Т	Evaluation of the short term effect of nursery treatments with phosphite-based products,
2	acibenzolar-S-methyl, pelleted Brassica carinata and biocontrol agents, against lettuce and
3	cultivated rocket Fusarium wilt under artificial inoculation and greenhouse conditions
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13 ABSTRACT

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Experimental trials have been carried out in order to evaluate the efficacy of preventative 15 treatments based on plant defense activator products, biocontrol agents, a microbial complex with 16 arbuscular mycorrhizal fungi, and Brassica carinata pellets against Fusarium oxysporum f. sp. 17 lactucae race 1 on lettuce and Fusarium oxysporum f. sp. raphani on cultivated rocket under 18 greenhouse conditions. These products were compared with fungicides known for their ability to 19 induce host resistance (phosethyl-Al and acibenzolar-S-methyl), and with azoxystrobin. Three and 20 four applications of the tested products were carried out on lettuce and rocket seedlings grown in 21 nursery conditions. Treated and untreated plants were transplanted into soil infested with Fusarium 22 wilt agents to obtain an average disease severity (DS) of 65.6-69.2 and of 56.9-62.1 on the untreated 23 lettuce and rocket plants, respectively. The best Fusarium wilt biocontrol was obtained after four 24

applications of Bacillus subtilis Qst713 and with the Glomas microbial complex (42 and 46.7%, 25 26 efficacy, respectively). Brassica carinata pellets provided a consistent control when applied 14 days before the rocket and lettuce were transplanted into the infested soil. Acibenzolar-S-methyl, applied 27 at 0.025 g/liter, showed a DS reduction in F. oxysporum f. sp. lactucae from 36 to 61% and of F. 28 oxysporum f. sp. raphani from 54 to 73%, thus showing statistically similar results to those of 29 azoxystrobin, which was used as a reference (DS reduction from 59 to 65%). Although the 30 Fusarium wilt control provided by such products was not complete in the present experimental 31 conditions, these products can be considered interesting components for an integrated pest 32 management of the Fusarium wilt of leafy vegetables, starting from nursey applications. Moreover, 33 34 the tested BCAs could become potentially useful, especially for plant monocultures. This study has been produced new information on the effects of potassium phosphite, applied at the nursery level, 35 on reducing lettuce and rocket fusarium wilt. An average efficacy of 69.5% was observed for 36 lettuce, while an average efficacy of 65.2% was observed for cultivated rocket. The good fungicidal 37 activity of the phosphite-based product, coupled with the positive effect on plant biomass, is of special 38 interest. 39

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41 Keywords: Lactuca sativa, Eruca vesicaria, Fusarium oxysporum f. p. lactucae; Fusarium
42 oxysporum f. sp. raphani, integrated control

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44 1. Introduction

In recent years, the economic relevance of lettuce (*Lactuca sativa* L.) and cultivated rocket [*Eruca vesicaria* (L.) Cav.] has increased, since many farms produce fresh cut and ready-to-eat salads. In these intensive systems, where leafy vegetables are continuously grown in the same soil, the phytopathological situation is constantly in evolution, as a consequence of the dynamism and specialization of such crops (Garibaldi and Gullino, 2010; Garibaldi et al., 2014). Fusarium wilt

incited by Fusarium oxysporum f. sp. lactucae on lettuce and by Fusarium oxysporum: ff. spp. 50 51 raphani and conglutinans on cultivated rocket, can lead to serious losses (Matheron and Gullino, 2012). The Fusarium wilt of lettuce was detected for the first time in Europe, in northern Italy, in 52 2002 (Garibaldi and Gullino, 2010). The two formae speciales that affect cultivated rocket, that is, 53 Fusarium oxysporum ff.spp. raphani and conglutinans, the first of which is more frequently 54 detected (Garibaldi et al., 2006; Srinivasan et al., 2012), also affect other genera belonging to 55 56 Brassicaceae, such as cabbage, brussel sprouts, broccoli, turnip, radish and stock (Garibaldi et al., 2006). The Fusarium wilt of lettuce and rocket are easily and frequently seed-transmitted (Gullino 57 et al., 2014), thus suggesting the importance of preventative disease management strategies. 58

Lettuce varieties that are resistant or at least tolerant to Fusarium wilt are available (Scott et al., 2010; Matheron and Gullino, 2012; Gilardi et al., 2014b), but their effective use is complicated by the presence of three races of the pathogen (Fujinaga, 2005). In the case of rocket, the use of resistant varieties is still very limited (Gilardi et al., 2007).

