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This is the author's manuscript

Original Citation:

Availability:

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7 was subsequently published in [Zhang D., Spadaro D., Garibaldi A., Gullino M.L. (2010) -
8 *Efficacy of the antagonist Aureobasidium pullulans PL5 against postharvest pathogens of peach,*
9 *apple and plum and its modes of action. Biological Control, 54, 172-180. DOI:*
10 *10.1016/j.biocontrol.2010.05.003*].

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22 **Efficacy of the antagonist *Aureobasidium pullulans* PL5 against postharvest pathogens of**
23 **peach, apple and plum and its modes of action**

24

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32 **Abstract**

33 The efficacy of *Aureobasidium pullulans* PL5 against different postharvest pathogens of fruits
34 (*Monilinia laxa* on plums and peaches, *Botrytis cinerea* and *Penicillium expansum* on apples)
35 were evaluated under storage conditions when applied at 10^8 cells ml^{-1} and their interactions
36 were studied *in vitro* and *in vivo* to discover the possible modes of action. Under 1.2°C and
37 95% relative humidity (RH) for 28 days, the efficacy of PL5 against *M. laxa* on plums was
38 45%, reducing disease incidence from 78% to 43%. Under 1°C and 95% RH for 21 days, the
39 efficacy against *M. laxa* on peaches was 63%, reducing disease incidence from 79% to 29%.
40 Under 4°C and 95% RH for 45 days, the efficacy against *B. cinerea* and *P. expansum* on
41 apples was 56% and 46%, respectively. In Lilly-Barnett minimal salt medium with the fungal
42 cell walls of pathogens as sole carbon source, PL5 produced β -1,3-glucanase, exo-chitinase
43 and endo-chitinase. Nutrient concentrations had significant effect on pathogen growth
44 reduction by PL5. No attachment was observed in antagonist-pathogen interactions *in vitro* or
45 *in vivo*. PL5 completely inhibited pathogen spore germination in PDB at 10^8 cells ml^{-1} ,
46 whereas at 10^6 cells ml^{-1} the efficacy was significantly decreased. However, inactivated cells
47 and culture filtrate of PL5 had no effect on pathogen spore germination and germ tube
48 elongation. Our results showed that *A. pullulans* PL5 could be introduced in commercial
49 formulations to control postharvest pathogens on fruits and its activity was based on secretion
50 of lytic enzymes and competition for nutrients.

51

52 **Keywords:** apple, *Aureobasidium pullulans*, β -1,3-glucanase, biological control, *Botrytis*
53 *cinerea*, chitinase, *Monilinia laxa*, peach, *Penicillium expansum*, plum.

54

55 1. Introduction

56 Fruit and vegetables are highly perishable products, especially during the postharvest
57 phase and major losses are caused by postharvest pathogens (Ippolito and Nigro, 2000; Chan
58 and Tian, 2005; Zhang et al., 2008). *Monilinia laxa*, *Botrytis cinerea* and *Penicillium*
59 *expansum* are among the most important postharvest pathogens on fruit and vegetables
60 (Snowdon, 1990). Among them, three species of *Monilinia* can cause severe losses on stone
61 fruits (Karabulut et al., 2002; Pellegrino et al., 2009), but *M. laxa* is the most dangerous in
62 European countries; *B. cinerea* could cause grey mold on pome and stone fruits, and *P.*
63 *expansum* can cause blue mold decay, which is one of the most destructive disease of pear and
64 apples and it is accompanied by the production of patulin, a mycotoxin with
65 immunosuppressive effects on humans (Spotts and Chen, 1987; Spadaro et al., 2007).
66 Chemical treatment is an important method for controlling postharvest diseases of fruits
67 (Eckert and Ogawa, 1988). However, pathogen resistance to fungicides (Holmes and Eckert,
68 1999) and concern for public safety have resulted in the cancellation of some of the most
69 effective fungicides in Europe (Directive 91/414 CE and Regulation) and the United States
70 (Food Quality Protection Act). In addition, the use of synthetic fungicides to control
71 postharvest diseases of peaches and plums is prohibited in European Union countries.
72 Therefore, researches have been focused on the development of alternative control that should
73 be both effective and economically feasible (El-Ghaouth et al., 1998). Biological control is an
74 effective alternative to fungicidal treatment in controlling postharvest diseases of fruits (Jijakli
75 and Lepoivre, 1998; Spadaro and Gullino, 2004). In 1995, the first commercial products were
76 registered in the United States by the U.S. Environmental Protection Agency (EPA) and are
77 sold under the names BioSave 100 and 110 to control postharvest rots of pome and citrus
78 fruit. In 2007, the biofungicide “Shemer” (based on a strain of *Metschnikowia fructicola*
79 Kurtzman & Droby) was registered in Israel, and is commercially used for the control of
80 sweet potato and carrot storage diseases (Blachinsky et al., 2007). In addition, a commercial

81 formulation of *Candida sake* was recently developed and registered for use on pome fruit in
82 Spain under the name Candifruit® (Droby et al., 2009).

