Freschet, G. T., Roumet, C., Comas, L. H., Weemstra, M., Bengough, A. G., Rewald, B.,
Bardgett, R. D., De Deyn, G., Johnson, D., Klimešo-vá, J., Lukac, M., McCormack, M. L.,
Meier, I. C., Pagès, L., Poorter, H., Prieto, I., Wurzburger, N., Zadworny, M.,
Bagniewska-Zadworna, A., ... Stokes, A., 2021. Root traits as drivers of plant and
ecosystem function-ing: Current understanding, pitfalls and future research needs.

567 New Phytologist. In press

Göttlein, A., 2006. Sampling of rhizosphere soil and collection of rhizosphere soil solution. In:
Luster, J., Finlay, R. (Eds.), Handbook of Methods Used in Rhizosphere Research.
Swiss Federal Research Institute WSL, Birmensdorf, pp. 25–29.

Hallett, P.D., Feeney, D.S., Bengough, A.G., Rillig, M.C., Scrimgeour, C.M., Young, I.M., 2009.
Disentangling the impact of AM fungi versus roots on soil structure and water
transport. Plant Soil 314, 183–196. https://doi.org/10.1007/s11104-008-9717-y

Hegyi, G., Garamszegi, L.Z., 2011. Using information theory as a substitute for stepwise
regression in ecology and behavior. Behav Ecol Sociobiol 65, 69–76.
https://doi.org/10.1007/s00265-010-1036-7

577 Henry A., Doucette W., Norton J., Bugbee B., 2007. Changes in Crested Wheatgrass Root
578 Exudation Caused by Flood, Drought, and Nutrient Stress. Journal of Environmental
579 Quality 36, 904-912.

Hinsinger, P., Bengough, A.G., Vetterlein, D. et al. Rhizosphere: biophysics, biogeochemistry
and ecological relevance. Plant Soil 321, 117–152 (2009).
https://doi.org/10.1007/s11104-008-9885-9

583 Jia, G., E. Shevliakova, P. Artaxo, N. De Noblet-Ducoudré, R. Houghton, J. House, K. Kitajima, 584 C. Lennard, A. Popp, A. Sirin, R. Sukumar, L. Verchot (2019). Land-climate 585 interactions. In: Climate Change and Land: an IPCC special report on climate change, 586 desertification, land degradation, sustainable land management, food security, and 587 greenhouse gas fluxes in terrestrial ecosystems [P.R. Shukla, J. Skea, E. Calvo Buendia, 588 V. Masson-Delmotte, H.-O. Pörtner, D.C. Roberts, P. Zhai, R. Slade, S. Connors, R. van 589 Diemen, M. Ferrat, E. Haughey, S. Luz, S. Neogi, M. Pathak, J. Petzold, J. Portugal 590 Pereira, P. Vyas, E. Huntley, K. Kissick, M, Belkacemi, J. Malley, (eds.)]. In press.

591Łabędzki L., 2016, Actions and measures for mitigation drought and water scarcity in592agriculture. Journal of Water and Land Development 29, 3-10.

593 Lê, S., Josse, J. & Husson, F. (2008). FactoMineR: An R Package for Multivariate Analysis.
594 Journal of Statistical Software. 25(1). pp. 1-18.

595 Lipiec, J., Doussan, C., Nosalewicz, A., Kondracka, K., 2013. Effect of drought and heat

596 stresses on plant growth and yield: a review. International Agrophysics 27, 463–477.

597 https://doi.org/10.2478/intag-2013-0017Lynch, J.M., Whipps, J.M., 1990. Substrate

flow in the rhizosphere. Plant Soil 129, 1–10. https://doi.org/10.1007/BF00011685

Marasco, R., Mosqueira, M.J., Fusi, M. et al., 2018.Rhizosheath microbial community
assembly of sympatric desert speargrasses is independent of the plant host.
Microbiome 6, 215. https://doi.org/10.1186/s40168-018-0597-y

Murtaugh, P.A., 2009. Performance of several variable-selection methods applied to real
ecological data. Ecology Letters 12, 1061–1068. https://doi.org/10.1111/j.14610248.2009.01361.x

Nazari M., 2021. Plant mucilage components and their functions in the rhizosphere.
Rhizosphere 18, 100344. https://doi.org/10.1016/j.rhisph.2021.100344

