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published in

Frontiers in Microbiology
2017

DOI (link to publisher)

[10.3389/fmicb.2017.00447](https://doi.org/10.3389/fmicb.2017.00447)

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Peer reviewed version

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citation for published version (APA)

Bosmans, L., de Bruijn, I., Gerards, S., Moerkens, R., Van Looveren, L., Wittemans, L., Van Calenberge, B., Paeleman, A., Van Kerckhove, S., Rozenski, J., de Mot, R., Rediers, H., Raaijmakers, J. M., & Lievens, B. (2017). Potential for biocontrol of hairy root disease by a *Paenibacillus* clade. *Frontiers in Microbiology*, 8, Article 447. <https://doi.org/10.3389/fmicb.2017.00447>

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Potential for biocontrol of hairy root disease by a *Paenibacillus* clade exhibiting antagonistic activity against rhizogenic *Agrobacterium* biovar 1 strains

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Submitted to Journal:
Frontiers in Microbiology

Specialty Section:
Plant Biotic Interactions

Article type:
Original Research Article

Manuscript ID:
243031

Received on:
16 Nov 2016

Frontiers website link:
www.frontiersin.org

Conflict of interest statement

The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest

Author contribution statement

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Keywords

Agrobacterium, Antagonistic Activity, biological control, High-Throughput Screening, Paenibacillus.

Abstract

Word count: 263

Rhizogenic *Agrobacterium* biovar 1 is the causative agent of hairy root disease (HRD) in the hydroponic cultivation of tomato and cucumber causing significant losses in marketable yield. In order to prevent and control the disease chemical disinfectants such as hydrogen peroxide or hypochlorite are generally applied to sanitize the hydroponic system and/or hydroponic solution. However, effective control of HRD sometimes requires high disinfectant doses that may have phytotoxic effects. Moreover, several of these chemicals may be converted to unwanted by-products with human health hazards. Here we explored the potential of beneficial bacteria as a sustainable means to control HRD. A large collection of diverse bacterial genera was screened for antagonistic activity against rhizogenic *Agrobacterium* biovar 1 using the agar overlay assay. Out of more than 130 strains tested only *Paenibacillus* strains showed antagonistic activity. Strikingly, phylogenetic analysis showed that antagonistic activity was restricted to a particular *Paenibacillus* clade, representing the species *P. illinoisensis*, *P. pabuli*, *P. taichungensis*, *P. tundrae*, *P. tylopili*, *P. xylanexedens* and *P. xylanilyticus*. Assessment of the spectrum of activity revealed that some strains were able to inhibit the growth of all 35 rhizogenic *agrobacteria* strains tested, while others were only active against part of the collection, suggesting a different mode of action. Preliminary characterization of the compounds involved in the antagonistic activity of two closely related *Paenibacillus* strains, tentatively identified as *P. xylanexedens*, revealed that they are water-soluble and have low molecular weight. Application of a combination of these strains in greenhouse conditions resulted in a significant reduction of HRD, indicating the great potential of these strains to control HRD.

Funding statement

We thank VLAIO (Flanders Innovation & Entrepreneurship) to support this research (project IWT-LA: 120761).

Ethics statements

(Authors are required to state the ethical considerations of their study in the manuscript, including for cases where the study was exempt from ethical approval procedures)

Does the study presented in the manuscript involve human or animal subjects: No

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28 **ABSTRACT**

29 Rhizogenic *Agrobacterium* biovar 1 is the causative agent of hairy root disease (HRD)
30 in the hydroponic cultivation of tomato and cucumber causing significant losses in
31 marketable yield. In order to prevent and control the disease chemical disinfectants such
32 as hydrogen peroxide or hypochlorite are generally applied to sanitize the hydroponic
33 system and/or hydroponic solution. However, effective control of HRD sometimes
34 requires high disinfectant doses that may have phytotoxic effects. Moreover, several of
35 these chemicals may be converted to unwanted by-products with human health hazards.
36 Here we explored the potential of beneficial bacteria as a sustainable means to control
37 HRD. A large collection of diverse bacterial genera was screened for antagonistic
38 activity against rhizogenic *Agrobacterium* biovar 1 using the agar overlay assay. Out of
39 more than 130 strains tested only *Paenibacillus* strains showed antagonistic activity.
40 Strikingly, phylogenetic analysis showed that antagonistic activity was restricted to a
41 particular *Paenibacillus* clade, representing the species *P. illinoisensis*, *P. pabuli*, *P.*
42 *taichungensis*, *P. tundrae*, *P. tylopili*, *P. xylanexedens* and *P. xylanilyticus*. Assessment
43 of the spectrum of activity revealed that some strains were able to inhibit the growth of
44 all 35 rhizogenic agrobacteria strains tested, while others were only active against part
45 of the collection, suggesting a different mode of action. Preliminary characterization of
46 the compounds involved in the antagonistic activity of two closely related *Paenibacillus*
47 strains, tentatively identified as *P. xylanexedens*, revealed that they are water-soluble
48 and have low molecular weight. Application of a combination of these strains in
49 greenhouse conditions resulted in a significant reduction of HRD, indicating the great
50 potential of these strains to control HRD.

51

52 **Key words:** *Agrobacterium*, antagonistic activity, biological control, high-throughput
53 screening, *Paenibacillus*.

54

55 INTRODUCTION

56 Since the early 1990s, in several European countries hydroponically grown
57 cucumber plants and tomato crops have been affected by a disorder called ‘hairy root
58 disease’ (HRD). The disease is characterized by extensive root proliferation leading to
59 strong vegetative growth and, in severe cases, substantial losses in marketable yield
60 (Weller *et al.*, 2006; Ludeking *et al.*, 2013). In hydroponic crops HRD is generally
61 associated with rhizogenic *Agrobacterium* biovar 1 strains (further referred to as
62 ‘rhizogenic agrobacteria’), harbouring a Ri-plasmid (root-inducing plasmid) (Gelvin,
63 2003). Symptoms arise following transfer of a portion of the Ri-plasmid (T-DNA;
64 transferred DNA) from the bacterium to plant cells, where it is integrated in the
65 chromosomal DNA and subsequently expressed (Hooykaas and Beijersbergen, 1994),
66 leading to excessive root development. Once plants are infected, HRD cannot be
67 controlled by curative means and rather preventative actions should therefore be taken,
68 such as preventing and/or removing *Agrobacterium* containing biofilms that are often
69 associated with the disease in the greenhouse irrigation system (Danhorn *et al.*, 2007;
70 Bosmans *et al.*, 2015). However, to effectively prevent the disease generally high
71 concentrations of chemical disinfectants are required, including levels that may be
72 phytotoxic (Bosmans *et al.*, 2016c). Moreover, several of these chemicals may be
73 converted to unwanted by-products with human health hazards (Damstra, 2002).
74 Therefore, there is currently a strong interest in alternative means to prevent and control
75 HRD such as the use of biocontrol organisms (BCO).