83 The comprehension of the modes of action of an antagonist is an important
84 prerequisite both for enhancing their biocontrol activity and establishing screening criteria in
85 search for new antagonists (Qin et al., 2003). Elucidation of the mechanisms of action is often
86 hampered by the complex interaction among host-pathogen-antagonist (Jijakli and Lepoivre,
87 1998). The mode of action of antagonists generally involves antibiotics (Bull et al., 1998),
88 nutrient competition and site exclusion (Bencheqroun et al., 2007), induced host resistance
89 (El-Ghaouth et al., 1998), and direct interactions between the antagonist and the pathogen
90 (Castoria et al., 1997). Additional modes of action including the production of lytic enzymes
91 viz., β -1,3-glucanase and chitinase were also reported (Ippolito et al., 2000; Saligkarias et al.,
92 2002; Yu et al., 2008).

93 *Aureobasidium pullulans* De Bary (Arnaud) is widely distributed in different
94 environments. Different strains of *A. pullulans* can produce amylase, proteinase, lipase,
95 cellulase, xylanase, mannanase, transferases, pullulan, siderophore, and single-cell protein.
96 Therefore it is a biotechnologically important yeast that can be used in different fields (Chi et
97 al., 2009). Moreover, different strains of *A. pullulans* showed wide efficacy against *B.*
98 *cinerea*, *P. expansum* and *Rhizopus stolonifer* on apple, sweet cherry, grapes, strawberry and
99 peach (Lima et al., 1997; Ippolito et al., 2000; Schena et al., 2003; Bencheqroun et al., 2007).
100 In particular, the strain PL5 of *A. pullulans* showed a high efficacy in the control of
101 postharvest diseases of peaches (Zhang et al., 2010). Strains of *A. pullulans* have been
102 reported to act against fungal pathogens through competition for nutrients (Bencheqroun et
103 al., 2007), secretion of exochitinase and β -1,3-glucanase (Castoria et al., 2001), or induction
104 of defence responses (Ippolito et al., 2000). Understanding the modes of action is essential for
105 developing appropriate commercial formulations and application methods to maximize the
106 potential use of microbial biocontrol agents.

107 The aim of this research was to evaluate the efficacy of *A. pullulans* PL5 against *B. cinerea*, *P.*
108 *expansum*, and *M. laxa* on postharvest fruits under storage conditions. Generally, biocontrol
109 agents are selected and optimized for their efficacy just against one pathogen on one fruit. In
110 our experiment, the antagonist was tested against three pathogens on three fruit species under
111 standard storage conditions. A second aim was to evaluate the production of hydrolytic
112 enzymes by *A. pullulans* PL5 *in vitro*, by studying the β -1,3-glucanase (EC 3.2.1.39),
113 exochitinase or *N*-acetyl- β -glucosaminidase (EC 3.2.1.52) and endochitinase (EC 3.2.1.14)
114 activities. A third aim was to investigate the effects of different nutrient concentrations on the
115 interactions with three postharvest pathogens (*M. laxa*, *B. cinerea* and *P. expansum*) *in vitro*
116 and *in vivo* to reveal their possible modes of action.

117

118 **2. Materials and methods**

119

120 *2.1 Microorganisms and fruit*

121 The yeast-like fungus *Aureobasidium pullulans* De Bary (Arnaud) PL5 was isolated from a
122 plum cv. Angeleno produced in Piemonte (Northern Italy) and maintained on potato dextrose
123 agar plates (PDA; 39 g l⁻¹, Merck) at 4°C for further studies. The antagonist, selected for its
124 efficacy (Zhang et al., 2009), was identified through microscopic observation of cell and
125 colony morphology, and by sequencing the internal transcribed spacer 1 (ITS1), 5.8S
126 ribosomal RNA gene, and internal transcribed spacer 2 (ITS2) according to White et al.
127 (1990). The sequence was deposited in GenBank (FJ919775). The strain was grown at
128 25±1°C in 300 ml YPD (10 g l⁻¹ of yeast extract; 20 g l⁻¹ of triptone-peptone of casein and 20
129 g l⁻¹ of D(+)-glucose) on a rotary shaker (250 rpm) for 48 h. Cells were harvested after
130 centrifugation at 5000×g for 10 min, resuspended in sterile Ringer's solution (pH 6.9±0.1;
131 Merck) and adjusted to the desired cell concentration (10⁸ cells ml⁻¹) with a Bürker chamber.

132 The pathogens *Monilinia laxa* (Aderhold & Ruhland) Honey, *Botrytis cinerea* (de Bary)
133 Whetzel, and *Penicillium expansum* Link were isolated from infected fruit, and pure cultures
134 were maintained on PDA plates at 4°C. The spore suspensions of the pathogens were
135 prepared from 7 days old mycelia by scraping the conidia on PDA plates with a sterile
136 Ringer's solution and adjusting to the desired conidial concentration with a Bürker chamber.
137 Fruit used throughout the experiments were apples (*Malus × domestica* Borkh.) cv. Golden
138 delicious, plums (*Prunus domestica* L.) cv. Angeleno, and peaches [*Prunus persica* (L.)
139 Batsch] cv. Redhaven harvested at commercial maturity and kept at 1°C until use. Fruit were
140 disinfected with 2% sodium hypochlorite for 2 min, washed with tap water and air dried prior
141 to wounding.