In general, the management of soil-borne pathogens is complicated by the limited number of 63 64 registered chemicals and by the restrictions in the use of pre-plant fumigants, including metam sodium and dazomet (Colla et al., 2012). Several approaches to soil-borne disease management 65 have been investigated intensively in an attempt to find an answer to the many practical problems 66 associated with the loss or limitation of use of effective fumigants encountered by growers. 67 Moreover, more emphasis is now given to crop and soil health instead of disease control (Barrière 68 et al., 2014). Among the exploited strategies for disease management, systemic acquired resistance 69 (SAR) and induced systemic resistance (ISR), which are mainly triggered by microorganisms, such 70 as plant growth-promoting rhizobacteria, by the metabolic products of affected plants, or by 71 chemicals (Sticher et al., 1997; Oostendorp et al., 2001; Vallad and Goodman, 2004; Shoresh et al., 72 2005) are at present attracting a great deal of interest. Induced systemic and localized resistance to 73 soil-borne pathogens, by means of Trichoderma treatments, has been well documented (Shoresh et 74

al., 2005; Vinale et al., 2008). Kloepper et al., (2004) proved the ability of Bacillus spp. to induce 75 systemic resistance, and in most cases, to also elicit plant growth promotion. Moreover, arbuscular 76 mycorrhizal fungi have been reported to be implicated in protecting plants against soil-borne 77 pathogens, through different mechanisms, including an improvement in plant nutrition, damage 78 compensation, competition, changes in the root system and activation of plant defense signaling 79 (Whipps, 2004; Pozo and Azcón-Aguliar, 2007). Moreover, among the chemical resistance 80 inducers, the phosphite-based fertilizers and acibenzolar-S-methyl have been shown to lead to a 81 reduction in disease against soil-borne pathogens, on vegetable and ornamental crops (Eikemo et 82 al., 2003; Elmer 2004; 2006; Hyeon et al., 2009; Bubici et al., 2006; Walters, 2012; 2013; Gilardi et 83 84 al., 2014a), but no data are available concerning their efficacy against the Fusarium wilts of lettuce and rocket. 85

This study has been carried out in order to evaluate the efficacy of preventative treatments, including SAR and IRS activator products, biocontrol agents, a microbial complex based on arbuscular mycorrhizal fungi, and *Brassica carinata* pellets against *Fusarium oxysporum* f. sp. *lactucae* race 1 on lettuce and *Fusarium oxysporum* f. sp. *raphani* on cultivated rocket under simulated nursery conditions, in a greenhouse.

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92 2. Material and methods

93 2.1. Plant material and experimental layout

Experimental trials (Table 1) were carried out in 2012 and 2013 under greenhouse conditions in order to test the efficacy of different products against the Fusarium wilts of lettuce and cultivated rocket. Lettuce (cv. Crispilla) and cultivated rocket seeds (cv. Coltivata), which are very susceptible to Fusarium wilt (Garibaldi et al., 2004; Gilardi et al., 2014,), were sown in 100-plug trays (2.5 cm Ø per pot, 4-L of soil capacity) filled with a steamed (90°C for 30 minutes) peat mix substrate 99 (blond peat:black peat 15:85, pH 5.5-6.0, 1,100 g m–3 of N:P:K and traces of molybdenum, Brill
100 Type 5, Georgsdorf, Germany).

Fifteen-day-old lettuce and cultivated rocket seedlings were transplanted into the same substrate in 12-L plastic pots. Ten plants/pot of each tested crop were kept in a greenhouse on benches with air temperatures ranging from 26 to 34°C during the day and from 20 to 25°C during the night (Table 1). The substrate was artificially infested with the Fusarium wilt agents, as described hereafter. Forty plants per treatment were arranged in a complete randomized block design in each trial, which represented the experimental unit, with three (Protocol 1) and four replicated trials (Protocol 2) as reported in Table 1.

