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Improving the efficacy of biocontrol agents against soilborne pathogens

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23 **Control of soilborne pathogens of tomato using a commercial formulation of *Streptomyces***
24 ***griseoviridis* and solarization**

25

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34

35 **Abstract**

36 Mycostop® is a commercial formulation of strain K61 of *Streptomyces griseoviridis*. This strain
37 was isolated from *Sphagnum* peat and can control or suppress some root rot and wilt diseases by
38 colonizing the rhizosphere prior to pathogens. The present work was carried out to test the ability of
39 the commercial formulation, combined or not with soil solarization, to control different diseases of
40 greenhouse-grown tomato. Data obtained from four trials carried out over two years (2001 and
41 2002) demonstrated that *S. griseoviridis* could play a role in the integrated control of tomato
42 soilborne diseases. This study is among the first to test *S. griseoviridis*'s effectiveness against corky
43 root rot caused by *Pyrenochaeta lycopersici* when it is applied throughout the irrigation system (10
44 l of water per m²). The biofungicide was very effective against *Fusarium oxysporum* f.sp.
45 *lycopersici* and *Verticillium dahliae* in 2002 in artificially infested soils; however, in 2001 there was
46 no statistically significant reduction of the vascular wilts compared to the control. Soil spraying was
47 more effective than soil irrigation to control tomato wilts. The bacterial antagonist was not effective
48 against *Fusarium* crown and root rot caused by *F. oxysporum* f.sp. *radicis-lycopersici*, when applied

49 alone, but was less effective when applied with *S. griseoviridis*. Soil solarization provided good
50 control of *V. dahliae* and *F. oxysporum* f.sp. *lycopersici*, but also was slightly less effective when
51 combined with *S. griseoviridis*. A significant increase in fruit mass and a higher yield m⁻² was
52 recorded when solarization and the biofungicide were applied together in 2001. This indicated there
53 may be a potential additive effect of the commercial biofungicide and solarization in increasing
54 tomato yield; however, it was not consistent and generally not significantly different from the
55 inoculated control. Metham sodium provided the most effective control of corky root and greatest
56 yield increase of all the treatments evaluated.

57

58 **Keywords:** biological control, *Fusarium oxysporum* f.sp. *lycopersici*, *Fusarium oxysporum* f.sp.
59 *radicis-lycopersici*, integrated control, *Pyrenochaeta lycopersici*, solarization, *Streptomyces*
60 *griseoviridis*, tomato, *Verticillium dahliae*.

61

62 **1. Introduction**

63 In the last thirty years, glasshouse cultivation of tomato (*Lycopersicon esculentum* Mill.) has greatly
64 increased in mild and warm climate regions (Albajes et al., 1999). The intensification of glasshouse
65 tomato production has created optimal conditions for many pathogens (Jones et al., 1991). Collar
66 rot, caused by *Rhizoctonia solani* Kühn and verticillium wilt, caused by *Verticillium* spp. (*V.*
67 *dahliae* in Italy), occur in all tomato growing regions. *Pyrenochaeta lycopersici* Schneider &
68 Gerlach, *Fusarium oxysporum* Schlechtend.: Fr. f.sp. *lycopersici* (Sacc.) Snyder & Hans. and *F.*
69 *oxysporum* f.sp. *radicis-lycopersici* Jarvis & Shoemaker are also widespread pathogens whose
70 severity varies with regional cultural practices.

71 Soilborne pathogens were effectively controlled by methyl bromide and tomato crops account for
72 about 30% of the methyl bromide used in the world (UNEP, 1995) and 43% of its use in Italy
73 (Gullino et al., 2003). Concern for the potential depletion of ozone by methyl bromide led to its
74 inclusion among the substances controlled by the Montreal protocol which stated that use by