76 The use of BCO has received great attention the last few decades because of the
77 ability of such antagonistic strains to suppress plant diseases with less environmental
78 impact than chemical pesticides, reduced off-target effects in microbiota linked to a
79 narrow activity spectrum, and the possibility to be integrated with other control methods
80 (Raaijmakers *et al.*, 2002; Rubino *et al.*, 2013). Especially rhizosphere bacteria are
81 generally considered ideal BCO of soilborne plant pathogens because of their effective
82 colonization of the rhizosphere providing a front-line defence against pathogen attack,
83 their versatility to protect plants under different conditions, and production of
84 antimicrobial compounds (Sharma *et al.*, 2009). However, so far, no bacterial
85 antagonists have been identified to control rhizogenic *Agrobacterium* biovar 1.

86 The objectives of this study were (i) to identify potential bacterial BCO of
87 rhizogenic agrobacteria using both laboratory and greenhouse experiments and (ii) to
88 perform a preliminary characterization of the compounds involved in the antagonistic
89 activity. To this end, a large collection of diverse bacterial isolates from rhizosphere soil
90 was screened for antagonistic activity using the agar overlay assay. Antimicrobial
91 compounds were determined using RP-HPLC and a quadrupole orthogonal acceleration
92 time-of-flight mass spectrometer. Further, biocontrol activity of a mixture of the most
93 promising strains was assessed under greenhouse conditions.

94

95 **MATERIALS AND METHODS**

96 **Culture collection and screening for antagonists of rhizogenic agrobacteria**

97 In a first screening, a collection of 130 phylogenetically different bacterial
98 strains isolated from soil habitats (de Ridder-Duine *et al.*, 2005) was used in this study
99 and subjected to high-throughput screening for antagonists of rhizogenic agrobacteria as

100 described previously (Tyc *et al.*, 2014) (Table S1, Supporting Information). The
101 collection consisted of strains from different phyla and different classes (Table 1), and
102 has previously been evaluated for antagonistic activity against two human pathogenic
103 model organisms, including *Escherichia coli* and *Staphylococcus aureus* (Tyc *et al.*,
104 2014). Additionally, *Streptomyces rimosus* DSM40260, a producer of oxytetracycline,
105 was included in the study as a reference strain. Strains were stored in glycerol at -80 °C
106 in two 96-well plates until further use. To this end, first wells of the 96-well plates were
107 filled with 150 µl lysogeny broth (LB) (10 g/L NaCl, 10 g/L Bacto™ Tryptone, 5 g/L
108 Bacto™ Yeast extract) and inoculated with the strains. Plates were then incubated for
109 two days at 25 °C with gentle agitation, after which 50 µl of 50 % (v/v) glycerol was
110 added to achieve a final glycerol concentration of 12.5 % (v/v).

111 For assessing the antagonistic properties of the collection, the 96-well plates
112 were thawed and isolates were spotted using the Genetix QPix 2 colony picking robot
113 (Molecular Devices, UK Limited, Wokingham, United Kingdom) on OmniTray-plates
114 (size 128 × 86 mm; capacity 90 mL; Greiner Bio-One B.V., Alphen a/d Rijn, The
115 Netherlands) with 15 mL solid bacterial growth medium (5 g/L NaCl, 1 g/L KH₂PO₄; 3
116 g/L Oxoid Tryptic Soy Broth (TSB); 20 g/L Merck Agar-agar). The plates were
117 incubated for 5 days at 20 °C and were used as source plates for spotting test plates
118 containing the same medium mentioned above. Importantly, Merck agar-agar was used
119 in our screening as this agar was shown to support bacterial antagonistic activity against
120 rhizogenic agrobacteria, while several other agars were not (Bosmans *et al.*, 2016b).
121 Spot-inoculated OmniTray plates were then incubated for 1 day at 25 °C. Subsequently,
122 15 mL melted LB agar containing *Agrobacterium* (about 6 × 10⁵ cells per mL) was
123 poured over the surface of the plate and incubated again at 25 °C. After 24 hours of
124 incubation, the diameter of the inhibition visible zones surrounding spotted colonies

125 was recorded (Tyc et al. 2014; Bosmans et al. 2016b). Experiments were performed for
126 one rhizogenic *Agrobacterium* biovar 1 strain (ST15.13/097, isolated from tomato;
127 Bosmans *et al.*, 2015), and were independently repeated twice.

128 In a second screening, several strains from the same genus as the only strain
129 showing antagonistic activity in the initial high-throughput screening mentioned above
130 (i.e. *Paenibacillus*) (Table 2) were evaluated for antagonistic activity against
131 *Agrobacterium* biovar 1 strain ST15.13/097 in an agar overlay assay using 9 cm-
132 diameter petri dishes as described by Bosmans *et al.* (2016b). For all strains showing
133 antagonistic activity the spectrum of activity was evaluated using 35 rhizogenic
134 *Agrobacterium* biovar 1 strains and 37 other strains from diverse phyla including
135 Actinobacteria, Firmicutes and Proteobacteria, among which several plant pathogens
136 (Table 3).

137

138 **Characterization of antagonistic strains**

139 For all strains with antagonistic activity the 16S ribosomal RNA (rRNA) genes
140 were partially amplified and sequenced as described by Bosmans *et al.* (2015). Obtained
141 sequences were individually trimmed for quality, using a minimum Phred score of 20,
142 and, in cases of ambiguous base calls, manually edited based on the obtained
143 electropherograms. A maximum likelihood tree was constructed using MEGA v5.2
144 (Tamura *et al.*, 2011) to assess the phylogenetic relatedness between the antagonistic
145 strains as well as their phylogenetic relationships with previously characterized
146 reference (type) strains for which the sequences were retrieved from EzTaxon
147 (www.ezbiocloud.net/eztaxon).

148 Antagonistic strains were subjected to a Bioscreen C analysis (Oy Growth
149 Curves Ab Ltd, Helsinki, Finland) to assess growth characteristics in different media.