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109 2.2. Fungal strains and artificial inoculation

A highly virulent strain of *F. oxysporum* f. sp. *lactucae*, isolated in 2002 from infected lettuce plants in north-western Italy, ATCCMYA3040, belonging to Race 1 of the pathogen (Garibaldi et al., 2002), and the FusRuc 13/03 strain, isolated from rocket grown in a commercial plastic greenhouse in northern-Italy in 2003 and identified as *F. oxysporum* f. sp. *raphani* (Garibaldi et al., 2006), were used throughout the experiments. The single-spore culture of each isolate was stored in glycerol at -80°C.

These strains were grown in potato dextrose broth (Sigma-Aldrich, St. Luis, USA) and kept in a rotatory shaker working at 90-100 rpm for 10 days at 25°C. The biomass produced after centrifugation (9,600 g at 4°C) was prepared as a dry talc-based powder (biomass: talc 2:1 w/w), as described by Locke and Colhoun (1974). After 20 days at 22-25°C, the number of chlamydospores per gram of talc was assessed by serial plating on potato dextrose agar, PDA (Merck, Darmstadt, Germany), which contained 25 mg L⁻¹ streptomycin sulphate. The talc formulations of *F*. *oxysporum* f.sp. *lactucae* (strain MYA3040) and of *F. oxysporum* f.sp. *raphani*, (strain FusRuc13/03) at 2 and $5x10^7$ chlamidospores/g, respectively, were used. These formulated pathogens (2 and 1 g liter/respectively) were mixed into the steamed substrate as chlamydospores dispersed in talc to achieve a final concentration of $5x10^4$ chlamydospores ml⁻¹ of substrate. A non-infested substrate was used as a control (Table 1).

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128 2.3. Products used in the test

Different compounds known for their capability to induce resistance in the host, that is, phosphitebased fertilizers, organic amendments, biocontrol agents (BCAs) and fungicides were tested (Table 2).

Among the considered phosphite-based products and organic amendments, a phosphite-based glucohumate complex (Glucoinductor + GlucoActivator, N 4%, P₂O₅ 18%, International patent PCT, IB2004\001905, Fertirev, Torino, Italy), a mineral fertilizer based on potassium phosphite (Alexin 95PS, P₂O₅ 52%, K₂O 42%, Massò, Spain), and a patented formulation of *Brassica carinata* defatted seed meal (Biofence, N organic 3%, P 2.2%, K 2%, organic C 52%, Triumph, Spain) were tested.

Among the BCAs, *Bacillus subtilis* QST 713 (Serenade, 14.6 % a.i., BayerCropScience, Italy), *Bacillus velezensis* (Cilus Plus IT45, 95%, Massò, Spain), *Trichoderma asperellum +T. gamsii*(Remedier WP, Isagro Ricerca, Milano, Italy), a product based on arbuscular mycorrhizal fungi
combined with a microbial complex of *Trichoderma* and *Bacillus* (Rizocore, *Glomus* spp.
5%+*Bacillus megaterium* 10⁴ UFCg⁻¹ +*Trichoderma* 10¹⁰ UFCg⁻¹, Biogard, division of CBCEurope,
Italy) were tested.

The chemicals known for their ability to induce resistance, that is, acibenzolar-S-methyl (Bion
50WG, 50% a.i., Syngenta Crop Protection, Italy) and phosethyl-Al (Alliette, 80% a.i, Bayer Crop

Science, Italy) were tested, while azoxystrobin (Ortiva, 23.2% a. i., Syngenta Crop Protection, Italy)
was used as traditional chemical control (Gullino et al., 2002).

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149 *2.4. Type and timing of the treatments*

The biocontrol agents, acibenzolar-S-methyl, the phosphite based products, as well as the tested fungicides phosethyl-Al and azoxystrobin were applied as a leaf spray with a high volume of water (1,500 L ha⁻¹), using a 1 L capacity hand sprayer.

