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**This is the author's final version of the contribution published as:**

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Not only priming: Soil microbiota may protect tomato from root pathogens

Plant Signaling & Behavior, VOL. 13, NO. 8, e1464855, 2018, (3 pages), 10.1080/15592324.2018.1464855

**The publisher's version is available at:**

<https://www.tandfonline.com/doi/full/10.1080/15592324.2018.1464855>

**When citing, please refer to the published version.**

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**2 2 Not Only Priming: Soil Microbiota May Protect Tomato from Root Pathogens**

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25 **Submitted:** 28 March 2018

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27 **Accepted:**

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31 12 **Keywords:** *Fusarium oxysporum* f. sp. *lycopersici*; arbuscular mycorrhizal fungi; defence

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34 13 responses; lignin biosynthesis; microbiota; suppressive and conducive soils; susceptible and

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36 14 resistant genotypes; tomato; gene expression.

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41 16 Correspondence to:

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44 17 Paola Bonfante

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50 20 **Abstract**

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52 21 An increasing number of studies have investigated soil microbial biodiversity. However, the

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54 22 mechanisms regulating plant responses to soil microbiota are largely unknown. A previous work

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56 23 tested the hypothesis that tomato plants grown on native soils with their complex microbiotas

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24 respond differently from tomato growing in a sterile substrate. Two soils, suppressive or conducive  
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225 to *Fusarium oxysporum* f. sp. *lycopersici* (FOL), and two genotypes susceptible and resistant to the  
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426 same pathogen were considered. The work highlighted that the two tested soil microbiotas,  
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727 irrespectively of their taxonomic composition, elicit the PAMP-triggered Immunity Pathway, the  
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928 first level of plant defence, as well as an increased lignin synthesis, leading to an active protection  
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1229 when FOL is present in the soil. Here, we tested the expression of a panel of genes involved in  
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1430 Effector-Triggered Immunity (ETI), demonstrating that soil microbiota, beside genotype, affects  
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1731 plant resistance to FOL also modulating this pathway.