150 The working volume in the wells of the Bioscreen plate was 200 μL , comprised of 5 μL
151 bacterial suspension (about 10^5 cells per mL LB medium) and 195 μL of one of the
152 following three media: TSB (Oxoid, Basingstoke, UK), LB and a minimal broth
153 medium (M70) containing 2 g/L Bacto™ Yeast extract and 10 g/L Mannitol (Sigma,
154 Missouri, US). The temperature was controlled at 25 °C, and the optical density of the
155 cell suspensions was measured automatically at 600nm in regular intervals of 15 min,
156 for three days. Before each measurement, the Bioscreen plate was automatically shaken
157 for 60 seconds. The experiments were performed two times independently, each with
158 three replicates. Tested culture medium without inoculum was used as a reference.
159 Growth curves were generated by monitoring the averaged optical density (OD_{600}) as a
160 function of incubation time.

161

162 **Preliminary characterization of the antagonistic compound(s)**

163 The two best performing strains (based on the size of the zone of inhibition,
164 specificity and growth in the previous assays), including AD117 (the same as
165 ST15.13/036, Bosmans *et al.*, 2016b) and ST15.15/027, were selected for preliminary
166 characterization of the antagonistic compounds. First, isolates were investigated for
167 production of volatile organic compounds (VOCs) having antagonistic activity against
168 *Agrobacterium*. To this end, two bottoms of a 9 cm-diameter petri dishe, one containing
169 a freshly spot-inoculated (15 μL per spot; about 10^5 cells per mL in TSB) antagonistic
170 bacterium (on the medium described above) and the other a rhizogenic *Agrobacterium*
171 biovar 1 isolate (ST15.13/097) (on TSA, Oxoid, Basingstoke, UK), were sealed facing
172 each other and incubated at 25 °C with the petri-dish containing the antagonistic
173 bacterium at the bottom. The experiments were carried out using two independent

174 repeats, each with three replicates. Growth inhibition was calculated by measuring the
175 zone of inhibition after 1, 2 and 3 days of incubation.

176 Secondly, to assess whether the antagonistic compounds are secreted to the
177 extracellular space, cell-free culture filtrates were prepared and tested for antibacterial
178 activity in a microtitre plate (Thermo Scientific™ Nunc™ MicroWell™ 96-Well
179 Microplates). To this end, antagonistic bacteria were cultured in liquid medium (100
180 mL) consisting of 3 g/L tryptic soy broth (TSB; Oxoid, Basingstoke, UK), 5 g/L NaCl,
181 and 1 g/L KH₂PO₄, and incubated at 25 °C for 2 days. Cultures of about 10⁴ cells per
182 mL were then filter-sterilized (0.2-µm filter, sterile mixed cellulose ester membrane,
183 Whatman, GE Healthcare Life Sciences, UK), and a portion of the filtrate was added to
184 the wells of the microtiter plate. More specifically, 100, 150 and 190 µL of the cell-free
185 filtrates were added to 100, 50 and 10 µL LB containing *Agrobacterium* biovar 1 isolate
186 ST15.13/097 (final concentration of 5 x 10² cells per mL for each condition),
187 respectively. In the control wells, the culture filtrate was replaced by LB. For all
188 treatments, plates were incubated with gentle agitation and growth was
189 photospectrometrically (OD₆₀₀) quantified after 24 h of incubation at 25 °C.
190 Experiments were independently repeated twice.

191

192 **Extraction and purification of the antagonistic compound(s)**

193 For the extraction and identification of the compounds responsible for the
194 antagonistic activity, the two best performing strains, AD117 and ST15.15/027, were
195 selected and spot-inoculated (15 µL per spot) on the agar medium mentioned above in 9
196 cm-diameter petri dishes (60 plates per strain). Following inoculation with
197 *Agrobacterium* (isolate ST15.13/097) (see above) and subsequent incubation for 1 day
198 at 25 °C, 60 agar pieces of approximately 1 cm² were excised from the zone of

199 inhibition, suspended in 65% methanol (65% methanol, 34.9% milliQ water and 0.1%
200 formic acid) and shaken for 3 h at room temperature. After centrifugation at 5000 g for
201 15 min, the liquid phase was transferred and the methanol was evaporated by air drying.
202 Subsequently, the aqueous phase was frozen and freeze-dried, and the dried extract was
203 dissolved again in 65% methanol prior to further analysis. Obtained extracts were
204 analysed by reversed-phase high-performance liquid chromatography (RP-HPLC)
205 (Waters Chromatography B.V., Etten-Leur, the Netherlands) equipped with a Waters
206 996 photodiode array detector. The separations were performed on a Waters Symmetry
207 C18RP column (5 μ m, 3.9 \times 150 mm) with a mobile phase of 70% methanol and
208 0.1% formic acid, and operated at a flow of 0.2 mL/min for 10 min (or 60 min for
209 improved resolution of peaks) with UV detection at 240 nm. Fractions were collected
210 each 5 min or by collecting particular peaks. For each collected fraction, methanol was
211 evaporated and the remaining (aqueous) phase was freeze-dried, dissolved again in 65%
212 methanol, and 20 μ L was spotted on a sterile filter paper and covered by an
213 *Agrobacterium* overlay. 20 μ L methanol, spotted on filter paper was used as an control.

214 For those HPLC fractions that had activity against *Agrobacterium*, mass spectra
215 were acquired in positive ionization mode on a quadrupole orthogonal acceleration
216 time-of-flight mass spectrometer (Syntapt G2, Waters, Milford, MA) equipped with a
217 standard electrospray probe and controlled by the MassLynx 4.1 software. Resolution of
218 the instrument was set to 15000 (resolution mode). The capillary voltage and cone
219 voltage were set to 3 kV and 35 V, respectively. Accurate masses were obtained using
220 the LockSpray source and leucine enkephalin (2 ng/ μ L in acetonitrile:water 1:1) as
221 reference compound infused at 3 μ L/min. The chromatographic system consisted of an
222 ultra-performance liquid chromatography (UPLC) system (Acquity H-class, Waters,
223 Milford, MA). Separations were performed on a reversed phase C18 column (Acquity

224 HSS T3 1.8 μm 1x50 mm) at a flow rate of 150 $\mu\text{L}/\text{min}$. The injection volume was 5
225 μL . A linear gradient of acetonitrile in water (2 to 22% in 10 min) was applied. Mass
226 spectra in the mass range m/z 100 to 700 were acquired at a rate of one spectrum per
227 second.