The temporal organization of the experimental trials, as well as, the timing of the artificial 153 infestation of the soil and treatments are summarized in Table 1. The timing and the application 154 dosages of the tested products, which were based on the manufacturer's suggestions, are given in 155 table 2. The lettuce and cultivated rocket seedlings grown in each tray were treated by leaf spraying 156 with three applications at 6-7 day intervals according to protocol 1 (Trial block 1 to 3), or with one 157 more spray applications in the 12 L plastic pots, as in protocol 2 (Trial block 4 to 7), seven days 158 after transplanting the lettuce and cultivated rocket seedlings. Azoxysrobin and *B. carinata* pellets 159 were only applied once (Tables 1 and 2). 160

The first treatment (T0) was carried out in a greenhouse, at a temperature of 22-24°C, on plants still
in the plug tray, at the second true leaf stage, 7-10 days after sowing.

163 The product based on arbuscular mycorrhizal fungi and the microbial complex (Rizocore) was 164 mixed with 4 L of the substrate that was used in the plug tray at T0, than leaf sprayed at 6-7 day 165 intervals (Table1). The patented formulation of *B. carinata* defatted seed meals was mixed with the 166 substrate used to fill the 12 L plastic pots at T0 (14 days before the crops transplant) and at the same 167 time as the artificial infestation of the soil at T7.

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The plants were monitored weekly and the data were recorded, starting 10-14 days after the lettuce 172 and rocket had been transplanted into the infested substrate, at the appearance of the first yellow 173 leaf symptoms and reduced growth. The number of infected plants showing wilting and stem 174 necrosis was counted to assess disease incidence. The totally wilted (dead) plants were removed. 175 The final disease rating was made four weeks after the soil infestation by dissecting each plant. The 176 used disease severity (DS) index was: 0 = healthy plant, 25 = initial leaf chlorosis, 50 = severe leaf 177 chlorosis and initial symptoms of wilting during the hottest hours of the day, 75 = severe wilting 178 179 and severe symptoms of leaf chlorosis; 100 = plant totally wilted, leaves completely necrotic. At the 180 end of the trials, the total biomass was weighed in order to evaluate the effect of the tested treatments on the yield. DS data were arcsine transformed in order to normalize the distribution of 181 variance. The data from the non-inoculated and untreated controls were not included in the 182 statistical evaluation of the DS data (Figure 1). 183

- The efficacy of the different treatments in controlling the Fusarium wilt of lettuce and rocket,corresponding to the percentage of DS reduction, was calculated as:
- 186
- 187 % efficacy =100 (DS $_t \ge 100/DS i_{control}$)
- 188
- 189 where,
- 190 i = inoculated and untreated control
- 191 t =tested treatments.
- 192

All the data were analysed by univariate ANOVA in SPSS 22.0, and the means were separated by means of Tukey's multiple comparison test (p=0.05). The standard errors are marked with error bars in all the figures.

196 **3. Results**

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198 Effect of pre-planting treatment on lettuce fusarium wilt

The average disease severity (DS) at the end of the lettuce trials in the inoculated non-treated 199 control plots was 62.1 and 56.9, respectively (Figures 1a and 2a). When at least three treatments 200 were carried out, the best control of Fusarium wilt on lettuce was provided by the phosphite-based 201 product Alexin (69.2% efficacy), and this was followed by acibezolar-S-methyl at 0.0125 (59.9% 202 efficacy), by phosethyl-Al (57.8% efficacy) and the phosphite-based glucohumate complex (52.4% 203 efficacy). These treatments were effective as one application of azoxystrobin, which was found to 204 205 protect the lettuce from Fusarium wilt with an efficacy 55.7%. The biocontrol agents Trichoderma asperellum + T. gamsii and B. subtilis provided a partial disease reduction, with statistically 206 different results compared to the inoculated and untreated control (32.8 and 30.7% efficacy, 207 208 respectively), while B. velenzensis and the microbial Glomus spp. + B. megaterium + Trichoderma complex (Rizocore) were similar to the inoculated and untreated control. The Brassica carinata 209 pellets, applied once seven days before transplanting, were not effective (Figure 1a). The higher 210 fresh weight of lettuce plants obtained with the first protocol reflected the greatest efficacy in 211 disease reduction, which was provided by phosphite-based products, by acibenzolar-S-methyl at 212 213 the lower tested dosages, and by phosethyl-Al (Figure 1b).