142

143 2.2 Efficacy of the antagonist PL5 against *M. laxa* on plums and peaches under storage 144 conditions

145 The cells of the antagonist PL5 was harvested by centrifugation at 5000 ×g for 10 min
146 after being grown in 300 ml YPD medium in 1000-ml Erlenmeyer flasks for 48 hours at 25°C
147 on a rotary shaker at 250 rpm, and were diluted with 30 L tap water in 50 L tank into a final
148 concentration of 1×10^8 cells ml⁻¹. Thirty L of tebuconazole solution (2.5 mL/L of Folicur,
149 Bayer Crop Science; 25.0 % a. i.) was prepared according to the manufacturer. The plums
150 and peaches were surface sterilized with 1% commercial sodium hypochlorite solution for 1
151 min followed by rinsing with tap water. After 2 h air-drying at 25°C, fruits were treated with
152 the antagonist suspension by dipping in tank for 1 min. Fruit surfaces were then air dried at
153 25°C for 2 h and 1 ml of a conidial suspension of *M. laxa* (5×10^4 spores ml⁻¹) was sprayed
154 universally onto each fruit. The fruits treated with tebuconazole constituted the chemical
155 control, while the fruits simply inoculated with the pathogen served as inoculated control.
156 Three replicates of 25 fruits were prepared for each treatment. Two hours after inoculation of
157 the pathogen on fruit surface, the plums were stored at 1.2°C and 95% RH and the peach

158 fruits were stored at 1°C and 95% relative humidity (RH) under storage conditions. After 21
159 and 28 days of storage, incidence of the rotten peaches and plums were measured,
160 respectively. The experiments were repeated twice.

161

162 2.3 Efficacy of the antagonist PL5 against *B. cinerea* and *P. expansum* on apples under 163 storage conditions

164 To evaluate the efficacy of the antagonist PL5 against *B. cinerea* and *P. expansum* on
165 apples, the trials were prepared in a similar way, as described above. Briefly, the cells of
166 antagonist *A. pullulans* PL5 were diluted with 30 L tap water in 50 L tank into a final
167 concentration of 1×10^8 cells ml^{-1} . The apples were surface sterilized with 1% commercial
168 sodium hypochlorite solution for 1 min followed by rinsing with tap water. After 2 h of air-
169 dry at 25°C, fruits were treated with the antagonist suspension (1×10^8 cells ml^{-1}) by dipping in
170 tank for 1 min. Fruit surfaces were air dried at 25°C for 2 h and 1 ml of a conidial suspension
171 of *B. cinerea* (5×10^4 spores ml^{-1}) or *P. expansum* (5×10^4 spores ml^{-1}) were sprayed onto each
172 fruit according to the different trials. The fruits treated with 2.5 mL/L of Folicur (Bayer Crop
173 Science; tebuconazole: 25.0 %) played as chemical controls, while the fruits only inoculated
174 with the pathogen served as inoculated controls. Three replicates of 25 fruits were prepared
175 for each treatment. Two hours after inoculation of the pathogen on fruit surface, the apples
176 were stored at 4°C and 95% RH under storage conditions. After 45 days of storage, the rotten
177 apples were counted and the infected percentage of the apples was recorded. The experiment
178 was repeated twice.

179

180 2.4 Pathogen cell wall preparation

181 Cell wall preparations (CWP) of each pathogen were prepared as described by
182 Saligkarias et al. (2002) with small modifications. Briefly, the pathogens were grown in
183 potato dextrose broth media (Sigma-Aldrich, USA) and the mycelium was collected with

184 four-fold cotton gauzes and washed twice with deionized water followed by two washes
185 through Whatman No.1 filter paper with deionized water and centrifuging (Centrifuger 6K15,
186 Sigma, Germany) at 480×g for 2 min. After discarding the supernatant, the fungal mycelia
187 were sonicated with a probe type sonicator (USC6000, VWR, Malaysia) for 20 min and
188 centrifuged at 480×g for 5 min. The supernatant was discarded and the pellet was resuspended
189 in deionized water. The same procedure was performed for six times. Then the crushed
190 mycelia were resuspended into an equal volume of Tris/HCl buffer (pH 7.2), centrifuged at
191 1900×g for 15 min, and the supernatant was discarded. The pellet was subjected to three
192 successive cycles of centrifugation and resuspension. The final pellet was frozen with liquid
193 N₂, lyophilized and stored at -20°C for further studies.

194

195 *2.5 Preparation of colloidal chitin*

196 Colloidal chitin was prepared according to the method described by Roberts and
197 Selitrennikoff (1988) from shrimp shell chitin. Five grams of chitin powder (Sigma, USA)
198 was added slowly into 100 ml of concentrated HCl and left overnight at 4°C with vigorous
199 stirring. The mixture was added to 2 L of ice-cold 95% ethanol with rapid stirring and kept
200 overnight at 25°C. The precipitate was collected by centrifugation at 3000×g for 20 min at
201 4°C and the precipitate was washed with sterile distilled water until the colloidal chitin
202 became neutral (pH 7.0). Colloidal chitin solution (5 mg ml⁻¹) was prepared and stored at 4°C
203 for further studies.