Nazari M, Riebeling S, Banfield CC, Akale A, Crosta M, Mason-Jones K, Dippold MA and
Ahmed MA, 2020. Mucilage Polysaccharide Composition and Exudation in Maize
From Contrasting Climatic Regions. Front. Plant Sci. 11:587610. https://doi:
10.3389/fpls.2020.587610

611 Pang, J., Ryan, M.H., Siddique, K.H.M. et al., 2017. Unwrapping the rhizosheath. Plant Soil
612 418, 129–139. https://doi.org/10.1007/s11104-017-3358-y

Patanè, C., Cosentino, S.L., 2010. Effects of soil water deficit on yield and quality of
processing tomato under a Mediterranean climate. Agricultural Water Management
97, 131–138. https://doi.org/10.1016/j.agwat.2009.08.021

616 Personeni E., Nguyen C., Marchal P., Pagès L., 2007. Experimental evaluation of an efflux–
617 influx model of C exudation by individual apical root segments. Journal of
618 Experimental Botany, 58, 8: 2091–2099. doi:10.1093/jxb/erm065

619 Poirier V, Roumet C, Munson AD. 2018. The root of the matter: linking root traits and soil 620 organic matter stabilization processes. Soil Biology and Biochemistry 120: 246–259.

621 Rasse, D.P., Smucker, A.J.M., Santos, D., 2000. Alfalfa Root and Shoot Mulching Effects on

Soil Hydraulic Properties and Aggregation. Soil Sci. Soc. Am. J. 64, 725–731.
https://doi.org/10.2136/sssaj2000.642725x

624 R Core Team (2017). R: A language and environment for statistical computing. R Foundation

625 for Statistical Computing, Vienna, Austria. URL https://www.R-project.org/.

Redmile-Gordon, M.A., Brookes, P.C., Evershed, R.P., Goulding, K.W.T., Hirsch, P.R., 2014.
Measuring the soil-microbial interface: Extraction of extracellular polymeric
substances (EPS) from soil biofilms. Soil Biology and Biochemistry 72, 163–171.
https://doi.org/10.1016/j.soilbio.2014.01.025

Rinaldi, M., Ventrella, D., Gagliano, C., 2007. Comparison of nitrogen and irrigation strategies
in tomato using CROPGRO model. A case study from Southern Italy. Agricultural
Water Management 87, 91–105. https://doi.org/10.1016/j.agwat.2006.06.006

Ripoll, J., Urban, L., Brunel, B., Bertin, N., 2015. Water deficit effects on tomato quality
depend on fruit developmental stage and genotype. Journal of plant physiology 190,
26-35. https://doi.org/10.1016/j.jplph.2015.10.006

Roberson, E.B., Firestone, M.K., 1992. Relationship between Desiccation and
Exopolysaccharide Production in a Soil Pseudomonas sp. Appl. Environ. Microbiol. 58,
1284–1291.

RStudio Team (2020). RStudio: Integrated Development for R. RStudio, PBC, Boston, MA URL
 http://www.rstudio.com/.

Schielzeth, H., 2010. Simple means to improve the interpretability of regression coefficients.
Methods in Ecology and Evolution 1, 103–113. https://doi.org/10.1111/j.2041210X.2010.00012.x

Sher, Y., Baker, N.R., Herman, D., Fossum, C., Hale, L., Zhang, X.-X., Nuccio, E., Saha, M.,
Zhou, J., Pett-Ridge, J., Firestone, M., 2019. Microbial extracellular polysaccharide
production and aggregate stability controlled by Switchgrass (Panicum virgatum) root
biomass and soil water potential. bioRxiv 724195. https://doi.org/10.1101/724195

van Veelen, A., Tourell, M.C., Koebernick, N., Pileio, G., Roose, T., 2018. Correlative
Visualization of Root Mucilage Degradation Using X-ray CT and MRI. Front. Environ.
Sci. 6. https://doi.org/10.3389/fenvs.2018.00032

Whittingham, M.J., Stephens, P.A., Bradbury, R.B., Freckleton, R.P., 2006. Why do we still use
stepwise modelling in ecology and behaviour? Journal of Animal Ecology 75, 1182–
1189. https://doi.org/10.1111/j.1365-2656.2006.01141.x