In general, when an extra treatment was applied to the potted plants after transplanting, according to protocol 2 (Figure 2a), all the tested biocontrol products differed significantly from the untreated control, and an improved efficacy for *Bacillus subtilis* (42.7% efficacy), *T. asperellum* + *T. gamsii* (38.3%, efficacy), and the microbial complex based *Glomus* spp complex (38.3%, efficacy) was observed. On the other hand, the extra treatment with acibenzolar-S-methyl and the phosphitebased products did not improve their efficacy. The application of *B. carinata* pellets led to a better control of Fusarium wilt when they were applied 14 days before transplanting at T0, with an average efficacy of 56% (Figure 2a). As far as the fresh weight of the lettuce plants at the end of
the trials is concerned, the best results, in terms of production, were observed in the plots in which
the best disease control was achieved (Figure 2b).

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225 Effect of the pre-planting treatment on rocket fusarium wilt

Similar results, in terms of disease control, to those observed on lettuce, were observed in the trials 226 performed to evaluate the effects of different pre-planting treatments against F. oxysporum f. sp. 227 raphani on cultivated rocket, with an average disease severity at the end of the trials in the infested 228 and untreated control of 62.1 and 56.9, respectively (Figures 3a and 4a). When at least three 229 treatments were carried out, according to protocol 1, that is, acibenzolar-S-methyl at 0.0125 g L⁻¹, 230 the phosphite based products (Alexin and the Glucohumate complex), provided similar results to 231 one application of azoxystrobin, which reduced Fusarium wilt by 65%. All the tested biocontrol 232 agents were ineffective, and showed similar results to the inoculated and untreated control (Figure 233 3a). In general, there were no significant differences in plant fresh weight between treatments and 234 the inoculated and non-treated control plants, with the exception of the results shown for Bacillus 235 subtilis, T. asperellum + T. gamsii and the Glomus spp. +Bacillus megaterium +Trichoderma 236 product, in one out of four trials (Figure 3b). 237

The tested products, which were applied in four treatments (protocol 2), as previously described, significantly reduced the Fusarium wilt of rocket compared to the infested and untreated control (Figure 4a). The best rocket fusarium wilt control result was provided by potassium phosphite (65%, efficacy), and this was followed by acibenzolar-S-methyl (57.5%, efficacy) and by the Glucohumate complex (56.8%, efficacy), which were statistically similar to azoxystrobin, applied at transplanting as a chemical reference, which showed 56.3% efficacy.

The *B. carinata* pellets also significantly reduced disease severity when they were applied 14 days before transplanting (60.3%, efficacy). In general, the four treatments carried out with the biocontrol agents *B. subtilis, B. velenzensis, T. asperellum +T. gamsii,* and by the microbial *Glomus* spp. + *B. megaterium* + *Trichoderma* complex, significantly reduced the disease, compared
to the untreated control (42.7%, 42.3%, 34.5%, 46.7% of efficacy, respectively).

The *Brassica carinata* pellets led to a positive effect on rocket fresh weigh, compared to the noninoculated and non-treated control, when applied 7-14 days before transplanting. The phosphitebased products led to an improvement in fresh weight after four treatments were applied (Figures 3b and 4b).

None of the products tested under the present experimental conditions affected the fresh weight ofthe lettuce or rocket plants, thus confirming the absence of phytotoxicity.

255

256 **4. Discussion**

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According to several authors, Fusarium wilt is a devasting disease that affects many economically
important vegetables and it is responsible for severe losses to growers worldwide (Elmer, 2006;
Katan et al., 2012; Matheron and Gullino, 2012; Walters, 2012; Garibaldi et al., 2014).