204

205 *2.6 Culture conditions for enzyme production by the antagonist*

206 The antagonist *A. pullulans* PL5 was cultured in modified Lilly-Barnett minimal salt
207 medium (LBMS; Lilly and Barnett, 1951) containing 2 mg ml⁻¹ CWP of each pathogen as
208 sole carbon source. A 30 ml of culture media in 100-ml flask was incubated at 25°C on a
209 rotary shaker at 150 rpm for 0, 24, 72, and 96 h. Culture filtrates from each individual culture

210 flask were collected by centrifuging at 7000×g for 8 min, and the supernatant was used for
211 enzyme assays.

212

213 *2.7 Determination of β -1,3-glucanase activity produced by A. pullulans PL5*

214 β -1,3-glucanase activity assays were carried out by measuring the amount of reducing
215 sugars released from laminarin (Sigma, USA), using glucose as a standard (Masih and Paul,
216 2002). A reaction mixture was prepared by adding 250 μ l of 0.005M potassium acetate buffer
217 (pH 5.0) containing 2.5 mg of laminarin per ml into 250 μ l of culture filtrate (Chan and Tian,
218 2005). The enzyme-substrate mixture was incubated for 2 h at 40°C in a water bath
219 (Melsungen, Germany). Then 0.5 ml of dinitrosalicylic acid reagent was added and boiled at
220 100°C for 5 min. After cooling, 2 ml of deionized water were added directly and measured
221 spectrophotometrically at 595 nm. Background levels of reducing sugars were determined
222 with a time 0 supernatant substrate just prior to boiling at 100°C for 5 min. The protein
223 concentration of the enzyme solution was determined according to Bradford (1976) by using
224 bovine serum albumin (Sigma, USA) as a standard. The specific activity was expressed as
225 micromoles of glucose per milligram protein per hour (μ mol Glucose Released/mg Protein/h;
226 Fan et al., 2002). Each treatment had three replications and the experiments were repeated
227 twice.

228

229 *2.8 Determination of chitinase activity produced by A. pullulans PL5*

230

231 The exo-chitinase was performed according to Abeles et al. (1970). A reaction mixture
232 was prepared by adding 0.5 ml of 5 mg ml⁻¹ colloidal chitin containing 1.2 μ mol l⁻¹ sodium
233 azide and 56 μ mol l⁻¹ sodium acetate to 0.5 ml enzyme supernatant. For the endo-chitinase
234 assay, a reaction mixture was prepared by adding 0.1 ml of 3% (w/v) desalted snail gut
235 enzyme cytohelicase (Sigma, USA) and 0.1 ml of 1 mol l⁻¹ potassium phosphate buffer (pH

236 7.0) into 0.5 ml of 5 mg ml⁻¹ colloidal chitin in a 2 ml tube, and then 0.5 ml of enzyme
237 supernatant was added to the same tube. The enzyme-substrate mixture was incubated for 2h
238 at 37°C in the water bath with constant shaking. The supernatant was collected from the
239 mixture by centrifuging at 7000×g for 8 min. In the following, 0.5 ml dinitrosalicylic acid
240 reagent was added to the mixture and boiled at 100°C for 5 min. After cooling, 1.5 ml of
241 deionized water was added directly and measured spectrophotometrically at 550 nm.
242 Background levels of reducing sugars and the protein concentration of the enzyme solution
243 was determined as described above. The specific activity was reported as micromoles of N-
244 acetyl-D-glucosamine per milligram protein per hour (μmol GlcNAc Released/mg Protein/h)
245 according to Reissig et al. (1955). Each treatment had three replications and the experiments
246 were repeated twice.

247

248 2.9 Biocontrol activity of antagonist against postharvest pathogens in vivo

249 To assess the biocontrol activity of *A. pullulans* PL5 against *B. cinerea* and *P.*
250 *expansum* on apples and against *M. laxa* on peaches, the fruits were surface sterilized with 1%
251 commercial sodium hypochlorite solution for 1 min and then rinsed with tap water. Three
252 artificial wounds (about 3mm wide× 3mm deep) along the equatorial zone of the each fruit
253 were made. Aliquots of 30 μl of antagonist suspension containing 10⁸cells ml⁻¹ were pipetted
254 into each wound. After 2 hours of incubation at 25±2°C, the wounds were inoculated with
255 30μl of pathogen conidial suspension containing 5×10⁴ conidia ml⁻¹. The fruits inoculated
256 with water served as control. Fifteen fruits with three wounds were used for each treatment.
257 The fruits were stored at 25±2°C and after 7 days, the diameter of decay on fruits was
258 measured. The experiments were repeated twice.