Zhang, Xuechen, Yakov Kuzyakov, Huadong Zang, Michaela A. Dippold, Lingling Shi, Sandra
Spielvogel, and Bahar S. Razavi. 2020. "Rhizosphere Hotspots: Root Hairs and
Warming Control Microbial Efficiency, Carbon Utilization and Energy Production." Soil
Biology and Biochemistry 148 (September): 107872.
https://doi.org/10.1016/j.soilbio.2020.107872

Zuluaga M.Y.A., Lima Milani K.M., Azeredo Gonçalves L.S., Martinez de Oliveira A.L., 2020.
Diversity and plant growth-promoting functions of diazotrophic/N-scavenging
bacteria isolated from the soils and rhizospheres of two species of Solanum. PLoS
ONE 15(1): e0227422. https://doi.org/10.1371/journal.pone.0227422

Figures and table

Table of abbreviations

Abbreviation	Description	Unit			
Α	Soil sampling date at the beginning of fructification stage (August 2. 2018).	-			
AIC	Akaike Information Criterion	-			
B (Bulk)	Soil sampling from bulk soil (without roots)	-			
CoY	Fresh Commercial Tomato Yield (including red and yellow tomato)	kg/ha			
DMY	Commercial Tomato Yield in Dry Matter (including red and yellow tomato)	kg/ha			
EPSac	ExoPolySaccharides	mg eq glc / g DrySoil			
F.Ala	Alanine inducted respiration for fungi communities in soil	CO2/gDrySoil/h			
F.Bas	Basal respiration for fungi communities in soil	CO2/gDrySoil/h			
F.Cell	Cellulobiose inducted respiration for fungi communities in soil	CO2/gDrySoil/h			
F.Gly	Glycine inducted respiration for fungi communities in soil	CO2/gDrySoil/h			
F.GlcN	Glucosamine inducted respiration for fungi communities in soil	CO2/gDrySoil/h			
F.Mal	Malate inducted respiration for fungi communities in soil	CO2/gDrySoil/h			
F.Tre	Trehalose inducted respiration for fungi communities in soil	CO2/gDrySoil/h			
н	Soil sampling from Heinz 1015 tomato cultivar	-			
l	Soil sampling date at the beginning of flowering stage (June 19. 2018).	-			
M.Ala	Alanine inducted respiration for fungi communities in soil	CO2/gDrySoil/h			
M.Bas	Basal respiration for fungi communities in soil	CO2/gDrySoil/h			
M.Cell	Cellulobiose inducted respiration for fungi communities in soil	CO2/gDrySoil/h			
M.Gly	Glycine inducted respiration for fungi communities in soil	CO2/gDrySoil/h			
M.GlcN	Glucosamine inducted respiration for fungi communities in soil	CO2/gDrySoil/h			
M.Mal	Malate inducted respiration for fungi communities in soil	CO2/gDrySoil/h			
M.Tre	Trehalose inducted respiration for fungi communities in soil	CO2/gDrySoil/h			
R (Rhizo)	Soil sampling from rhizosheath soil (root adhering)	-			
RoY	Rotten Tomato Yield	t/ha			
т	Soil sampling from Terradou tomato cultivar	-			
VIF	Variable Inflation Factor	-			
W0.3b	Water Retention Capacity at 0.3b (at Field Capacity)	g/g			
W15b	Water Retention Capacity at 15b (at Permanent Wilting Point)	g/g			
WD	Water deficit irrigation (57% of ETP losses)	-			
WUE	Plant Water Use Efficiency	kg/m3			
ww	Well watered irrigation (87% of ETP losses)	-			

Localisation	44°11′N/4°48′E
Texture	loam
Clay (g kg ⁻¹)	290
Loam (g kg ⁻¹)	331
Sand (g kg ⁻¹)	379
SOC (g kg ⁻¹)	12.5
TN (g kg ⁻¹)	1.1
C/N	11
CaCO3 (g kg ⁻¹)	484
Ca ²⁺ (cmol+ kg ⁻¹)	14.6
Mg ²⁺ (cmol+ kg ⁻¹)	0.595
CEC (cmol+ kg ⁻¹)	12.7
pH (water)	8.5
WSA (%)	36

Table 1: Main physical, chemical and microbial characteristics of the soils used for this study on the Piolenc experimental site. SOC: Soil Organic Carbon, TN: Total Nitrogen, C/N: carbon over nitrogen ratio CEC: Cation Exchange Capacity, WSA: Water Stable Aggregates