75 industrialized countries should have been eliminated by the end of 2004 (Bell et al., 1996; Gullino
76 et al., 2003). Alternatives of biological control and soil solarization, alone or in combination, are
77 promising methyl bromide replacement methods for controlling soilborne diseases of tomato.
78 Microbial antagonists have been widely studied as biological controls the last few years.
79 *Penicillium oxalicum* reduced the incidence of *F. oxysporum* f. sp. *lycopersici* (Cal et al., 1997).
80 *Trichoderma harzianum* and *T. koningii* controlled Fusarium root and crown rot (Bourbos et al.,
81 1997). Non-pathogenic strains of *F. oxysporum*, obtained from suppressive soil, controlled
82 Fusarium wilts (Alabouvette, 1988; Minuto et al., 1995a; Minuto et al., 1997). Many of these
83 biological control agents, however, are still being tested and are not commercially available.
84 Actinomycetes have been recognized as sources of several secondary metabolites, antibiotics, and
85 lytic enzymes that affect fungal growth (Goodfellow and Williams, 1983). A strain of *Streptomyces*
86 sp., isolated from the rhizosphere of field-grown tomato, has been reported to suppress damping-off
87 of tomato transplants caused by *R. solani* in a peat-based, soilless potting mix under greenhouse
88 conditions (Sabaratnam and Traquair, 2002). Isolates of *Streptomyces* spp. were assessed in Finland
89 for 20 years for their ability to control fungal diseases (Tahvonen, 1982) and a selected *S.*
90 *griseoviridis* has been developed as a biofungicide by Verdera OY and tested on a wide range of
91 organisms on glasshouse and field crops (Tahvonen and Avikainen, 1987; Lahdenperä, 1987).
92 Mycostop® is a biofungicide based on the K61 strain of *S. griseoviridis* isolated from *Sphagnum*
93 peat (White et al., 1990). Streptomycetes are active in the rhizosphere and the modes of action of
94 species tested include antibiosis, lysis of fungal cell walls, competition and hyperparasitism
95 (Mohammadi and Lahdenperä, 1992; Tapio and Pohto-Lahdenperä, 1991). The commercial product
96 based on strain K61 can control or suppress some root rot and wilt diseases, caused by *Pythium*
97 spp., *Fusarium* spp., *Rhizoctonia* spp. and *Phytophthora* spp., if it colonizes the rhizosphere prior to
98 the pathogens. Mycostop® is registered in many European countries for use on different vegetables,
99 herbs, and ornamentals, such as basil, cucumber, eggplant, melon, pepper, tomato, pumpkin,
100 gerbera, cyclamen, and carnation (in Italy: Registration 10506 of the Italian Ministry of Health).

101 Solarization, carried out by covering the soil with plastic film during the hot season, has been
102 widely exploited in a number of warm countries as well as in climatically marginal ones (Katan and
103 DeVay, 1991). In Northern Italy, its efficacy is improved when applied with transparent mulch (30-
104 40 μm thick) in the greenhouse (Garibaldi and Gullino, 1991). Many growers are skeptical about its
105 effectiveness since it requires soil free of cultivation for at least four weeks. A practical possibility,
106 often adopted to increase soil solarization efficiency and, especially, to enlarge the spectrum of
107 efficacy of biocontrol agents, is the combination of both methods (Katan and DeVay, 1991).
108 Previous studies clearly demonstrate the feasibility of a combination of soil solarization and
109 biocontrol agents that target reducing the mulching period (Minuto et al., 1995b).

110 The present work was carried out under semi-commercial conditions to evaluate the effectiveness of
111 *Streptomyces griseoviridis* isolate K61 for control of different soilborne pathogens of greenhouse-
112 grown tomato. Its combination with soil solarization also was investigated in order to evaluate
113 potential beneficial effects of *S. griseoviridis* applied after mulching and to explore the possibility
114 of reducing its duration. All of these activities were used to better define the formulation label in
115 terms of application methods, according to European legislation (Directive 91/414 CE) related to
116 pesticide registration. In this regard, different dosages and application methods were assessed for
117 their disease control effectiveness. A general objective was to study the relationship between plant
118 productivity and disease incidence as they were affected by the different treatments applied. To
119 increase the incidence of some diseases during specific experiments, some target pathogens were
120 inoculated into the soils.

121

122 **2. Materials and methods**

123

124 *2.1. Layout of tomato trials*

125 During 2001-2002, four experimental trials were carried out with different pathogens in four
126 different glasshouses of the Experimental Station (CeRSAA) of the Chamber of Commerce of

127 Savona located in Albenga (Italian Riviera). Experiments used tomato cv. Cuore di Bue planted in
128 soil systems (sandy loam: sand 75%, silt 20%, clay 5%; pH, 8.1; organic matter content, 2.5%;
129 cation exchange capacity, 8.5 meq 100g⁻¹ soil) to assess the efficacy of biological and physical
130 methods against different soilborne pathogens. A randomized complete block design was used with
131 four replicates per trial. The cultivation density was 2 plants m⁻². During the first and second trial,
132 16 plants per replicate were used; during the third and fourth trial, the number of plants per replicate
133 was increased to 40. The efficacy against tomato soilborne pathogens of a commercial formulation
134 of *Streptomyces griseoviridis* alone, or combined with solarization, was compared to the efficacy of
135 chemical pesticides used for soil disinfestations in intensive horticulture (metham sodium) or
136 known for its efficacy against vascular diseases (benomyl). The layout of the four trials is
137 summarized in Table 1.