259

260 2.10 Testing of antagonist-pathogen direct interaction in vitro and in vivo

261 The interaction between the antagonist and the pathogen hyphae was assessed in Petri
262 dishes (90 mm, containing 20 ml PDA for *B. cinerea* and *P. expansum* or 20 ml peach juice
263 agar for *M. laxa*). The plates were divided into three sections, 30 mm wide each, along the
264 diameter. Pathogen mycelial plugs (5mm in diameter) were corked from a 4 day old PDA
265 culture plate and fixed upon one point on the agar surface. After 48h at 25±2°C, 50 µl aliquots
266 of the antagonist cell suspension containing 10⁸ cells ml⁻¹ were streaked onto PDA and peach
267 juice agar at 30 mm from the pathogen plug (Spadaro et al., 2002). The dual cultures were
268 washed with deionized water for about 2 min after 48h co-culture at 25±2°C. The interactions
269 of the antagonist and pathogen were directly observed under light microscope (Axioskop 40,
270 Germany). The experiments were repeated twice.

271 To test the interaction between antagonist and pathogen *in vivo* the fruits were
272 inoculated as described above. The fruits were stored at 25°C and the interaction of the
273 antagonist and pathogens were directly observed under light-microscope 7 days after
274 inoculation. Only the interactions between the antagonist and *B. cinerea* are presented in this
275 paper.

276

277 2.11 Effect of nutrient concentration on antagonist-pathogen interaction in vitro

278 To characterize the nutrimental mechanism, different concentrations of PDA (39 g L⁻¹,
279 19.5 g L⁻¹, 9.7 g L⁻¹, and 3.9 g L⁻¹) for *B. cinerea* and *P. expansum* and peach juice agar (50%,
280 25%, 10%, and 5%) for *M. laxa* were used to examine the effect of nutrients on mycelial
281 growth reduction by *A. pullulans* PL5. The co-inoculation of *A. pullulans* PL5 (10⁸ cells ml⁻¹)
282 and the pathogens was performed as described above (2.9). After 7 days of incubation at
283 25°C, the pathogen mycelial growth reduction was measured and recorded. Each treatment
284 had three replications and the experiments were repeated twice.

285

286 2.12 Effect of antagonist on pathogen spore germination in vitro

287 The effect of the antagonist *A. pullulans* PL5 on *M. laxa*, *B. cinerea* and *P. expansum*
288 spore germination was assessed in potato dextrose broth (PDB) according to the method
289 described by Spadaro et al. (2002). Yeast cells grown at $25\pm 2^{\circ}\text{C}$ for 48h in YPD broth were
290 harvested by centrifugation at $5000\times g$ for 10 min, and then resuspended in sterile ringer
291 solution. The culture media of YPD were filtered with $22\ \mu\text{m}$ nitro-cellulose filter for further
292 use. Living cells of the antagonist ($100\ \mu\text{l}$ of a suspension containing 5×10^7 , 5×10^8 and 5×10^9
293 cells ml^{-1}) or cells ($100\ \mu\text{l}$ of a suspension containing 5×10^9 cells ml^{-1}) inactivated by
294 irradiation for 30' with a germicidal lamp (General Electric, G15T8) that emitted
295 predominantly UV light of a wavelength of 254 nm at fluence of $1.5\ \text{W/m}^2$ posed at 5 cm
296 from the cell suspension layer (2 mm thick) were added to 5 ml PDB in 35 ml tubes. The final
297 concentrations of living yeast cells were 1×10^6 , 1×10^7 , and 1×10^8 cells ml^{-1} respectively. One
298 ml of culture filtrate was transferred into the 5 ml PDB tubes for the culture filtrate treatment.
299 Pathogens were then inoculated into each tube containing 5×10^4 conidia ml^{-1} . The PDB tubes
300 inoculated only with the pathogens served as control. Then the tubes were incubated at 25°C
301 on a rotary shaker at 200 rpm for 20 h. One hundred spores were randomly selected from each
302 treatment. The spore germination of each pathogen was assessed and the germ tube length
303 was measured using light microscope. Three replications were maintained for each treatment
304 and the experiments were repeated twice.

305

306 2.13 Statistical analysis

307 Homogeneity of variances was tested by Levene's test ($P = 0.05$). When the variances
308 were considered homogenous, data were subjected to analysis of variance (ANOVA) using
309 SPSS (version 13.0) and statistical significance was assessed at the level of $P < 0.05$. Duncan's
310 multiple Range Test was used to separate the means.

311

312 3. Results

313

314 *3.1 Efficacy of the antagonist PL5 against M. laxa on plums and peaches under storage*
315 *conditions*

316 After storage at 1.2°C and 95% RH for 28 days, brown rot disease incidence was recorded
317 on plums (Table 1). Compared with that (78%) of inoculated control, the disease incidence
318 caused by *M. laxa* of plums treated with *A. pullulans* PL5 was reduced to 43%.
319 Correspondingly, the efficacy of PL5 against *M. laxa* on plums was 45% .

320 After the peach fruits were stored at 1°C and 95% RH for 21 days under storage conditions,
321 the disease incidence was reduced from 79% (inoculated control) to 29% and the efficacy of
322 PL5 against *M. laxa* on peaches was 63%. The biocontrol efficacy of tebuconazole to control
323 brown rot on peaches and plums was 67% and 84%, respectively (Table 1).