228

229 **Evaluation of the antagonistic activity in greenhouse conditions**

230 A greenhouse experiment was performed to assess the biocontrol activity of a
231 mixture of the two selected bacteria (AD117 and ST15.15/027) against *Agrobacterium*
232 biovar 1 in a commercial hydroponic tomato production system in Belgium (Research
233 Centre Hoogstraten, Belgium). Experiments were performed using the tomato cultivar
234 ‘Rebelski’ (De Ruiter, The Netherlands), rootstock Maxifort (De Ruiter, The
235 Netherlands). Four plants were planted in one rockwool mat with a plant density of 2.5
236 plants / m^2 . From the start of the experiment, i.e. from the moment of planting of ~60-
237 day-old tomato seedlings (January 2016), a set of 20 plants (5 rockwool mats) were
238 treated by adding a mixture of 50 mL of the two candidate BCO (10^8 cells/mL each) to
239 the rockwool mat daily for 10 days, while another set of 20 plants remained untreated.
240 From day ten of the experiment, all 40 plants were artificially infected by applying a
241 rhizogenic *Agrobacterium* biovar 1 strain (isolate ST15.13/097) (50 mL of a suspension
242 of 10^8 cells/mL) once a week for a total of six weeks to the rockwool mats. Plants were
243 visually evaluated every two weeks for a total examination period of eight weeks (until
244 17 weeks after infection) for development of aberrant root formation. Plants were
245 considered infected when visual HRD symptoms were confirmed by a positive qPCR
246 analysis of the pathogen from investigated root material (Bosmans *et al.*, 2016a).

247

248 **RESULTS**

249

250 **Antagonistic activity against rhizogenic agrobacteria**

251 Out of 130 tested bacterial strains belonging to different phyla and different
252 classes, *Paenibacillus* strain AD117 showed antibacterial activity against the tested
253 rhizogenic *Agrobacterium* strain (ST15.13/097) (Table 1 and Table S1, Supporting
254 Information). Additional screening of other *Paenibacillus* strains resulted in four
255 additional antagonistic strains, including the type strain of *Paenibacillus xylanilyticus*
256 (DSM17255^T) and three *Paenibacillus* strains that were not yet assigned to the species
257 level (ST15.15/027, ST15.15/031 and ST15.15/032) (Table 2). Overall, for these strains
258 the average diameter of the inhibition zones varied between 1.57 cm and 2.88 cm, with
259 the largest zones of inhibition for strains AD117 (2.88 cm) and ST15.15/027 (2.79 cm)
260 (Fig S1, Supporting Information). 16S rRNA gene sequence analysis using the
261 EZTaxon database showed that the strains AD117, ST15.15/027, ST15.15/031 and
262 ST15.15/032 had highest sequence homology with *Paenibacillus illinoisensis*
263 (ST15.15/031 and ST15.15/032) and *P. xylanexedens* (AD117 and ST15.15/027) (Table
264 2). Examination of the growth characteristics of the five selected strains revealed
265 highest growth rates for AD117, DSM17255^T and ST15.15/027, irrespective of the
266 growth medium used (data not shown). Phylogenetic analysis with all validly named
267 *Paenibacillus* species (163 species) revealed that these five strains clustered tightly with
268 *P. illinoisensis*, *P. xylanilyticus*, *P. taichungensis*, *P. pabuli*, *P. tundra*, *P. tylopili* and *P.*
269 *xylanexedens* (Fig. 1). When also the type strains of these species were subjected to the
270 agar overlay assay, all strains demonstrated antagonistic activity, while strains that were
271 less related to this cluster did not (Table 2).

272 Assessment of the spectrum of antagonistic activity of strains AD117,
273 DSM15255^T, ST15.15/027, ST15.15/031 and ST15.15/032 revealed that three strains

274 (AD117, DSM17255^T and ST15.15/027) showed antagonistic activity against all
275 rhizogenic *Agrobacterium* biovar 1 strains (35) tested (Table 3). In contrast, the isolates
276 corresponding to *P. illinoisensis*, ST15.15/031 and ST15.15/032, showed a different
277 activity spectrum and were only able to inhibit the growth of 19 and 17 *Agrobacterium*
278 biovar 1 strains, respectively (Table 3). Furthermore, strains AD117, DSM17255^T and
279 ST15.15/027 were able to suppress the growth of one or more rhizogenic *Agrobacterium*
280 biovar 2 strains causing HRD on Rosaceae. Additionally, strain ST15.15/027 showed
281 antagonistic activity against *Rhizobium vitis* LMG256, a plant pathogen causing crown
282 gall of grapevine (Table 3).

283

284 **Preliminary characterization of the antagonistic compound(s)**

285 Based on the results described above (size of the zone of inhibition, spectrum of
286 activity and general growth characteristics), both AD117 and ST15.15/027 were
287 selected for further experiments to identify the active substances mediating the
288 antagonistic effects observed. First, strains were evaluated for the production of volatile
289 organic compounds (VOCs) with antagonistic activity against rhizogenic agrobacteria,
290 but no VOC-dependent activity could be detected. In contrast, when the cell-free culture
291 filtrates were tested, a dose-dependent growth inhibition of *Agrobacterium* was
292 observed (Fig. 2), suggesting that the selected bacteria secrete water-soluble
293 antibacterial compounds. HPLC fractionation of an extract from the agar cut from the
294 inhibition zones in the agar overlay assay was performed and gave one fraction with
295 antagonistic activity. For each isolate, mass spectrometry analysis of this HPLC fraction
296 showed the presence of four specific peaks having a mass number of $m/z = 463.2030$,
297 477.1830, 504.2669 and 578.2324.

298

299 **Greenhouse experiments**

300 A mixture of AD117 and ST15.15/027 was evaluated for its biocontrol potential
301 of rhizogenic agrobacteria in greenhouse conditions. To this end, two sets of 20 plants
302 were scored weekly for development of excessive root formation. Nine weeks after
303 artificial infection with *Agrobacterium*, the first symptoms of HRD were observed.
304 After 17 weeks about 75% of all control plants artificially infected with *Agrobacterium*
305 showed HRD. When plants were treated with a mixture of AD117 and ST15.15/027
306 incidence of HRD dropped to 45% (Fig. 3), suggesting high biocontrol potential of the
307 used inoculum. Observation of HRD symptoms was always confirmed by a positive
308 qPCR analysis targeting *Agrobacterium* biovar 1 DNA.

309

310 **DISCUSSION**

311 HRD caused by rhizogenic *Agrobacterium* biovar 1 strains is an economically
312 important disease in the hydroponic cultivation of cucurbits and tomato leading to
313 significant losses in marketable yield. As different lineages of rhizogenic
314 *Agrobacterium* strains are able to form biofilms in which they can be protected from
315 chemical disinfectants (Bosmans *et al.*, 2015), or are able to tolerate high disinfectant
316 concentrations (Bosmans *et al.*, 2016c) or even diverse antibiotics (Khodykina *et al.*,
317 2014), there is an urgent need for alternative, effective means to prevent and control the
318 disease including the use of biocontrol organisms.