The tactics adopted to control this plant disease mainly involve the use of preventative measures, 261 such as the minimization of dissemination in the field, the use of plant resistant cultivars whenever 262 possible, the adoption of a proper crop rotation, and the removal and destruction of infested plant 263 material. Like other soil-borne pathogens, Fusarium wilt is difficult to manage with a single 264 approach and/or with a single product (Katan et al., 2012). Moreover, different measures are needed 265 for different cropping systems. Gordon and Koike (2015) have reviewed different ways of 266 managing lettuce fusarium wilt and they have pointed out the importance of the adoption of proper 267 268 practices in order to reduce the inoculum level. Among these measures, it has been found that 1month of solarization, with an average temperature of 47 to 49°C, at a depth of 5 cm, provides a 269

reduction of 41 to 92% of Fusarium wilt on lettuce in naturally infested fields (Matheron and 270 271 Porchas, 2012). Even in sub-optimal temperature regimes (50 to 48°C for 6 h, 45 to 43°C for 8 h and 40 to 38°C for 10 h/day), 14 days of thermal treatment under controlled conditions has provided 272 very valuable results, in terms of disease control on rocket (Gilardi et al., 2014). Moreover, recent 273 investigations on the antagonistic microorganisms Trichoderma asperellum and T. gamsii, with a 274 Glomus spp. +Bacillus megaterium +Trichoderma product used as a dressing on lettuce seeds, have 275 shown promising results in the control of fusarium wilt on lettuce (Lopez et al., 2014). On the 276 contrary, crop rotation is not an effective measure, because F. oxysporum f.sp. lactucae can 277 colonise the roots of other crops and, there is evidence of an expanded host range of Fusarium 278 279 oxysporum f.sp. raphani (Garibaldi et al., 2006; Scott et al., 2014).

Among the various alternatives to chemicals that have been tested so far, it has here been shown 280 that any solution on its own is able to provide a viable level of disease control of the Fusarium wilt 281 282 of lettuce and rocket, under short-cycles and for high levels of infestation. The success of the biological control strategies used against Fusarium wilt agents, through the adoption of various 283 antagonist, organic amendments and biofumigation, in fact depends on many different factors, 284 including the pathogen infestation level in the soil and the type of inoculum (Termorshuizen and 285 Jeger, 2014). Soil temperature also has an important effect on the disease expression of both of 286 287 these Fusarium wilt agents, and it has been shown to be a key factor that can influence the success of the control measures (Bosland et al. 1988; Scott et al., 2010a). 288

The present results provide evidence of the capability of commercial BCAs to reduce lettuce and rocket Fusarium wilt by 42 to 47%, respectively, under high disease pressure, when applied four times, starting from the nursery. The limited efficacy of lettuce and rocket Fusarium wilt control, after three applications of *Glomus* spp. +*Bacillus megaterium* +*Trichoderma*, could be due to an unsuccessful establishment of the mycorrhizal symbiosis. However, the mechanism involved in the control of Fusarium wilt by *Glomus* is complex and seems to be host specific. In the present study, four applications of the *Glomus* spp. +*Bacillus megaterium* +*Trichoderma* product provide a better Fusarium wilt reduction on rocket with 47% of efficacy. Martinez-Medina et al., (2010) have suggested that the mechanism involved in *Fusarium oxysporum* f.sp. *melonis* control by *Glomus intraradices* is independent of the SA and JA pathways, while the Fusarium wilt control of tomato by means of root colonization with mycorrhizae has been attributed to a possible plant-mediated phenomenon (Kapoor, 2008).

301 The present study provides new information on the effect of resistance inducers, based on either phosphites or acibenzolar-S-methyl, applied as a pre-plant treatment in the nursery, against F. 302 oxysporum f. sp. lactucae and F. oxysporum f. sp. raphani on lettuce and cultivated rocket, 303 304 respectively. Among the tested chemical resistance inducers, acibenzolar-S-methyl has been found to be effective in controlling the Fusarium wilt of both lettuce and rocket at the lowest tested 305 dosage, with a positive effect on the yield. Encouraging results pertaining to the control of 306 307 Verticillium wilt on eggplants and Fusarium wilt on ornamental plants by means of acibenzolar-Smethyl have already been reported by Elmer (2004; 2006) and by Bubici (2006). 308