324

325 *3.2 Efficacy of the antagonist PL5 against B. cinerea and P. expansum on apples under*
326 *storage conditions*

327 After storage at 4°C and 95% RH under storage conditions for 45 days, the incidence of rotten
328 apples caused by *B. cinerea* and *P. expansum* was recorded. The results are shown in Table 2.
329 *A. pullulans* PL5 reduced gray mold incidence from 45% to 20% and reduced blue mold
330 incidence from 47% to 25%. Correspondingly, the efficacy of *A. pullulans* PL5 against *B.*
331 *cinerea* and *P. expansum* on apples was 56% and 46%, respectively.

332

333 *3.3 Production of β -1, 3-glucanase by A. pullulans PL5 and its activity*

334 The antagonist *A. pullulans* PL5 produced extracellular β -1,3-glucanase in culture
335 medium in presence of pathogen cell walls as sole carbon source. *A. pullulans* PL5 began to
336 produce extracellular β -1,3-glucanase immediately after cultivation in LBMS medium with *B.*
337 *cinerea* and *P. expansum* cell walls as sole carbon source. When *M. laxa* was supplied as sole
338 carbon source, at 0 h incubation, no extracellular β -1,3-glucanase was detected. At 48 h

339 incubation, the extracellular β -1,3-glucanase activity reached the maximum level, and the β -
340 1,3-glucanase activity in the minimal salt medium with *M. laxa*, *B. cinerea* and *P. expansum*
341 cell walls as sole carbon source were 46.9, 66.1 and 80.1 U (μmol Glucose Released/mg
342 Protein/h), respectively. However, at 72 h the β -1,3-glucanase activities began decreasing. At
343 96 h incubation, the β -1,3-glucanase activity in the minimal salt medium with *M. laxa*, *B.*
344 *cinerea* and *P. expansum* cell walls as sole carbon source were 32.9, 8.0, and 19.6 U (μmol
345 Glucose Released/mg Protein/h), respectively (Fig. 1 a).

346

347 3.4 Production of chitinases by *A. pullulans* PL5 and their activities

348 The assay of chitinase activity showed that exo-chitinase activity of *A. pullulans* PL5,
349 when cultured in minimal salt medium with the different fungal cell walls as sole carbon
350 source, showed the different trends, according to different incubation periods and different
351 pathogens. The maximum level of exo-chitinase activity in the medium with *M. laxa* and *P.*
352 *expansum* as sole carbon source was detected after 48 h incubation. However, the maximum
353 level of exo-chitinase activity in the medium with *B. cinerea* cell wall as sole carbon source
354 appeared before 48 h incubation (Fig. 1 b). The maximum level of exo-chitinase activity in
355 the medium with *P. expansum* as sole carbon source was 0.96 U (μmol GlcNAc Released/mg
356 Protein/h) which was higher than the maximum levels of those in the medium with *M. laxa*
357 and *B. cinerea* cell walls as sole carbon source (Fig. 1 b).

358 The endo-chitinase activity of *A. pullulans* PL5 cultured in LBMS medium with *P.*
359 *expansum* and *M. laxa* cell walls as sole carbon sources reached the maximum level after 48 h
360 incubation. However, when *A. pullulans* PL5 was cultured in the medium with *B. cinerea* cell
361 walls as sole carbon source, the maximum level of endo-chitinase activity was detected at
362 about 30 h incubation. Also the endo-chitinase activity assay showed that the maximum
363 activity with *P. expansum* cell walls as sole carbon source was higher than the activity with *B.*
364 *cinerea* or *M. laxa* cell walls (Fig. 1 c.).

365

366 3.5 Biocontrol activity of *A. pullulans* PL5 and antagonist-pathogen direct interaction *in vitro*
367 and *in vivo*

368 After 7 days of co-incubation in PDA plates at $25\pm 2^{\circ}\text{C}$, direct interaction *in vitro* was
369 observed microscopically. *A. pullulans* PL5 significantly inhibited the hyphae elongation of
370 the pathogens. However, no attachment of fungal hyphae was observed (Fig. 2) and the
371 hyphae end close to the antagonist grew normally (Fig. 2b).

372 After 7 days of co-incubation in artificial wounds of apples, the pathogen spore
373 germination was almost completely inhibited by *A. pullulans* PL5 (Fig. 2d) and the population
374 of *A. pullulans* PL5 greatly increased. However, no attachment to the pathogen spore surfaces
375 was observed.

376 In artificial wound inoculation tests, *A. pullulans* PL5 was effective in controlling
377 decays of fruits caused by *M. laxa*, *B. cinerea* and *P. expansum* (Table 3). After 7 days of
378 storage at $25 \pm 2^{\circ}\text{C}$, *A. pullulans* PL5 significantly reduced the decay of apples caused by *B.*
379 *cinerea* and *P. expansum* from 61.9 mm and 35.3 mm to 9.5 mm and 7.1 mm, respectively.
380 While the diameter of rotten lesions caused by *M. laxa* on peaches was significantly reduced
381 from 49.4 mm to 24.6 mm.