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## 21 2233 **TEXT**

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2434 Next-generation sequencing (NGS) has enabled in-depth investigations of the microbial  
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2635 communities associated with animals, plants, and fungi. The awareness that multicellular  
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2936 eukaryotes host thousands of microbes, many beneficial, some essential and only a few deleterious  
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3137 has led to a paradigm shift in our knowledge of microbial–eukaryote interactions. NGS approaches  
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3438 helped us to reply to basic questions of traditional microbiology, as: ‘Which are the microbes  
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3639 thriving in that niche?’, and ‘What are they doing?’. Focusing on the plant side and starting from  
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3940 the pioneering researches by Bulgarelli et al.<sup>1</sup> and Lundberg et al.<sup>2</sup>, many other studies revealed the  
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4141 extraordinary diversity of microbes present on both roots, shoots, leaves, fruits<sup>3,4</sup>, and demonstrated  
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4442 how different parameters affect the composition of the microbiota: plant genotype, soil features,  
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4643 environmental parameters<sup>5,6</sup>. Interestingly, the environment resulted to be the driving force also for  
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4944 human microbiota, where it dominates over host genetics in shaping human gut microbiota<sup>7</sup>. The  
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5145 strict relationship existing between microbiota and their eukaryotic host has also led to the  
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5346 development of the *holobiont* concept<sup>8,9</sup>. Host-microbial systems, being a complex assembly of  
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5647 diverse organisms, constitute unique biological entities, defined as ‘meta-organisms’ or holobionts<sup>10</sup>.  
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5848 However, metagenomic sequencing has only given indirect responses to the questions opened by  
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49 these new scenarios: ‘How the host responds to its extended microbiota, which represents its second  
1 genome?’.  
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752 Chialva et al.<sup>11</sup> focused on tomato (*Solanum lycopersicum*), testing the hypothesis that plants grown  
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1053 on native soils display different responses to soil microbiotas. Using transcriptomics, proteomics,  
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1254 and biochemistry, the study has described the responses of two tomato genotypes (susceptible or  
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1455 resistant to FOL) grown on two native soils (conducive and suppressive to FOL) and an artificial  
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1756 substrate. Results showed that native soils, particularly the suppressive one, affect tomato responses  
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1957 by modulating pathways involved in responses to oxidative stress, phenol biosynthesis, lignin  
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2258 deposition, and PAMP-triggered Immunity (PTI). By contrast, in tomato plants grown on steam-  
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2459 disinfected soils, total phenols and PTI responses significantly decreased, suggesting a crucial role  
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2760 of soil microbiota in eliciting a priming effect. To validate those observations, the mycorrhizal  
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2961 fungus *Funnelliformis mosseae*, was selected as one of the most abundant AM fungi in both soils,  
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3262 and inoculated in tomato growing on steam-disinfected soils: the fungal inoculation partly rescued  
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3463 some of the local and systemic responses, which were identified as a part of the priming response.  
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3664 Martinez-Medina et al.<sup>12</sup> have neatly identified different conditions where plant defence priming  
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3965 takes place and have acknowledged many beneficial microbes as a source for priming stimuli.  
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4166 Indeed, under the tested experimental conditions (native soils vs sterile substrate), tomato activates  
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4467 several genes involved in PTI, such as those encoding for PR proteins, WRKY transcription factors,  
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4668 ROS burst signalling and calcium signalling, which are involved in immune response<sup>13</sup>. To  
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4969 understand whether such an adaptive measure leads the plant to an enhanced defence readiness<sup>11</sup>  
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5170 tomato plants were inoculated with FOL. As expected, reduced disease symptoms were detected in  
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5371 the resistant genotype ('Battito') in both soils; but surprisingly the susceptible genotype 'Cuore di  
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5672 Bue' was partially protected from FOL on the suppressive soil. However, it is still unknown whether  
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73 the Effector-Triggered Immunity (ETI), *i.e.* the second barrier against pathogens, responds to soil  
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274 microbiota.  
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776 Here, we hypothesized that the priming status raised in tomato by soil microbiota could elicit the  
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1077 expression of genes directly involved in ETI in the presence of FOL. With this aim, we selected a  
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1278 panel of genes involved in the ETI pathway (Table 1) and tested their expression by using RT-  
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1479 qPCR in FOL-inoculated plant roots according to the set-up and methods described in Chialva et  
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1780 al.<sup>11</sup>.  
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2282 Results indicate that soil microbiota promoted the ETI response of plants after FOL infection (Fig.  
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2483 1): while in RNA-seq experiment, where FOL was not present, ETI genes were not differentially  
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2784 expressed, in FOL-inoculated plants RT-qPCR experiment detected gene modulation<sup>11</sup>. Both  
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2985 genotypes significantly upregulated the expression of *RIN4* ( $p < 0.05$ ) in both native soils compared  
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3286 to the control substrate. This protein is a target of type III pili effector proteins (virulence factors)  
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3487 from bacterial pathogens and interacts with RPS2 and RPM1 R protein leading to hypersensitive  
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3688 response<sup>14,15</sup>. Moreover, we tested the expression of two previously described ETI-marker genes<sup>16</sup>  
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3989 and found that one of them coding for a UDP-glucosyltransferase family 1 protein (UDP) is  
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4190 upregulated in both soils ( $p < 0.05$ ) with the exception of the susceptible cultivar in the conducive  
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4491 soil. However, the other marker gene tested (UDP1) did not show differential expression across  
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4692 conditions. By contrast, the expression of the *I-2* R gene, directly involved in FOL race 2<sup>17</sup>, was  
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4993 upregulated only in the resistant genotype grown in the suppressive soil, while it remained  
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5194 consistent for the susceptible genotype in all the substrates. These results suggest a synergy between  
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5395 the genotype (presence of Resistance genes), the soil biological features, and – mechanistically –  
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5696 the ETI response. The 'Cuore di Bue' susceptible genotype has a more modulated response: FOL-  
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5897 suppressive soil with its microbiota activates the ETI response, while this action is not elicited in  
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98 the conducive soil. This well explains the modulation of *I-2 R* gene: to be activated, plant defences  
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299 require the suppressive soil microbiota acting on the resistant genotype, while the synergy between  
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5100 these two conditions is not satisfied in the susceptible genotype. The hypothesis may have an  
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8101 experimental validation by the presence of many bio-control *Fusaria* strains isolated in the  
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10102 Albenga soil<sup>18</sup>.

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14104 Our previous experiments demonstrated that soil microbiota leads to a priming ('state of alert') in  
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17105 tomato eliciting the PTI, which represents the first level of plant defence. When challenged by a  
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19106 pathogen, the alerted plant activates a new set of more specific genes related to the ETI, which is  
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22107 the second specific defence level (Fig. 2). This mechanism leads to a partial protection from the  
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24108 pathogen attack, even in the absence of specific resistance genes (as for the cultivar 'Cuore di Bue').