138

139 2.2 Soil infestation with pathogen inocula

140 In order to evaluate the effectiveness of Mycostop® under semi-commercial conditions, the
141 experimental activities were carried out in greenhouses where, without any soil disinfestation,
142 tomato had been grown at least the two previous growing seasons and showed some symptoms of
143 vascular wilt (*F. oxysporum* f.sp. *lycopersici* and *V. dahliae*) and basal root rot (*R. solani*, *F.*
144 *oxysporum* f.sp. *radicis-lycopersici*). Nevertheless, to achieve a uniform soil infestation and higher
145 disease pressure before solarization, fumigation or Mycostop® application, artificial soil
146 inoculations of some pathogens were executed. Two strains of each pathogen (*F. oxysporum* f.sp.
147 *radicis-lycopersici*, *F. oxysporum* f.sp. *lycopersici*, *V. dahliae*, or *P. lycopersici*), freshly isolated
148 from tomato plants, were grown on autoclaved wheat kernels. Inocula were incorporated into the
149 soil by rototilling to a depth of 15 cm 7-10 d prior to soil treatments at the dosages reported in Table
150 1. Soil was kept moist for one week by periodic sprinkler irrigation (5-10 mm).

151

152 2.3. Soil treatments

153 In trial 1, the efficacy of the commercial biocontrol agent *S. griseoviridis* (Mycostop®; Verdera
154 OY, Helsinki, Finland) applied at 10^7 colony forming units (cfu) m^{-2} was compared with the
155 efficacy of solarization and a combination of the two methods. *S. griseoviridis* was delivered as
156 suspension of the powder formulation ($0.1\text{ g }m^{-2}$ with 0.5 l of water per m^2) immediately after
157 transplanting and repeated three times every four to five weeks.

158 Solarization was accomplished by covering the soil for 26 d with standard low-density polyethylene
159 (LDPE, Eiffel, Fontanellato, Italy, $40\text{ }\mu\text{m}$ thick). Soil was irrigated ($30\text{-}35\text{ mm}$ of water m^{-2}) the day
160 before mulching. In the combination antagonist and solarization, the biofungicide was applied to the
161 soil immediately at the end of solarization. In trial 2 the same treatments were compared but *S.*
162 *griseoviridis* was applied at 10^7 cfu. and 5×10^6 cfu. m^{-2} (0.05 g of commercial formulation in 0.25
163 l of water m^{-2}).

164 Soil solarization was not evaluated in 2002, and three different methods of application of *S.*
165 *griseoviridis* were compared with fumigation by metham sodium (trial 3) or benomyl (trial 4).
166 Mycostop® was applied at 10^7 cfu m^{-2} ($0.1\text{ g }m^{-2}$) or at 5×10^6 cfu m^{-2} ($0.05\text{ g }m^{-2}$) by irrigation or at
167 10^7 cfu m^{-2} ($0.1\text{ g }m^{-2}$) by spraying. Irrigation was accomplished by drenching the soil with $5\text{ l }m^{-2}$
168 of water immediately before applying the biofungicide in $2\text{ l }m^{-2}$ with another $3\text{ l }m^{-2}$ immediately
169 afterwards. When spraying, the commercial formulation was delivered with $0.5\text{ l }m^{-2}$ of water.
170 Metham sodium (Vapam, SIPCAM, 32.7% a.i. corresponding to 380 g a.i./ l formulation) was
171 applied as a water suspension ($76\text{ g a.i. }m^{-2}$) using $15\text{ l }m^{-2}$ of water. Benomyl (Benlate, DuPont,
172 50% a.i.) was distributed as a water suspension ($2\text{ g a.i. }m^{-2}$) using $10\text{ l }m^{-2}$ of water.