382

383 3.6 Effect of nutrient concentration on antagonist-pathogen interaction *in vitro*

384 The effect of nutrient concentrations on the interaction of *A. pullulans* PL5 with the
385 three pathogens (*M. laxa*, *B. cinerea* and *P. expansum*) was studied *in vitro*. After incubation
386 at $25\pm 2^{\circ}\text{C}$ for 7 days *A. pullulans* PL5 co-cultured with *B. cinerea* on different concentrations
387 of PDA (39 g L^{-1} , 19.5 g L^{-1} , 9.7 g L^{-1} , and 3.9 g L^{-1}) reduced the mycelial growth by 8.0 mm,
388 10.7 mm, 12.0 mm and 14.8 mm, respectively (Table 4). When co-cultured with *M. laxa* on
389 50%, 25%, 10%, and 5% peach juice agar, *A. pullulans* PL5 reduced the mycelial growth of
390 *M. laxa* by 4.0 mm, 8.2 mm, 11.5 mm and 14.2 mm, respectively. After 7 days of co-

391 incubation at $25 \pm 2^\circ\text{C}$ on 9.7 g L^{-1} and 3.9 g L^{-1} PDA, the pathogen *P. expansum* mycelial
392 growth reduction by the antagonist was 3.3 mm and 7.2 mm. However, on 39 g L^{-1} and 19.5 g
393 L^{-1} PDA, no mycelial growth reduction of *P. expansum* was observed (Table 4).

394

395 3.7 Effect of antagonist on pathogen spore germination in vitro

396 By co-culturing in PDB, the effect of different concentrations of cell suspension,
397 inactivated cells and culture filtrate of *A. pullulans* PL5 were investigated on *M. laxa*, *B.*
398 *cinerea* and *P. expansum* spore germination and germ tube length (Table 5). After 20 h
399 incubation at $25 \pm 2^\circ\text{C}$ almost complete inhibition of the spore germination was observed in
400 presence of 1×10^8 cells ml^{-1} of the antagonist. The percentage of spore germination of *M.*
401 *laxa*, *B. cinerea*, and *P. expansum* were 1.7%, 1.7% and 2.7%, respectively. Correspondingly,
402 at 1×10^8 cells ml^{-1} , the antagonist inhibition efficacies were 98.1%, 98.2% and 96.9%,
403 respectively. When co-cultured with 1×10^7 cells ml^{-1} of the antagonist, the percentage of
404 spore germination of *M. laxa*, *B. cinerea* and *P. expansum* were 31.7%, 28.7% and 26.7%,
405 respectively. Whereas at 1×10^6 cells ml^{-1} of the antagonist applied, the percentage of spore
406 germination were reduced only into 60.0%, 61.7% and 62.7%, respectively. No significant
407 difference of the percentage of spore germination was observed when the pathogens were co-
408 cultured with inactivated cells or culture filtrate.

409 Germ tube elongation of *M. laxa*, *B. cinerea* and *P. expansum* in PDB was greatly
410 controlled by the living cells of *A. pullulans* PL5. In the inoculated control, length of the germ
411 tubes of *M. laxa* and *B. cinerea* were 123.1 μm and 148.4 μm . However, the length of germ
412 tubes of *M. laxa* co-cultured with 1×10^8 , 1×10^7 , and 1×10^6 cells ml^{-1} of the antagonist were
413 4.2 μm , 37.7 μm and 58.1 μm , respectively, and the length of the germ tubes of *B. cinerea* co-
414 cultured with 1×10^8 , 1×10^7 , and 1×10^6 cells ml^{-1} of the antagonist was reduced by 85.9%,
415 70.8% and 38.1%, respectively. The length of germ tubes of *P. expansum* in the presence of
416 1×10^8 , 1×10^7 and 1×10^6 cells ml^{-1} of the antagonist was reduced from 117.5 μm to 13.4 μm ,

417 20.1 μm and 65.3 μm , respectively. When the pathogens were co-cultured with inactivated
418 cells or culture filtrate of the antagonist, no significant difference in germ tube length was
419 observed (Table 5).

420

421 **4. Discussion**

422 The efficacy of the antagonistic yeast-like fungus *A. pullulans* PL5 against three
423 postharvest pathogens on three fruit species (*M. laxa* on plums and peaches, *B. cinerea* and *P.*
424 *expansum* on apples) was evaluated under storage conditions.

425 Though much work has been done to develop biocontrol agents against postharvest
426 pathogens of fruits, none of the commercialized biofungicides is effective against brown rot
427 on stone fruit (Zhang et al., 2010). The antagonist *A. pullulans* PL5 was effective in
428 controlling brown rot on peaches as well as on plums under storage conditions. Few
429 antagonists have been selected for their biocontrol potential on different fruit species. A
430 broader range of efficacy gives more opportunities for the commercial development of
431 antagonists. Our research demonstrated that *A. pullulans* PL5 had also a high efficacy in
432 controlling grey mould and blue mould on apples, beyond its strong biocontrol activity
433 against brown rot on peaches and plums.

434 The lack of secretion of antibiotic compounds in the culture filtrate is an important and
435 positive factor for potential registration of the biocontrol agents. In this research, the trials,
436 carried out *in vitro* showed that neither inactivated cells nor culture filtrate of the antagonist
437 had effect on the pathogens spore germination or germ tube elongation, suggesting that
438 production of antibiotics were not involved in the modes of action of *A. pullulans* PL5 against
439 the pathogens.