The good fungicidal activity of the phosphite-based product, coupled with the positive effect on 309 plant biomass, is of special interest. In the present study, the phosphite-based products have 310 generally caused a consistent disease control of F. oxysporum f. sp. lactucae, that is, from 61 to 311 69%, and of F. oxysporum f. sp. raphani, from 54 to 65%, and have shown statistically similar 312 results to those of azoxystrobin, which has been used as a reference (59 to 65% efficacy), with a 313 significant effect on the yield. A previous study carried out on lettuce reported conflicting data on 314 plant growth for the use of phosphite-phosphate as a fertilizer. However, this may have been due to 315 the different cultivars that were examined, which could have different capabilities of absorbing 316 phosphite (Thao et al., 2009). The rate and timing of application of resistance inducers are 317 considered critical factors that can affect both the level of disease control and the yield (Walters, 318 2012). 319

321 The effectiveness of soil applications of the patented formulation of Brassica carinata defatted seed meals has shown conflicting results against the Fusarium wilt of lettuce and rocket. In the present 322 study, Brassica carinata pellets have provided a consistent control, when applied 14 days before 323 transplanting the rocket and lettuce into the infested soil, with an efficacy that ranged from 56 to 324 60%. The present results are consistent with previous research, because organic amendments need 325 326 protracted periods of time to become effective, since their activity is due to decomposition and the release of volatiles. (Mazzola et al., 2007; Bonanomi et al. 2010). The combination of Brassica 327 carinata pellets with solarization can be effective, as already observed, in infested soils, against the 328 329 Fusarium wilt of basil, lettuce and rocket (Garibaldi et al., 2011; Gilardi et al., 2014b).

Although the Fusarium wilt control provided by such products was not complete in the present experimental conditions, it should be considered that disease severity is generally lower in the field. Since the BCA treatments have not offered a complete Fusarium control, they could be integrated with other strategies, and their contribution to disease management could be interesting, because they can be used to complement other control measures. For instance, in the case of lettuce, the use of resistant or partially resistant commercial cultivars (Garibaldi et al., 2004; Matheron et al., 2005; Scott et al., 2010b; Gilardi et al., 2014b) could be combined with pre-plant treatments.

The four application of *Bacillus subtilis* Qs713 and the microbial *Glomus* spp. + *B. megaterium* + *Trichoderma* complex, as well as the three phosphite-based commercial products, which may be technically effective, especially at the nursery level, as well as economically feasible. The positive effect of these products against leaf pathogens should also be considered. *Bacillus* spp., from the rizhosphere of different vegetable plants, has resulted in induced resistance to *B. cinerea* (Levy et al., 2015). In a preventative application, phosphite and acibenzolar-S-methyl induced resistance to downy mildew on basil (Gilardi et al., 2013). Such products are easy and safe to apply under nursery conditions, and could represent an effective tool for the management of foliar and soil-borne pathogens. Moreover, the use of these products before they are applied at a farm level could help to standardize their applications in an integrated pest management model.

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489 Tables

Protocol, Sowing Tray treatment Artificial Transplanting Plot Treatment End of the trial (trial) inoculation (Day) 1(1) 6/04/2012 11/04 (T0)^a; 17/04 (T7); 24/04 (T14) 17/04//2012 25/04/2012 6/6/2012 -21/06(T0); 28/06(T7); 5/07 (T14) 1(2) 14/06/2012 28/06/2012 6/07/2012 4/09/2012 1(3) 10/01/2013 21/01(T0); 28/01(T7); 4/02(T14) 28/01/2013 5/02/2013 18/03/2013 _ 2 (4) 2/04/2013 10/04(T0); 17/04(T7); 24/04 (T14) 17/04/2013 25/04/2013 30/04/2013 28/05/2013 2 (5) 7/06/2013 14/06 (T0);21/06 (T7); 27/06 (T14) 21/06/2013 28/06/2013 4/07 /2013 23/07/2013 1/08 (T0); 8/08 (T7); 12/08 (T14) 2 (6) 25/07/2013 12/08/2/2013 13/08/2013 19/08/2013 17/09/2013 4/10/2013 2 (7) 9/09/2013 20/09 (T0); 27/09 (T7); 3/10 (T14) 27/09/2013 10/10/2013 28/10/2013

490 Table 1. General information on the trials and timing of the operations carried out on lettuce and rocket.

^aT0 corresponding to the first treatment carried out as leaf spraying at the development stage of 1-2 true leaves and soil

492 mixing application.