173 In every trial a not treated control was introduced.

174

175 *2.4. Tomato transplant and cultural practices*

176 Tomato plants, 40-50 d old, belonging to the cultivar Cuore di Bue, were transplanted 7 d after soil
177 treatment. Plants were placed in two rows per bed approximately 15-20 cm from the bed edge and
178 40-50 cm apart. Plants were fertilized in the drip irrigation system five times at 10 day intervals

179 with a solution of N:P₂O₅:K₂O (20:10:10) at 100kg/ha each time. Three insecticide sprays to reduce
180 the presence of virus (TSWV, TYLC) vectors were applied at 7-10 day intervals after transplanting.

181

182 2.5. Data collection and analysis

183 Infection by *F. oxysporum* f.sp. *radicis-lycopersici* (FORL), *P. lycopersici* and the two vascular
184 pathogens, *F. oxysporum* f.sp. *lycopersici* (FOL) and *V. dahliae*, were assessed at the beginning of
185 each trial and several times later to correctly identify the pathogens by plating vascular tissues from
186 diseased plants on potato dextrose agar (PDA, Merck) or Komada's semi-selective medium
187 (Komada, 1975). Natural infections of *Rhizoctonia solani*, which frequently occurred during the
188 cropping period, were also evaluated. Disease development was evaluated every 10-14 d by
189 counting and eliminating symptomatic and collapsed plants (disease pressure). The final data is
190 reported for all but trial 1 and also indicates the number of healthy plants. The yield was evaluated
191 on healthy plants by weighing and counting the number of fruits per plant. Data are expressed as
192 number and mass of marketable fruits per plant. All data collected were statistically analyzed
193 according to Duncan's Multiple Range Test (P=0.05).

194

195 3. Results

196

197 3.1 Trial 1

198 The inoculated control plants (Table 2) at the end of the trial had a high level of infection by the
199 three pathogens inoculated (*F. oxysporum* f.sp. *radicis-lycopersici*, *F. oxysporum* f.sp. *lycopersici*
200 and *V. dahliae*). In addition, a very high percentage (31.3%) of the control plants were naturally
201 infected by corky root rot (*P. lycopersici*). The application of *S. griseoviridis* at 10⁷ cfu m⁻² partially
202 controlled the natural infection of corky root rot (18.8%), but was ineffective against the wilt
203 pathogens (12.5%) and Fusarium crown and root rot (20.8%). Nevertheless, two months after
204 planting, the antagonist had partially reduced the incidence of the inoculated pathogens (Table 2).

205 Solarization, applied alone, significantly reduced vascular wilts (2.1%), and also Fusarium crown
206 and root rot (2.1%). The plots treated with solarization and *S. griseoviridis* were severely infected
207 by *F. oxysporum* f.sp. *radicis-lycopersici* (18.8%), but the wilt pathogens (2.1%) and corky root
208 (18.8%) were controlled satisfactorily (Table 2).

209 No significant difference was observed in the number of fruits produced per healthy plant and in the
210 mass of fruits per healthy plant (Table 3). The average mass of tomato fruits was significantly
211 higher with solarization and *S. griseoviridis* (138.8 g), than the control (110.0 g).

212

213 3.2 Trial 2

214 The second trial was carried out at the same time as the first one, from July to December 2001. In
215 this trial, the plots were inoculated with *F. oxysporum* f.sp. *radicis-lycopersici*, which infected
216 21.2% of the control plants (Table 4). Corky root natural infestation was very low. Only a few
217 plants were attacked by *R. solani* at the beginning of the cropping season and no differences were
218 observed between treated and untreated plots.

219 Neither the physical or biological control method, nor their combination, significantly reduced the
220 incidence of Fusarium crown and root rot. This confirms the highly aggressive nature of the causal
221 agent and its difficulty to control. Neither formulation of *S. griseoviridis*, alone, controlled the wilt
222 pathogens, while solarization alone and the combination of antagonist and physical treatment were
223 able to control them.

224 Solarization, alone or combined with *S. griseoviridis* controlled the wilt pathogens, while neither
225 formulation of *S. griseoviridis* controlled the wilt pathogens.

226 A lower mass and number of fruits per plant were observed in the solarized plots (Table 5).
227 Moreover, the yield per m² data showed that the application of *S. griseoviridis* combined with
228 solarization increased yield (428 g) statistically more than the control (270 g).