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26109 The modulation of the ETI-related genes indicates that native soil microbiota also affects plant  
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29110 response to FOL via ETI, in addition to the crucial role played by the genotype.

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31111 In conclusion, the investigation of the mechanisms operating in plants in native soils and in the  
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34112 presence of complex soil microbiota has revealed new unexpected responses. It seems that - just  
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36113 like humans - the tomato plant living in non-sterile conditions can better activate its immunity  
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39114 defence via the interaction with its microbiota.

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#### 42 43 44116 **Disclosure of potential conflicts of interest**

45  
46117 No potential conflicts of interest were disclosed.

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#### 50 51119 **Acknowledgments**

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53120 The authors thank the members of the Mycoplant Consortium for the support in FOL infection  
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56121 experiments.

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**Funding**

This research has received funding from the Mycoplant project (Root Microbiome for Plant Health: dissecting the role of soil fungi, D15E13000350003, Fondazione Compagnia di San Paolo Torino) and by the European Union’s Horizon 2020 research and innovation programme under grant agreement No 727929 (A novel and integrated approach to increase multiple and combined stress tolerance in plants using tomato as a model – TOMRES). M.C. was funded by a TOMRES fellowship.

**References**

1. Bulgarelli D, Rott M, Schlaeppi K, Ver Loren van Themaat E, Ahmadinejad N, Assenza F, Rauf P, Huettel B, Reinhardt R, Schmelzer E, et al. Revealing structure and assembly cues for Arabidopsis root-inhabiting bacterial microbiota. *Nature* 2012; 488:91–5.
2. Lundberg DS, Lebeis SL, Paredes SH, Yourstone S, Gehring J, Malfatti S, Tremblay J, Engelbrektson A, Kunin V, Rio TG del, et al. Defining the core Arabidopsis thaliana root microbiome. *Nature* 2012; 488:86–90.
3. Bai Y, Müller DB, Srinivas G, Garrido-Oter R, Potthoff E, Rott M, Dombrowski N, Münch PC, Spaepen S, Remus-Emsermann M, et al. Functional overlap of the Arabidopsis leaf and root microbiota. *Nature* 2015; 528:364–9.
4. Coleman-Derr D, Desgarenes D, Fonseca-Garcia C, Gross S, Clingenpeel S, Woyke T, North G, Visel A, Partida-Martinez LP, Tringe SG. Plant compartment and biogeography affect microbiome composition in cultivated and native Agave species. *New Phytol* 2016; 209:798–811.

5. Zgad Zaj R, Garrido-Oter R, Jensen DB, Koprivova A, Schulze-Lefert P, Radutoiu S. Root nodule symbiosis in *Lotus japonicus* drives the establishment of distinctive rhizosphere, root, and nodule bacterial communities. *Proc Natl Acad Sci* 2016; 113:E7996–8005.
6. Hamonts Kelly, Trivedi Pankaj, Garg Anshu, Janitz Caroline, Grinyer Jasmine, Holford Paul, Botha Frederik C., Anderson Ian C., Singh Brajesh K. Field study reveals core plant microbiota and relative importance of their drivers. *Environ Microbiol* 2018; 20:124–40.
7. Rothschild D, Weissbrod O, Barkan E, Kurilshikov A, Korem T, Zeevi D, Costea PI, Godneva A, Kalka IN, Bar N, et al. Environment dominates over host genetics in shaping human gut microbiota. *Nature* 2018; 555:210–5.
8. Bordenstein SR, Theis KR. Host Biology in Light of the Microbiome: Ten Principles of Holobionts and Hologenomes. *PLOS Biol* 2015; 13:e1002226.
9. Theis KR, Dheilly NM, Klassen JL, Brucker RM, Baines JF, Bosch TCG, Cryan JF, Gilbert SF, Goodnight CJ, Lloyd EA, et al. Getting the Hologenome Concept Right: an Eco-Evolutionary Framework for Hosts and Their Microbiomes. *mSystems* 2016; 1:e00028-16.
10. Rosenberg E, Koren O, Reshef L, Efrony R, Zilber-Rosenberg I. The role of microorganisms in coral health, disease and evolution. *Nat Rev Microbiol* 2007; 5:355–62.
11. Chialva M, Salvioli di Fossalunga A, Daghino S, Ghignone S, Bagnaresi P, Chiapello M, Novero M, Spadaro D, Perotto S, Bonfante P. Native soils with their microbiotas elicit a state of alert in tomato plants. *New Phytol* 2018;
12. Martinez-Medina A, Flors V, Heil M, Mauch-Mani B, Pieterse CMJ, Pozo MJ, Ton J, van Dam NM, Conrath U. Recognizing Plant Defense Priming. *Trends Plant Sci* 2016; 21:818–22.
13. Chiang Y-H, Coaker G. Effector Triggered Immunity: NLR Immune Perception and Downstream Defense Responses. *Arab Book* 2015; 13:e0183.