	W0.3b	W15b	EPSac	F.Glc	F.Tre	F.Cell	F.GlcN	F.Ala	F.Gly	F.Mal	FB	M.Bas	M.Glc	M.Tre	M.Cell	M.GlcN	M.Ala	M.Gly	M.Mal	MB
Global																				
SOIL	R>B	R>B***	R>B	R>B***	R>B**	R>B***	R>B***	R>B***	R>B***	R>B***	R>B***	R>B**	R>B***							
CULT	T>H	H>T	T>H	T>H	T>H	T>H	T>H	H>T	T>H	H>T	T>H**	T>H	T>H*	T>H**	T>H*	T>H	T>H	T>H	H>T	T>H*
SEAS	J>A	A>J	A>J***																	
SOIL:CULT	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-
SOIL:SEAS	***	-	-	-	-	-	***	*	-	-	-	-	-	-	*	-	-	-	-	-
CULT:SEAS	-	*	**	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	*	-
Global Rhizo																				
CULT	T>H	H>T	T>H	T>H***	T>H	T>H	T>H	H>T	H>T	T>H	T>H**	T>H	T>H	T>H*	T>H	T>H	T>H	H>T	H>T	T>H
SEAS	J>A***	A>J**	A>J***	A>J*	A>J***															
CULT:SEAS	-	*	-	-	-	-	-	*	-	-	-	-	-	-	-	-	-	*	**	-
June																				
SOIL	R>B***	R>B	R>B**	R>B***	R>B	R>B*	R>B	R>B	R>B	R>B*	R>B***	R>B	R>B	R>B**	R>B*	R>B	R>B*	R>B*	R>B**	R>B
CULT	H>T	T>H	H>T	T>H																
SOIL:CULT	-	-	-	*	-	-	-	-	-	-	**	-	-	-	-	-	-	-	-	-
June Rhizo																				
CULT	H>T	T>H	T>H	T>H*	T>H	T>H	T>H	T>H	T>H	T>H	T>H*	T>H								
August WW																				
SOIL	B>R	R>B*	R>B	R>B**	R>B*	R>B*	R>B***	R>B**	R>B**	R>B*	R>B**	R>B*	R>B***	R>B***	R>B***	R>B***	R>B**	R>B***	R>B***	R>B***
CULT	T>H	H>T*	T>H*	T>H	T>H	H>T	T>H	H>T	H>T	H>T	T>H	T>H	T>H	T>H	T>H	T>H	H>T	H>T	H>T	T>H
SOIL:CULT	-	-	-	-	-	-	-	*	-	-	-	-	-	-	*	-	**	*	-	-
August WW Rhizo																				
CULT	T>H	H>T	T>H	T>H	H>T	H>T	T>H	H>T*	H>T	H>T	H>T	H>T	H>T	T>H	H>T	H>T	H>T	H>T**	H>T**	H>T
August WD																				
SOIL	B>R**	R>B***	R>B	R>B	R>B	R>B	R>B***	R>B***	R>B***	R>B*	R>B	R>B*	R>B***							
CULT	T>H***	T>H	T>H	T>H*	T>H	T>H	T>H*	H>T	H>T	T>H	T>H*	T>H	H>T	T>H						
SOIL:CULT	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-
August WD Rhizo																				
CULT	T>H*	H>T	T>H	T>H*	T>H	T>H	T>H	T>H	H>T	T>H	T>H*	T>H	T>H	T>H	T>H	T>H	T>H	H>T	H>T	H>T

Table 2: Analyses of variance (ANOVAs) results for the soil parameters for the three factors : the type of soil rhizosheath or bulk (SOIL), the season (SEAS) and the cultivar (CULT). Main effect and their interactions are calculated on global dataset (whatever the rhizosheath or bulk soil, the season, the cultivar) and on rhizosheath soil dataset. Bellow are analysis for June, August-WD irrigation and August-WW irrigation dataset. X>Y signifies that the parameter X is greater than Y. Level of significance: *p<0.05; **p<0.01;***p<0.001. See table of abbreviations for more information.