229

230 3.3 Trial 3

231 This trial was carried out in a soil artificially infested with *P. lycopersici* where severe infection by
232 corky root rot had been observed in the previous tomato crop. During the experiment, a low
233 incidence of *F. oxysporum* f.sp. *radicis-lycopersici* and *F. oxysporum* f. sp. *lycopersici* was
234 observed (Table 6). The incidence of *F. oxysporum* f.sp. *radicis lycopersici* was not statistically
235 different between the treatments and the inoculated control. Attacks of *F. oxysporum* f.sp.
236 *lycopersici* registered in the control were low (1.3%) but they were statistically higher in the plots
237 treated with metham sodium.

238 At the end of the cropping cycle, the average percentage of the root surface of surviving plants
239 attacked by *P. lycopersici* was calculated to determine the severity of infection (Table 6). In the
240 inoculated control, 10.3% of the root surface showed symptoms of corky root compared with only
241 5.9% in the plots treated with metham sodium. All of the treatments containing *S. griseoviridis*
242 provided medium levels of efficacy regardless of the mode of application or the dosage applied.
243 Disease severity in the plots treated with the bacterial antagonist was not different from the plots
244 treated with metham sodium or the control.

245 In contrast, the incidence of *P. lycopersici* was very high in the control (55.3%) and lowest in the
246 plots fumigated with metham sodium (7.4%). Application of the biofungicide through the irrigation
247 system at 0.05 and 0.1 g per m², significantly reduced the incidence of infections compared with the
248 untreated control. The application of *S. griseoviridis* by spraying at 0.1 g per m² was not effective.

249 None of the treatments significantly affected either the quality or the yield, represented by the
250 average mass of single fruit or their quality (Table 7). The total mass and number of fruits produced
251 per plant were significantly higher (yield) in the plots fumigated with metham sodium. *S.*
252 *griseoviridis* given by irrigation at the rate of 10⁷ cfu m⁻² nearly doubled the yield (mass/plant)
253 compared to the untreated inoculated control, even though the yield increase was not statistically
254 significant.

255

256 *3.4 Trial 4*

257 Natural infection by *R. solani* in 2002, caused slight damage to the crop with no significant
258 differences among the treatments (Table 8). No infection by *V. dahliae* and only a low infection of
259 *F. oxysporum* f.sp. *radicis-lycopersici* were recorded in 2002 and there were no significant effects of
260 any treatment on the incidence of disease. All of the treatments generally limited the incidence of
261 Fusarium wilt but only *S. griseoviridis* applied by spraying at 10^7 cfu m⁻² provided a protection
262 significantly different from the control. Heavy yield losses caused by a strong reduction in fruit set
263 due to adverse temperature conditions (data not published). Tomato cv. Cuore di Bue is particularly
264 sensitive to strong temperature changes.

265

266 **4. Discussion**

267 The aim of this work was to test the ability of a commercial formulation of *S. griseoviridis*
268 combined or not with soil solarization to control several diseases of greenhouse-grown tomato.

269 Tomato provides a good example of how the use of biocontrol agents can be introduced into
270 practice as an IPM strategy (Albajes et al., 1999). Examples of integrated methods are provided by
271 the combination of biocontrol agents applied as seed dressing, with soil solarization (Gullino, 1998;
272 Minuto et al., 1995b; Spadaro and Gullino, 2005). Sivan and Chet (1993) combined *Trichoderma*
273 *harzianum* with soil solarization under field conditions to obtain significant control of Fusarium
274 crown and root rot of tomato and a significant yield increase.

275 Data obtained over two years (2001 and 2002) demonstrated that *S. griseoviridis* could play a role
276 in integrated control of tomato diseases. In some cases, the effect of biological control was more
277 pronounced early after transplanting (Table 2), but were less evident at the end of the experiment
278 (Table 2). This seemed to confirm the need for information related to the microorganism's ability to
279 survive in a natural soil.

280 This study shows that biocontrol agents can be effective against corky root rot (Bochow, 1989). *S.*
281 *griseoviridis* is a root colonizer and stimulates root growth during rhizosphere colonization
282 (Kortemaa et al., 1994). In some cases, stimulated plant growth could explain the enhanced yield

283 results (trial 1 and trial 2) when the antagonist was combined with soil solarization.