319 After an extensive evaluation of a diverse bacterial collection several
320 *Paenibacillus* strains were found to have antagonistic activity against rhizogenic
321 *Agrobacterium* biovar 1 strains. Antagonistic strains included the type strain of *P.*
322 *xylanilyticus* (DSM17255^T), two strains putatively identified as *P. illinoisensis*
323 (ST15.15/031 and ST15.15/032) and two strains putatively identified as *P. xylanexedens*

324 (AD117 and ST15.15/027). *Paenibacillus* species have been isolated from various
325 ecological habitats including soil, air, rhizosphere, and extreme environments such as
326 floral plant nectar, warm water springs and glaciers (McSpadden Gardener, 2004;
327 Jacquemyn *et al.*, 2013). The wide range of habitats from which the identified strains
328 have been previously isolated include air (DSM17255^T), rhizosphere (AD117; de
329 Ridder-Duine *et al.*, 2005), malting wheat kernels (ST15.15/031, ST15.15/032 (Malfliet
330 *et al.*, 2013)), and oak bourbon casks used to age beer (ST15.15/027), which suggests
331 that antagonistic activity against rhizogenic agrobacteria is not related to the original
332 (natural) habitat of the strains, and that antagonistic activity is not dependent on a
333 history of previous contact with the pathogen (see also Duffy *et al.*, 2003). However,
334 positioning of these strains in a phylogenetic tree containing 16S rRNA gene sequences
335 of the reference (type) strains of all validly named *Paenibacillus* species revealed that
336 these five strains clustered tightly together with the type strains of *P. illinoisensis*, *P.*
337 *xylanilyticus*, *P. taichungensis*, *P. pabuli*, *P. tundra*, *P. tylopili* and *P. xylanexedens*.
338 Similar results were obtained when a phylogenetic analysis was performed using *rpoB*
339 sequences (encoding the β subunit of the bacterial RNA polymerase) (although less
340 sequences were available for type strains; data not shown), confirming their close
341 phylogenetic relatedness. When also these strains were subjected to the agar overlay
342 assay, they all exhibited antagonistic activity against rhizogenic agrobacteria,
343 suggesting phylogenetic conservation in antagonistic activity. Also other studies have
344 reported on a correlation between antimicrobial activity and phylogeny. For example,
345 Satheeja and Jebakumar (2011) showed that the antimicrobial activities of *Streptomyces*
346 isolates were linked to their phylogenetic position. Likewise, Wilson *et al.* (2010) found
347 a correlation between the antimicrobial activities of marine bacteria and the phylogeny
348 of the isolates investigated. Several studies have shown antagonistic properties of

349 *Paenibacillus* species or demonstrated their potential as biocontrol agents to control
350 plant diseases caused by bacteria, fungi and oomycetes (Tjamos *et al.*, 2004; Jung *et al.*,
351 2005; Haggag and Timmusk, 2008; Timmusk *et al.*, 2009; Algam *et al.*, 2010; Sato *et*
352 *al.*, 2014). However, to the best of our knowledge, our study is the first in which a
353 correlation was found between a distinct phylogenetic clade and antagonistic activity
354 against a particular bacterial pathogen. All antagonistic strains were found to have the
355 following 16S rRNA gene signature sequence differentiating antagonistic from non-
356 antagonistic strains: 5'-
357 TTGGGACAACACTACCGGAAACGGTAGCTAATACCGAATA-3'.

358 Strikingly, differences were observed between the activity spectrum of the
359 phylogenetically-clustered antagonistic *Paenibacillus* strains. More particularly, while
360 isolates AD117, DSM17255^T and ST15.15/027 showed antagonistic activity against all
361 rhizogenic *Agrobacterium* biovar 1 isolates tested (35 isolates), the two isolates
362 identified as *P. illinoisensis*, (ST15.15/031 and ST15.15/032) showed a different
363 activity spectrum inhibiting the growth of different *Agrobacterium* strains and were
364 only antagonistic against part (approximately 50%) of the strains tested. Rhizogenic
365 *Agrobacterium* biovar 1 comprises a group of different genetic lineages exhibiting
366 substantial genetic diversity (Bosmans *et al.*, 2015). Nevertheless, no correlation could
367 be found between the genetic background of the tested *Agrobacterium* strains and their
368 vulnerability/resistance to these *Paenibacillus* strains. This also suggests that different
369 modes of action are at play explaining antagonistic activity against rhizogenic
370 *Agrobacterium* biovar 1. Interestingly, strains AD117, DSM17255^T and ST15.15/027
371 were also able to suppress the growth of one or more rhizogenic *Agrobacterium* biovar 2
372 strains, causing HRD on other crops such as Rosaceae (Cervera *et al.*, 1998).
373 Additionally, strain ST15.15/027 showed antagonistic activity against *Rhizobium vitis*.

374 Preliminary characterization of the antagonistic compounds of AD117 and ST15.15/027
375 revealed that the compounds are water-soluble molecules of low molecular weight
376 (<600 Da). There also seems to be an important role of Ca²⁺ to produce and/or secrete
377 potential toxins/antibiotics against rhizogenic agrobacteria (Bosmans *et al.*, 2016c).
378 Further research, however, is necessary to structurally identify and characterize these
379 compounds. The fact that they are water-soluble opens perspectives towards their
380 application and efficacy in hydroponic systems. Indeed, when the paenibacilli were
381 evaluated in a commercial hydroponic tomato production system, a significant reduction
382 in incidence of HRD (45% versus 75% for the control treatment) was obtained when
383 plants were evaluated over a period of about 4 months. Although these results are highly
384 promising, it can be assumed that biocontrol efficacy can even be enhanced by frequent
385 application of the BCO.

386 Altogether, we have shown that *Paenibacillus* holds great potential to control
387 HRD. Furthermore, we have shown that its antagonistic activity against rhizogenic
388 agrobacteria is related with the phylogeny of the *Paenibacillus* strains, but not with the
389 phylogeny of the agrobacteria. Together with its plant-growth promoting traits (Lamsal
390 *et al.*, 2013), this makes *Paenibacillus* an excellent candidate for practical applications
391 in the hydroponic cultivation of cucurbits and tomato crops.