Datasat	Factor	D ²	n_value	Sign.
Dalasel	Factor	N	p-value	code
Global	MB	0.20	4.09E-05	* * *
n=76	FB	0.32	1.28E-07	* * *
Rhizo	MB	0.16	7.39E-03	**
n=38	FB	0.19	3.53E-03	**
Bulk	MB	0.21	2.38E-03	**
n=38	FB	0.39	1.71E-05	* * *
June	MB	-0.02	6.12E-01	
n=38	FB	0.22	1.60E-03	**
June rhizo	MB	-0.06	9.63E-01	
n=19	FB	0.11	9.07E-02	•
June bulk	MB	-0.03	4.79E-01	
n=19	FB	5E-4	3.29E-01	
August	MB	5E-4	3.19E-01	
n=38	FB	-0.03	7.83E-01	
August rhizo	MB	-0.06	8.48E-01	
n=19	FB	-0.03	5.05E-01	
August bulk	MB	0.07	1.50E-01	
n=19	FB	-0.005	3.54E-01	

Table 3: Results of linear regression between EPSac and the microbial or fungal biomasses (MB, FB). Considering global soil data or subsets of the soil data. a: slope ; b: intercept of linear regression. Level of significance: *p<0,05; **p<0,01;***p<0,001. See table of abbreviations for more information. See table of abbreviations for more information.

Variables selected	AIC	VIF	Final AIC	Final p- value	Signif. Code	Final R ²
W0.3b global			-445.06	3.69E-04	***	0.21
F.Ala	-433.78	5				
M.Ala	-441.62	3.06				
F.Cell	-444.49	5.43				
EPSac	-445.06	1.73				
W15b global			-562.14	6.14E-05	***	0.21
F.Tré	-548.36	3.67				
M.Tré	-550.61	3.67				
W0.3b June			-215.13	7.91E-03	**	0.23
EPSac	-212.62	1.13				
F.Cell	-212.71	5.09				
F.Tré	-215.13	4.87				
W15b June			-284.35	8.24E-03	**	0.16
M.Cell	-284.35	nn				
W0.3b August-WW			-122.85	4.15E-03	**	0.51
FB	-112.45	1.66				
M.Cell	-114.28	2.73				
F.Ala	-122.85	1.86				
W15b August-WW			-120.62	1.04E-02	*	0.3
F.Ala	-120.62	nn				
W0.3b August-WD			-129.78	3.94E-04	***	0.65
F.Ala	-115.33	2.61				
EPSac	-119.84	1.07				
M.Tré	-123.31	1.98				
F.Gly	-129.78	2.99				
W15b August-WS			-164.75	1.50E-05	***	0.74
F.D.Glc	-157.97	5.2				
M.Gly	-161.08	2.94				
M.D.Glc	-164.75	7.1				

Table 4: Multiple linear regression model between water retention (W0.3b and W15b) and both physico-chemical (EPSac) and microbial parameters (catabolic profile, biomass) for the global dataset and subsets. Best model issued from stepwise regression using AIC selection criteria. The final model include variable with VIF<10. restricted to a maximum 4 explanatory variables. Level of significance: *p<0.05; **p<0.01;***p<0.001. See table of abbreviations for more information.



Figure 1: Principal Component Analysis (PCA) of fungal catabolic profile of the rhizosheath and bulk soil data. Projection on axes 1 and 2 (73% of variance). Distribution of individuals and correlations shown for cultivar (H and T). soil type (R and B) and sampling season (J and A).



Figure 2: Principal Component Analysis (PCA) of microbial catabolic profile of the rhizosheath and bulk soil data. Projection on axes 1 and 2 (> 64% variance). Microbial catabolic profile for June. August-WW and August-WD data. Distribution of individuals and correlations shown for cultivar (H and T).

2-c: August-WD





3-c: August-WD



3-b: August-WW



Figure 3: Principal Component Analysis (PCA) of fungal catabolic profile for June. August-WW and August-WD data. Projection on axes 1 and 2 (> 65% variance). Distribution of individuals and correlations shown for soil types (R and B).



Fig. 4: Linear regression between EPSac amount and the microbial or fungal biomasses (MB, FB) for the global soil data set.



Figure 5: Box plots comparing H and T varieties for the WW irrigation dataset (Fig.5-a), and the WD irrigation dataset (Fig.5-b), with Rotten tomato Yield (RoY, in t ha-1), fresh Commercial Yield (CoY, in t ha-1) and Dry Matter Yield (DMY, in 10 kg ha-1) and Water Use Efficiency (WUE, in kg m–3). Level of significance: *p<0.05. See table of abbreviations for more information.