284 The effectiveness of Mycostop® against Fusarium wilt of tomato was satisfactory in 2002 (Table 8)

285 when applied to artificially infested soil. It was not effective against Fusarium and Verticillium

286 wilts of tomato in 2001 (Tables 2 and 4). Spraying was a more effective method of application than

287 irrigation, contrary to the results obtained with corky root. In trials carried out previously, strain

288 K61 formulated as Mycostop® had exhibited partial efficacy against *F. oxysporum* f.sp. *basilici*

289 (Minuto et al., 1997), but the same formulation was not effective when tested as a root dip (10^6 cfu

290 ml^{-1}) and soil drench (10^8 cfu ml^{-1}) in glasshouse experiments to control *F. oxysporum* f.sp. *dianthi*

291 on carnation (Garibaldi et al., 1990). Biocontrol efficacy is apparently determined by the method of

292 application of the biocontrol agents to the ecosystem, the number of treatments and the ability of the

293 selected strain to adapt to different environments.

294 The ineffectiveness of *S. griseoviridis* against Fusarium crown and root rot could be explained

295 because *F. oxysporum* f.sp. *radicis-lycopersici* has airborne microconidia that reinfest disinfested

296 soils (Rowe et al., 1977). Soil treatments, therefore, could *a priori* be expected to provide

297 inadequate protection against this pathogen (Rowe and Farely, 1981).

298 The combination of the biocontrol agent with soil solarization generally increased disease control

299 and yield. Soil solarization, as previously documented before (Katan and DeVay, 1991), provided

300 good control of *V. dahliae* and *F. oxysporum* f.sp. *lycopersici*. In trial 1, solarization was also

301 effective against Fusarium crown and root rot, but the result was not confirmed by trial 2. The

302 combination of soil solarization and *S. griseoviridis* was effective against Fusarium and

303 Verticillium wilts and corky root. Previous experiments showed that soil solarization could also be

304 exploited for corky root control (Garibaldi and Tamietti, 1983).

305 In general, no significant differences in fruit yield or quality were observed when solarization and

306 Mycostop® were applied together.

307 Metham sodium fumigation provided a high level of control of *P. lycopersici* and a superior mass

308 and number of fruits per plant produced. The efficacy of *S. griseoviridis*, although encouraging, is

309 not competitive with the effectiveness of this fumigant or as effective as methyl bromide,
310 chloropicrin, or dazomet. The incidence of Fusarium wilt in the fumigated plots was significantly
311 higher than the other treatments and the control. This may have been because of the low incidence
312 of corky root, but also could have been because metham sodium does not always guarantee a
313 complete protection of the fumigated soil when used at the low dosage of 250 ml m⁻² of the
314 commercial formulation (32.7 % a.i.).

315 *S. griseoviridis* could play a role in integrated control of different soilborne diseases but alone could
316 not control the main soilborne diseases of tomato. Its adoption after a solarization treatment may be
317 used to reduce the period of solarization. Solarization is mainly inconvenient by preventing use of
318 the soil during the hot season, but possesses great potential as an alternative to fumigation for soil
319 disinfection. Solarization alone controlled at least two serious soilborne pathogens; however, the
320 combination of soil solarization and *S. griseoviridis* was effective against Fusarium and
321 Verticillium wilts and somewhat against corky root even though the biofungicide did not improve
322 control of the individual pathogens. Moreover, at least in one case, the combination of physical and
323 biological methods increased the yield of tomato plants per area unit.

324

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329

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405

406 **Tables**

407

408 Table 1 – Experimental protocols for the four trials carried out against different soilborne pathogens

409 on tomato cv. Cuore di bue.

	1st trial	2 nd trial	3 rd trial	4 th trial
Artificial inoculation*	FORL (30 g m ⁻²), FOL (30 g m ⁻²), <i>Verticillium dahliae</i> (15 g m ⁻²)	FORL (20 g m ⁻²)	<i>Pyrenochaeta lycopersici</i> (30 g m ⁻²)	FOL (35 g m ⁻²)
Treatments with <i>Streptomyces griseoviridis</i> by irrigation (I) or by spraying (S)	10 ⁷ cfu m ⁻² (S)	10 ⁷ cfu m ⁻² (S) 5x10 ⁶ cfu m ⁻² (S)	10 ⁷ cfu m ⁻² (I) 5x10 ⁶ cfu m ⁻² (I) 10 ⁷ cfu m ⁻² (S)	10 ⁷ cfu m ⁻² (I) 5x10 ⁶ cfu m ⁻² (I) 10 ⁷ cfu m ⁻² (S)
Solarization (26 d)	alone and followed by Mycostop® application	alone and followed by Mycostop® application	---	---
Chemical control	---	---	Metham sodium (76 g a.i. m ⁻²)	Benomyl (2 g a.i. m ⁻²)
Planting	July 2001	July 2001	April 2002	July 2002
Harvest	December 2001	December 2001	July 2002	October 2002

410 *All artificially inoculated pathogens were propagated on wheat kernels. Dose expressed as mass of

411 infected kernels m⁻². FOL: *Fusarium oxysporum* f.sp. *lycopersici*. FORL: *F. oxysporum* f.sp.412 *radicis-lycopersici*.