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394 **Acknowledgements**

395 We thank VLAIO (Flanders Innovation & Entrepreneurship) to support this research
396 (project IWT-LA: 120761).

397

398 **Conflict of Interest**

399 No conflict of interest declared.

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In review

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520 **FIGURES LEGENDS**

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522 **Figure 1.** Phylogenetic positioning of *Paenibacillus* strains showing antagonistic
523 activity against rhizogenic *Agrobacterium* biovar 1 strains. A maximum likelihood
524 (ML) tree was constructed based on 16S rRNA gene sequences (1390 bp) for all
525 reference (type) strains of all *Paenibacillus* species (EZtaxon) currently described (163
526 species) and all other *Paenibacillus* strains included in this study (Table 2). Only
527 members of a tight cluster of *Paenibacillus* strains were found to have antagonistic
528 activity against rhizogenic agrobacteria, while strains that were less related to this
529 cluster were not antagonistic. *Paenibacillus* strains that were tested for antagonistic
530 activity against *Agrobacterium* biovar 1 (isolate ST15.13/097) are marked with a green
531 or red dot, representing antagonistic or no antagonistic strains, respectively. Strains
532 without coloured dots were not tested for antagonistic activity against *Agrobacterium*
533 biovar 1. Major bootstrap values (> 85 %; 1000 replications) are shown at the nodes of
534 the tree.

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541 **Figure 2.** Antagonistic activity of cell-free culture filtrates of selected *Paenibacillus*
542 strains (AD117 and ST15.15/027) grown in LB against rhizogenic *Agrobacterium*
543 biovar 1 (isolate ST15.13/097). *Paenibacillus* cultures of 10^4 cells per mL were filter-
544 sterilized and 100 μ L (blue), 150 μ L (red) and 190 μ L (green) of the cell-free filtrates
545 were added to 100, 50 and 10 μ L *Agrobacterium*-containing LB (final concentration of
546 5×10^2 cells per mL), respectively. The yellow bar represents the control treatment (200
547 μ L LB medium; no culture filtrate). Bacterial growth ((OD₆₀₀) was measured after 24 h
548 of incubation at 25 °C. Presented data are means of two independent experiments (two
549 replicates per experiment) and error bars represent standard error of the mean. The
550 asterisk indicates a statistically significant difference (Student t-test) with the control (P
551 < 0.05).

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564 **Figure 3.** Biocontrol activity of a mixture of *Paenibacillus* strains (AD117 and
565 ST15.15/027) against rhizogenic agrobacteria causing HRD (isolate ST15.13/097) in

566 greenhouse conditions. Incidence of HRD (calculated as the ratio of infected tomato
567 plants) is plotted in function of time (weeks after initial infection with *Agrobacterium*):
568 red, control plants (n=20); green, plants treated with the BCO mixture (n=20). Since day
569 10 of the experiment, all hydroponically grown plants were weekly infected with
570 *Agrobacterium* (isolate ST15.13/097) for six weeks in total. Plants were visually
571 evaluated every two weeks for development of excessive root formation. Observation of
572 symptoms was confirmed by a positive qPCR analysis specifically targeting
573 *Agrobacterium* biovar 1 DNA.

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591 **TABLES**

592 **Table 1.** Overview of antagonistic activity screening of 130 bacterial soil isolates^a against rhizogenic *Agrobacterium* biovar 1 (strain
 593 ST15.13/097)^b. For more details, the reader is referred to Table S1 (Supporting Information).

Phylum / Class	Number of strains tested	Strains with antagonistic activity
Actinobacteria		
Actinobacteria	9	0
Bacteroidetes		
Flavobacteria	15	0
Sphingobacteria	1	0
Firmicutes		
Bacilli	7	1 ^c
Proteobacteria		
Alpha-proteobacteria	12	0
Beta-proteobacteria	61	0
Gamma-proteobacteria	25	0
Total	130	1

594 ^aThe collection consisted of 130 isolates from soil habitats (de Ridder-Duine *et al.*, 2005) and has previously been evaluated for antagonistic
 595 activity against *Escherichia coli* and *Staphylococcus aureus* (Tyc *et al.*, 2014).

596 ^bAntagonistic activity was evaluated using the agar overlay assay (Bosmans *et al.*, 2016b). The strain with antagonistic activity produced a clear
 597 zone of inhibition where *Agrobacterium* growth was inhibited.

598 *Paenibacillus* sp. AD117.

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In review

Table 2. Antagonistic activity^a of diverse *Paenibacillus* strains against rhizogenic *Agrobacterium* biovar 1 (strain ST15.13/097).

Isolate^b	<i>Paenibacillus</i>	Antagonistic activity
DSM5050 ^T	<i>Paenibacillus alginolyticus</i>	-
DSM15478	<i>Paenibacillus barcinonensis</i>	+
DSM13188 ^T	<i>Paenibacillus borealis</i>	-
DSM17253 ^T	<i>Paenibacillus favisporus</i>	-
DSM22343 ^T	<i>Paenibacillus glacialis</i>	-
LMG12239 ^T	<i>Paenibacillus glucanolyticus</i>	-
DSM17608 ^T	<i>Paenibacillus glycanilyticus</i>	-
DSM15220 ^T	<i>Paenibacillus graminis</i>	-
LMG23886 ^T	<i>Paenibacillus humicus</i>	-
DSM13815 ^T	<i>Paenibacillus jamilae</i>	-
DSM7030	<i>Paenibacillus larvae</i>	-
LMG6324 ^T	<i>Paenibacillus macerans</i>	-
LMG6935 ^T	<i>Paenibacillus macquariensis</i>	-
LMG15970	<i>Paenibacillus pabuli</i>	+
ST15.15/027	<i>Paenibacillus</i> sp. ^c	+
ST15.15/031	<i>Paenibacillus</i> sp. ^d	+
ST15.15/032	<i>Paenibacillus</i> sp. ^e	+
AD117	<i>Paenibacillus</i> sp. ^f	+
DSM19942	<i>Paenibacillus taichungensis</i>	+
DSM7262 ^T	<i>Paenibacillus thiaminolyticus</i>	-
DSM21291	<i>Paenibacillus tundrae</i>	+

DSM18927	<i>Paenibacillus tylopili</i>	+
LMG9817 ^T	<i>Paenibacillus validus</i>	-
DSM16970 ^T	<i>Paenibacillus xinjiangensis</i>	-
DSM17255	<i>Paenibacillus xylanilyticus</i>	+

601 ^aAntagonistic activity was evaluated using the agar overlay assay (Bosmans *et al.*, 2016b). Strains with antagonistic activity produced a clear
602 zone of inhibition where *Agrobacterium* growth was inhibited (+). -, no inhibition zone observed.