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14. Mackey D, Belkhadir Y, Alonso JM, Ecker JR, Dangl JL. Arabidopsis RIN4 is a target of the type III virulence effector AvrRpt2 and modulates RPS2-mediated resistance. *Cell* 2003; 112:379–89.
  15. Liu J, Elmore JM, Coaker G. Investigating the functions of the RIN4 protein complex during plant innate immune responses. *Plant Signal Behav* 2009; 4:1107–10.
  16. Pombo MA, Zheng Y, Fernandez-Pozo N, Dunham DM, Fei Z, Martin GB. Transcriptomic analysis reveals tomato genes whose expression is induced specifically during effector-triggered immunity and identifies the Epk1 protein kinase which is required for the host response to three bacterial effector proteins. *Genome Biol* 2014; 15. Available from: <https://www.ncbi.nlm.nih.gov/pmc/articles/PMC4223163/>
  17. Simons G, Groenendijk J, Wijbrandi J, Reijans M, Groenen J, Diergaarde P, Van der Lee T, Bleeker M, Onstenk J, de Both M, et al. Dissection of the fusarium I2 gene cluster in tomato reveals six homologs and one active gene copy. *Plant Cell* 1998; 10:1055–68.
  18. Poli A, Lazzari A, Prigione V, Voyron S, Spadaro D, Varese GC. Influence of plant genotype on the cultivable fungi associated to tomato rhizosphere and roots in different soils. *Fungal Biol* 2016; 120:862–72.

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**Figure Legends**

**Figure 1.**

**RT-qPCR relative expression levels of gene involved in ETI in tomato plants (*Solanum lycopersicum*) infected with *Fusarium oxysporum* f. sp. *lycopersici* (FOL).**

*Ubiquitin* gene was used as reference for RT-qPCR. Letters indicate statistically supported differences (Kruskal–Wallis test at  $P < 0.05$ ). Data are means  $\pm$  SE ( $n = 3$ ). AL, ‘Albenga’ suppressive soil; RO, ‘Rosta’ conducive soil; CONT, Control 'Neutral' soil. B, 'Battito' FOL-resistant genotype; C, 'Cuore di Bue' FOL-susceptible genotype. (A) *RIN4*, RPM1 interacting protein 4; (B) *I-2*, CC-NBS-LRR, resistance protein 1; (B,C) *UDP*, *UDP1*, UDP-glucosyltransferase family 1 proteins.

**Figure 2**

**Scheme of defence responses activated by tomato (*Solanum lycopersicum*) in the presence of a complex native soil microbiota.**

(1) According to the models proposed by Chialva et al.,<sup>11</sup> in native soils microbial-associated molecular patterns (MAMPs) such as flagellin (flg22) and chitin are perceived by tomato plant. Those events elicit the PTI pathway (Plant-triggered Immunity) as a first defence level with the activation of calcium signalling (CNGCs, cyclic nucleotide-gated channels; CaM/CaM-like (CML), calmodulin-like proteins; CDPKs, calcium-dependent protein kinases) and WRKY transcription factors. This brings to the downstream activation of pathogenesis-related proteins genes (PR), such

166 as PR1, and to cell-wall fortification and lignin synthesis. (2) Since PTI-related defence is elicited, a  
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267 “continuative priming” by soil microbiota components occurs, maintaining plant defence active. (3)  
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568 When plant is attacked by *Fusarium oxysporum* f. sp *lycopersici* (FOL) the plant is already primed  
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769 and activates stronger ETI (Effector-triggered Immunity) defence. In both genotypes, effectors are  
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1070 strongly perceived (e.g. by *RIN4*): only in the FOL-resistant one a specific resistance mediated by *I*-  
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1271 2 is activated leading to the activation of the downstream ETI responses (such as UDP  
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1572 upregulation). However, in the susceptible genotype even if *I*-2 upregulation was not observed,  
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1773 FOL-suppressive soil induced the activation of downstream ETI pathway with the upregulation of a  
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1974 marker UDP gene.