413

414 Table 2 – Effect of treatments with *Streptomyces griseoviridis*, soil solarization or their combination
 415 on the number of tomato plants infected with *Rhizoctonia solani*, *Fusarium oxysporum* f.sp. *radicis-*
 416 *lycopersici* (FORL), *F. oxysporum* f.sp. *lycopersici* (FOL), *Verticillium dahliae* and *Pyrenochaeta*
 417 *lycopersici* after two months (left) and six months (right) (trial 1, 2001)

Treatment	% plants infected by (2 months after planting)				% plants infected by (end of the trial)				
	<i>R. solani</i>	FORL	FOL + V. <i>dahliae</i>	Total	<i>R. solani</i>	FORL	FOL + V. <i>dahliae</i>	<i>P. lycopersici</i>	Total
Inoculated control	2.1 a*	0 a	12.5 b	14.6	2.1 a*	16.7 b	18.8 b	31.3 B	68.9
<i>S. griseoviridis</i> 10^7 cfu m ⁻² in 0.5 l m ⁻²	2.1 a	0 a	4.2 ab	6.3	2.1 a	20.8 b	12.5 b	18.8 A	54.2
Solarization	0 a	0 a	0 a	0	0 a	2.1 a	2.1 a	27.1 Ab	31.3
Solarization + <i>S. griseoviridis</i> 10^7 cfu m ⁻² in 0.5 l m ⁻²	0 a	0 a	0 a	0	0 a	18.8 b	2.1 a	18.8 A	39.7

418 *Means of the same column followed by the same letter do not differ according to Duncan's

419 Multiple Range Test ($P=0.05$)

420

421 Table 3 – Effect of treatments with *Streptomyces griseoviridis*, soil solarization or their combination
 422 on tomato yield (trial 1, 2001)

Treatment	Mass (g)/fruit	n° fruit/plant	Mass (g)/plant
Inoculated control	110.0	6.09 a	661.1 a
<i>S. griseoviridis</i> 10^7 cfu m ⁻² in 0.5 l m ⁻²	110.2	7.64 a	831.6 a
Solarization	123.8	6.02 a	702.2 a
Solarization + <i>S. griseoviridis</i> 10^7 cfu m ⁻² in 0.5 l m ⁻²	138.8	5.77 a	781.9 a

423 *See Table 2

424

425 Table 4 – Effect of treatments with *Streptomyces griseoviridis*, soil solarization or their combination
 426 on the number of tomato plants infected with *Rhizoctonia solani*, *Fusarium oxysporum* f.sp. *radicis-*
 427 *lycopersici* (FORL), *F. oxysporum* f.sp. *lycopersici* (FOL), *Verticillium dahliae* and *Pyrenochaeta*
 428 *lycopersici* (trial 2, 2001)

Treatment	% plants infected by				
	<i>R. solani</i>	FORL	FOL + <i>V. dahliae</i>	<i>P. lycopersici</i>	Total
Inoculated control	0.0 a*	21.2 a	28.8 b	0.0 a	50.0
Solarization	1.9 a	27.0 a	2.9 a	2.9 a	34.7
<i>S. griseoviridis</i> 10 ⁷ cfu m ⁻² in 0.5 l m ⁻²	1.9 a	22.1 a	23.1 b	1.0 a	48.1
<i>S. griseoviridis</i> 5x10 ⁶ cfu m ⁻² in 0.25 l m ⁻²	4.2 a	25.3 a	24.2 b	1.3 a	55.0
Solarization + <i>S. griseoviridis</i> 10 ⁷ cfu m ⁻² in 0.5 l m ⁻²	0.0 a	26.0 a	7.8 a	1.0 a	34.8

429 *See Table 2

430

431 Table 5 – Effect of treatments with *Streptomyces griseoviridis*, soil solarization or their combination
 432 on the yield of tomato plants (trial 2, 2001)