603 ^bAD, NIOO culture collection, Wageningen, The Netherlands; DSM, Deutsche Sammlung von Mikroorganismen und Zellkulturen,
604 Braunschweig, Germany; LMG, Laboratory of Microbiology, Ghent University, Ghent, Belgium; ST, PME&BIM culture collection, Sint-
605 Katelijne Waver, Belgium.

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607 ^crRNA gene analysis (1390 bp) using EzTaxon revealed highest sequence identity (99.65%) with *Paenibacillus xylanexedens* DSM21292^T
608 (GenBank Accession N° EU558281).

609 ^drRNA gene analysis (1390 bp) using EzTaxon revealed highest sequence identity (99.88%) with *Paenibacillus illinoisensis* NBRC15959^T
610 (GenBank Accession N° AB681007).

611 ^erRNA gene analysis (1390 bp) using EzTaxon revealed highest sequence identity (99.72%) with *Paenibacillus illinoisensis* NBRC15959^T
612 (GenBank Accession N° AB681007).

613 ^frRNA gene analysis (1390 bp) using EzTaxon revealed highest sequence identity (99.85%) with *Paenibacillus xylanexedens* DSM21292^T
614 (GenBank Accession N° EU558281).

Table 3. Activity spectrum of selected *Paenibacillus* strains^a.

Phylum / Class	Species	Isolate ^b	Antagonistic activity					
			AD117	ST15.15/027	DSM17255	ST15.15/031	ST15.15/032	
Actinobacteria								
Actinobacteria	<i>Mycobacterium peregrinum</i>	LMG19256	-	-	-	-	-	
Bacteroidetes								
Flavobacteria	<i>Flavobacterium breve</i>	ST01.08/026	-	-	-	-	-	
Firmicutes								
Bacilli	<i>Bacillus amyloliquefaciens</i>	ST12.14/143	-	-	-	-	-	
	<i>Bacillus bataviensis</i>	EMI_2_2	-	-	-	-	-	
	<i>Bacillus endophyticus</i>	EMI_1_27	-	-	-	-	-	
	<i>Bacillus megaterium</i>	EMI_2_14	-	-	-	-	-	
	<i>Bacillus muralis</i>	EMI_1_24	-	-	-	-	-	
	<i>Bacillus pumilus</i>	ST12.14/241	-	-	-	-	-	
	<i>Bacillus subtilis</i>	ST01.08/012	-	-	-	-	-	
	<i>Bacillus thuringiensis</i>	ST12.14/323	-	-	-	-	-	
	<i>Staphylococcus aureus</i>	ST01.08/020	-	-	-	-	-	
Proteobacteria								
Alpha-proteobacteria	<i>Agrobacterium tumefaciens</i>	LMG187	-	-	-	-	-	
	<i>Rhizobium larrymoorei</i>	LMG21410	-	-	-	-	-	
	<i>Rhizobium meliloti</i>	LMG4290	-	-	-	-	-	
	<i>Rhizobium rubi</i>	LMG294	-	-	-	-	-	
	<i>Rhizobium vitis</i>	LMG256	-	+	-	-	-	
	Rhizogenic <i>Agrobacterium</i> biovar 1 O^c	MAFF106580	+	+	+	-	+	

Rhizogenic <i>Agrobacterium</i> biovar 1	○	MAFF106587	+	+	+	-	+
Rhizogenic <i>Agrobacterium</i> biovar 1	○	MAFF301724	+	+	+	-	+
Rhizogenic <i>Agrobacterium</i> biovar 1	○	MAFF210265	+	+	+	-	-
Rhizogenic <i>Agrobacterium</i> biovar 1	○	MAFF210268	+	+	+	+	+
Rhizogenic <i>Agrobacterium</i> biovar 1	○	NCPPB2655	+	+	+	-	-
Rhizogenic <i>Agrobacterium</i> biovar 1	○	NCPPB2656	+	+	+	+	+
Rhizogenic <i>Agrobacterium</i> biovar 1	○	NCPPB2659	+	+	+	+	+
Rhizogenic <i>Agrobacterium</i> biovar 1	○	NCPPB2660	+	+	+	-	-
Rhizogenic <i>Agrobacterium</i> biovar 1	○	NCPPB4043	+	+	+	-	-
Rhizogenic <i>Agrobacterium</i> biovar 1	○	NCPPB4042	+	+	+	-	-
Rhizogenic <i>Agrobacterium</i> biovar 1	○	ST15.13/001	+	+	+	+	+
Rhizogenic <i>Agrobacterium</i> biovar 1	○	ST15.13/006	+	+	+	-	-
Rhizogenic <i>Agrobacterium</i> biovar 1	○	ST15.13/007	+	+	+	-	-
Rhizogenic <i>Agrobacterium</i> biovar 1	○	ST15.13/012	+	+	+	+	+
Rhizogenic <i>Agrobacterium</i> biovar 1	○	ST15.13/013	+	+	+	+	-
Rhizogenic <i>Agrobacterium</i> biovar 1	○	ST15.13/039	+	+	+	+	+
Rhizogenic <i>Agrobacterium</i> biovar 1	○	ST15.13/040	+	+	+	+	-
Rhizogenic <i>Agrobacterium</i> biovar 1	○	ST15.13/042	+	+	+	+	-
Rhizogenic <i>Agrobacterium</i> biovar 1	○	ST15.13/046	+	+	+	+	+
Rhizogenic <i>Agrobacterium</i> biovar 1	○	ST15.13/048	+	+	+	+	+
Rhizogenic <i>Agrobacterium</i> biovar 1	○	ST15.13/054	+	+	+	+	+
Rhizogenic <i>Agrobacterium</i> biovar 1	○	ST15.13/056	+	+	+	+	+
Rhizogenic <i>Agrobacterium</i> biovar 1	○	ST15.13/057	+	+	+	+	-
Rhizogenic <i>Agrobacterium</i> biovar 1	○	ST15.13/059	+	+	+	-	-
Rhizogenic <i>Agrobacterium</i> biovar 1	○	ST15.13/060	+	+	+	+	+