	BCA or active ingredient	Commercial	Dosage	Time (days) of application in tray	Time (days) of
		formulation	a. i.	conditions and type of application	application in
			g L ⁻¹		plastic pot (2-L
					and 12-L) and
					type of
					application
	Bacillus subtilis QST713	Serenade Max	0.58	T0ª,T7,T14 (T21), leaf spray	-
	Bacillus velezensis	Cilus Plus ^b	0.4 ^b	T0,T7,T14, (T21)leaf spray	-
	T. asperellum + T. gamsii	Remedier	0.04	T0,T7,T14, (T21)leaf spray	-
	Acibenzolar-S-methyl	Bion 50 WG	0.025	T0,T7,T14, (T21) leaf spray	-
			0.0125		
	Phosethyl-Al	Aliette	1.6	T0,T7,T14, (T21)leaf spray	-
	Glomus spp. +Bacillus megaterium	Rizocore ^b	0.08 ^b	T0,T7,T14, (T21) leaf spray	-
	+Trichoderma				
	Potassium phosphite P:K 52:42	Alexin	1.3+1.06	T0,T7,T14, (T21) leaf spray	-
	Phosphite based glucohumate N:P 4:18	Glucohumate complex	1.6+0.72	T0,T7,T14, (T21)leaf spray	-
	Azoxystrobin	Ortiva	0.19	T14, (T21)	-
				leaf spray	
	Brassica carinata pellet	Biofence	0.15+0.055+0.0	-	T0 - T7,
	N:P:K: C organic		5+1.13		soil mixing ^c
495	^a T0 corresponds to the leaf spraying	carried out, starting a	t the second tru	ue leaf stage, and at seven-day	intervals (T7
496	and T14).				
497	^b Corresponds to the dosage (g L ⁻¹) o	of the commercial form	nulation.		
498	^c Corresponds to the treatment carrie	ed out at 14 day before	transplanting (T0) and 7 days before transplan	ting the crops.
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494 Table 2. List of the tested products and of the experimental protocol.

Figure 1a. Effect of three tray-treatments on the Fusarium wilt of lettuce (cv. Crispilla). The data
are expressed as DS at the end of the trials and compared with azoxystrobin and *Brassica carinata*pellets applied once.



Figure 1b. Effect of three tray-treatments on the yield of lettuce plants in the presence of *F. oxysporum* f. sp. *lactucae*. The data are expressed as fresh weight and compared with azoxystrobin and *Brassica carinata*pellets applied once (g).



Figure 2a. Effect of four treatments on Fusarium wilt of lettuce (cv. Crispilla). The data are expressed as DS at the end of the trials and compared with azoxystrobin and Brassica carinata pellets applied at transplanting (T7) and 14 days before transplanting the lettuce (T0).





540 Figure 2b. Effect of four treatments on the yield of lettuce plants in the presence of *F. oxysporum* f. sp.

lactucae. The data are expressed as fresh weight and compared with azoxystrobin and *Brassica carinata*



542 pellets applied once (g).

Figure 3a. Effect of three tray-treatments against the Fusarium wilt of rocket (cv. Coltivata). The data are

expressed as DS at the end of the trials and compared with azoxystrobin and *Brassica carinata*



557 pellets applied at transplanting (T7).



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Figure 4a. Effect of four treatments on the Fusarium wilt of rocket. The data are expressed as DS at the end of the trials and compared with azoxystrobin and *Brassica carinata* pellets applied at transplanting (T7) and 14 days before transplanting rocket, at the same time as the artificial infestation (T0).



Figure 4b. Effect of four treatments on the yield of rocket plants in the presence of *F. oxysporum* f. sp. *raphani*. The data are expressed as fresh weight and compared with azoxystrobin and *Brassica carinata*pellets applied once (g).