Treatment	Mass (g) / fruit	n° fruit / plant	Mass (g) / plant	Mass (g) / m ²
Inoculated control	76.7 a*	3.0 a	241.5 a	270 b
Solarization	79.4 a	2.6 b	201.1 b	359 b
<i>S. griseoviridis</i> 10 ⁷ cfu m ⁻² in 0.5 l m ⁻²	83.4 a	3.0 a	251.6 a	314 b
<i>S. griseoviridis</i> 5x10 ⁶ cfu m ⁻² in 0.25 l m ⁻²	84.1 a	3.1 a	253.4 a	326 b
Solarization + <i>S. griseoviridis</i> 10 ⁷ cfu m ⁻² in 0.5 l m ⁻²	85.5 a	3.2 a	272.6 a	428 a

433 *See Table 2

434

435 Table 6 - Effect of treatments with *Streptomyces griseoviridis* and fumigation with metham sodium
 436 on infection of tomato plants with *Fusarium oxysporum* f.sp. *radicis-lycopersici* (FORL), *F.*
 437 *oxysporum* f.sp. *lycopersici* (FOL) and *Pyrenochaeta lycopersici*, and the severity of infection by *P.*
 438 *lycopersici* (trial 3, 2002)

Treatment	% plants infected by			Severity (%) in the tomato roots infected by <i>P. lycopersici</i>	
	FORL	FOL	<i>P. lycopersici</i>	Total	
Inoculated control	5.8a*	1.3a	55.3c	62.4	10.3b
<i>S. griseoviridis</i> 5x10 ⁶ cfu m ⁻² by irrigation ^a	6.3a	1.9a	37.5b	45.7	9ab
<i>S. griseoviridis</i> 10 ⁷ cfu m ⁻² by irrigation ^a	8.2a	0.6a	38.4b	47.2	7.5ab
<i>S. griseoviridis</i> 10 ⁷ cfu m ⁻² by spraying ^b	3.8a	2.5a	46.2bc	52.5	8.2ab
Metham sodium (32.7% a.i.) 76 g a.i. m ⁻²	9.6a	6.3b	7.4a	23.3	5.9a

439 *See Table 2; ^a Five l of water applied per m² immediately before inoculation, 2 with and 3
 440 immediately after applying the commercial product; ^b 0.5 l of water distributed per m² with the
 441 biological product.

442

443 Table 7 – Effect of treatments with *Streptomyces griseoviridis* and fumigation with metham sodium
 444 on tomato yield (trial 3, 2002)

Treatment	Mass (g) / fruit	n° fruit / plant	Mass (g) / plant
Inoculated control	75.4a*	9b	639b
<i>S. griseoviridis</i> 5x10 ⁶ cfu m ⁻² by irrigation ^a	71.7a	12b	881b
<i>S. griseoviridis</i> 10 ⁷ cfu m ⁻² by irrigation ^a	77.4a	16b	1268b
<i>S. griseoviridis</i> 10 ⁷ cfu m ⁻² by spraying ^b	76.3a	14b	1087b
Metham sodium (32.7% a.i.) 76 g a.i. m ⁻²	73.6a	34a	2491a

445 *See Table 2; ^{a,b} See Table 6.

446

447 Table 8 – Effect of treatments with *Streptomyces griseoviridis* and drenching with benomyl on
 448 infection of tomato plants with *Rhizoctonia solani*, *Fusarium oxysporum* f.sp. *radicis-lycopersici*
 449 (FORL), *F. oxysporum* f.sp. *lycopersici* (FOL) (trial 4, 2002)

Treatment	% plants infected by				
	<i>R.solani</i>		FORL	FOL	Total
Inoculated control	3.3	a*	7.6 a	35.9 b	46.8
<i>S. griseoviridis</i> 5x10 ⁶ cfu m ⁻² by irrigation ^a	1.9	a	7.6 a	27.6 ab	37.1
<i>S. griseoviridis</i> 10 ⁷ cfu m ⁻² by irrigation ^a	1.4	a	10.0 a	17.0 ab	28.4
<i>S. griseoviridis</i> 10 ⁷ cfu m ⁻² by spraying ^b	0.6	a	14.5 a	13.9 a	29.0
Benomyl (50% a.i.) 2 g a.i. m ⁻²	1.3	a	14.0 a	21.7 ab	37.0

450 *See Table 2; ; ^{a,b} See Table 6.