	Rhizogenic <i>Agrobacterium</i> biovar 1	○	ST15.13/064	+	+	+	+	+
	Rhizogenic <i>Agrobacterium</i> biovar 1	○	ST15.13/077	+	+	+	+	+
	Rhizogenic <i>Agrobacterium</i> biovar 1	○	ST15.13/090	+	+	+	-	-
	Rhizogenic <i>Agrobacterium</i> biovar 1	○	ST15.13/091	+	+	+	-	-
	Rhizogenic <i>Agrobacterium</i> biovar 1	○	ST15.13/095	+	+	+	-	+
	Rhizogenic <i>Agrobacterium</i> biovar 1	○	ST15.13/097	+	+	+	-	-
	Rhizogenic <i>Agrobacterium</i> biovar 1	○	ST15.13/098	+	+	+	-	-
	Rhizogenic <i>Agrobacterium</i> biovar 1	○	NCPPB4062	+	+	+	+	-
	Rhizogenic <i>Agrobacterium</i> biovar 1	○	ST15.13/045	+	+	+	+	-
	Rhizogenic <i>Agrobacterium</i> biovar 2		NCPPB2991	+	+	+	-	-
	Rhizogenic <i>Agrobacterium</i> biovar 2		LMG150	-	-	-	-	-
	Rhizogenic <i>Agrobacterium</i> biovar 2		NCPPB2303	-	-	-	-	-
	Rhizogenic <i>Agrobacterium</i> biovar 2		LMG149	-	-	-	-	-
	Rhizogenic <i>Agrobacterium</i> biovar 2		LMG138	+	+	-	-	-
	Rhizogenic <i>Agrobacterium</i> biovar 2		ST15.13/027	-	-	-	-	-
Beta-proteobacteria	<i>Burkholderia bryophila</i>		ST15.15/021	-	-	-	-	-
	<i>Burkholderia insulsa</i>		ST15.15/014	-	-	-	-	-
	<i>Collimonas arenae</i>		ST15.15/017	-	-	-	-	-
	<i>Collimonas fungivorans</i>		ST15.15/016	-	-	-	-	-
	<i>Collimonas pratensis</i>		ST15.15/019	-	-	-	-	-
	<i>Janthinobacterium lividum</i>		ST15.15/039	-	-	-	-	-
Gamma-proteobacteria	<i>Escherichia coli</i>		ST08.12/001	-	-	-	-	-
	<i>Pseudomonas aeruginosa</i>		ST01.08/008	-	-	-	-	-
	<i>Pseudomonas fluorescens</i>		ST12.14/123	-	-	-	-	-
	<i>Pseudomonas lurida</i>		EPU_2_30	-	-	-	-	-

<i>Pseudomonas orientalis</i>	ST12.14/122	-	-	-	-	-
<i>Pseudomonas plecoglossicida</i>	ST12.14/336	-	-	-	-	-
<i>Pseudomonas poae</i>	9.1.2-B1	-	-	-	-	-
<i>Pseudomonas putida</i>	ST12.14/260	-	-	-	-	-
<i>Pseudomonas veronii</i>	EHE_1_3	-	-	-	-	-

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617 ^aAntagonistic activity was evaluated using the agar overlay assay (Bosmans *et al.*, 2016b). Antagonistic effects were observed as a clear zone of
618 inhibition where growth of the tested bacterium was inhibited (+). -, no inhibition zone observed.

619 ^bAD, NIOO culture collection, Wageningen, The Netherlands; LMG, Laboratory of Microbiology, Ghent University, Ghent, Belgium; MAFF,
620 NIAS Genebank (National Institute of Agrobiological Sciences), Ibaraki, Japan; NCPPB, National Collection of Plant Pathogenic Bacteria, York,
621 UK; EMI, EPU, EHE and ST, PME&BIM culture collection, Sint-Katelijne Waver, Belgium.

622 ^c*Agrobacterium* biovar 1 strains isolated from Cucurbitaceae (melon, cucumber) and Solanaceae (tomato crops) (for more information, see
623 Bosmans *et al.*, 2015) are indicated by green and red circles, respectively.

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Figure 2.JPEG

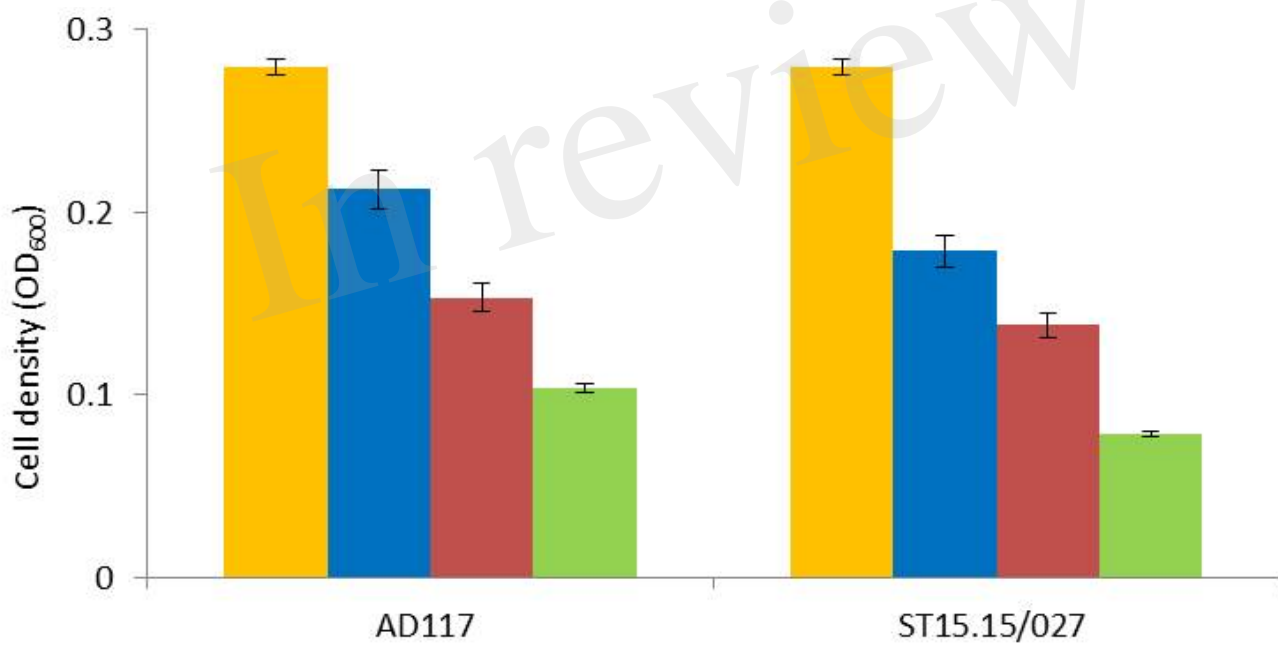


Figure 3.JPEG