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**Table 1. Table of primers used in RT-qPCR experiment.**

Gene	Transcript ID	Forward primer (5'-3')	Reverse primer (5'-3')	Reference
RPM1	Solyc11g0120	TCCTTCTGTAGAGTCGG	TCTTCTTCGTCGTGTTG	<sup>11</sup>
interacting protein 4 ( <i>RIN4</i> )	10.1	GCCA	GTTGGT	
CC-NBS-LRR, resistance protein 1 ( <i>I-2</i> )	Solyc11g0714	TTTGAAAGGGTCCCAA	TGCAGAGGGGTGTCAA	This study
UDP-glucosyltransferase family 1 protein (UDP)	Solyc10g0858	CAAAGCTGAAAGAGGG	TAACCCAAGCCCTAGCT	This study
UDP-	Solyc09g0925	GGTGCAACCCCATGTC	ATCAGAGAATGCCGCC	This

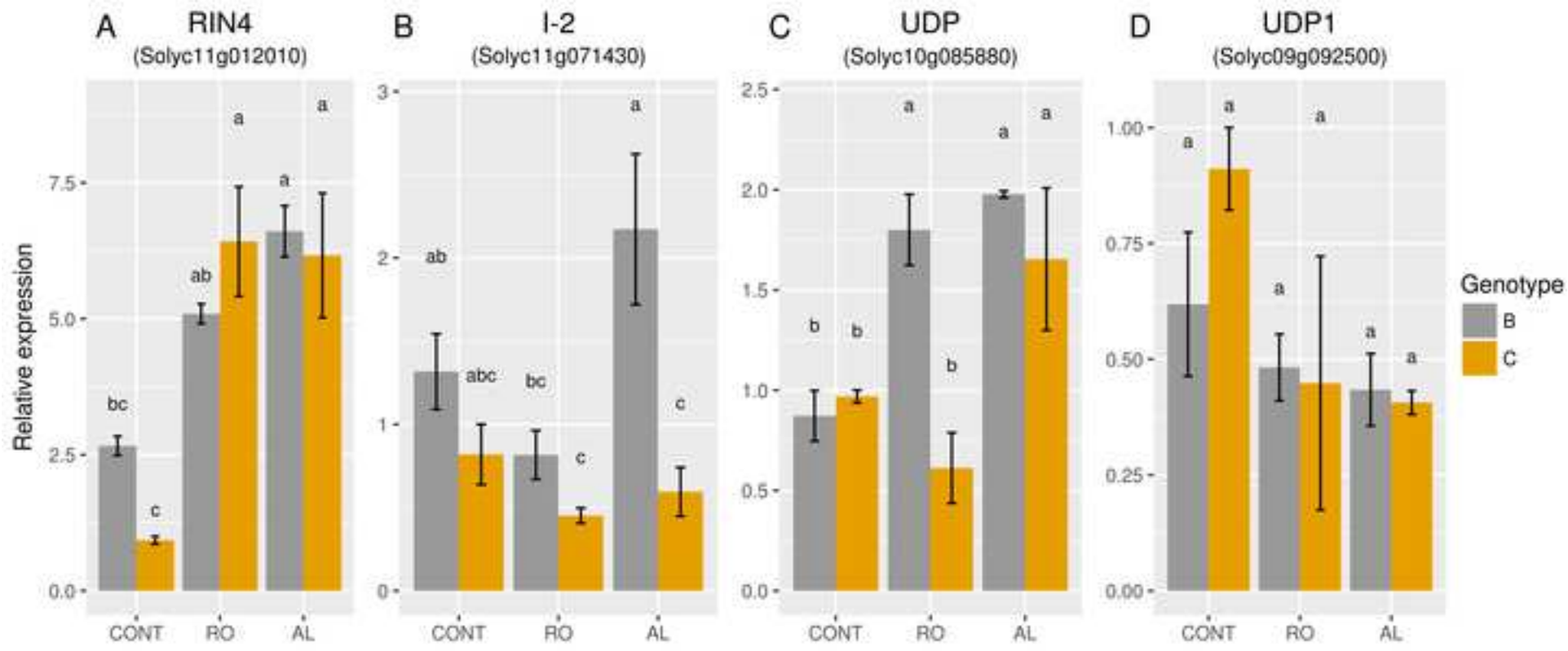
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