440 Mycoparasitism of antagonists is associated with the production of cell wall-degrading
441 enzymes and induction of host defense. Concurrent induction of chitinase and glucosidase has
442 been described in plant as a response to infection by microbial pathogen (Lorito et al., 1994),

443 and the two classes of enzymes exhibit synergistic activity against growth of several fungi
444 (Qin et al., 2003). Peroxidase in combination with chitinase and β -1,3-glucanase has also
445 shown to inhibit the growth of several pathogenic fungi *in vitro* (Schlumbaum et al., 1986).
446 Lorito et al. (1994) anyway proved that a synergistic antifungal effect of either exo- or endo-
447 chitinases in combination with glucosidase of *T. harzianum* was able to inhibit *B. cinerea*.
448 The strain PL5 showed a strong antagonistic activity in controlling postharvest pathogens,
449 such as *M. laxa* on peaches, *B. cinerea* and *P. expansum* on apples. *A. pullulans* PL5, when
450 applied as a wound treatment, was effective in controlling postharvest decay of apple and
451 peach. Moreover, when co-cultured with the pathogens *in vitro* or *in vivo*, it was also effective
452 in inhibiting the pathogen mycelial growth.

453 It has been suggested that extensive production of the extracellular lytic enzymes,
454 especially β -1,3-glucanase and chitinase, by yeast or filamentous fungi may play an important
455 role either by enhancing nutrient competition or by some other unknown mechanisms (Jijakli
456 and Lepoivre, 1998; Fan et al., 2002; Masih and Paul, 2002; Chan and Tian, 2005). In our
457 investigation, the *in vitro* assays showed that the antagonist *A. pullulans* PL5 produced high
458 β -1,3-glucanase, exo-chitinase and endo-chitinase, especially when co-cultured in the LBMS
459 medium supplied with *P. expansum* cell wall as sole carbon source. Our results *in vitro* are in
460 accordance with previous researches of Castoria et al. (1997 and 2001) and Ippolito et al.
461 (2000) that demonstrated the capacity of some strains of *A. pullulans* of producing β -1,3-
462 glucanase and chitinase. However, the strain *A. pullulans* PL5 used in this research produced
463 higher β -1,3-glucanase than the strain *A. pullulans* L-30 tested by Castoria et al. (2001), when
464 the two strains were cultured in a medium with the same pathogen cell walls as sole carbon
465 source. Anyway, the biocontrol efficacy cannot be compared among the two strains because
466 Ippolito et al. (2000) worked on artificially wounded and inoculated apples, while we tested
467 PL5 on intact fruits under standard storage conditions. Moreover, in this paper, the dynamics
468 of time-course changes of enzyme activities produced by the antagonist was studied. The

469 production of β -1,3-glucanase and chitinase by *A. pullulans* PL5 and their activities
470 demonstrated a time-curve trend of the enzyme activities. *A. pullulans* PL5 began to produce
471 extracellular β -1,3-glucanase immediately after inducing the treatment in LBMS medium with
472 *B. cinerea* and *P. expansum* cell walls used as sole carbon source, while when *M. laxa* was
473 supplied as sole carbon source, no extracellular β -1,3-glucanase was detected at 0 h. At 24 h,
474 the activity of β -1,3-glucanase began increasing and reached the maximum level at 48 h.
475 However, at 72 h the β -1,3-glucanase activity began decreasing. Even if the exo- and endo-
476 chitinase activities of *A. pullulans* PL5 showed a similar trend to that of β -1,3-glucanase
477 activities, the maximum activities in LBMS medium with the three fungal cell walls as sole
478 carbon source were different. Probably *A. pullulans* PL5 showed a different response to the
479 presence of the cell walls of the three pathogens in producing chitinase. At present, the
480 production pathways of β -1,3-glucanase and chitinase by *A. pullulans* PL5 are still poorly
481 understood. Therefore, it is necessary to carry out more molecular research about the
482 regulation and function of the genes related to the enzyme production.

483 Attachment to fungal hyphae or mycoparasitism has also been proposed as a mode of
484 action of biocontrol agents against pathogens of fruits and vegetables (Wisniewski et al.,
485 1991; Arras, 1996; Wan and Tian, 2002). However, in our research, the observation of the
486 antagonist-pathogen interactions *in vitro* and *in vivo* indicated the lack of attachment to the
487 pathogen hyphae. Interaction test *in vitro* showed that the mycelia of the three pathogens were
488 greatly reduced by *A. pullulans* PL5, while no attachment of the antagonist to the pathogens
489 was observed on PDA, proving that inhibition of pathogen by *A. pullulans* PL5 was not based
490 on the mycoparasitism. Attachment of fungal hyphae by *A. pullulans* PL5 was not even
491 detected in the interaction tests performed *in vivo*.

492 Our results indicated that competition for nutrients played a significant effect on the
493 biocontrol activities of *A. pullulans* PL5 against the three pathogens, in accordance to

494 previous studies of Lima et al. (1997). When co-cultured with the pathogens on culture
495 medium with lower nutrient concentrations, the ability of *A. pullulans* PL5 to inhibit the
496 mycelial growth of pathogens was greatly improved. More consistent evidence was supplied
497 by the co-culture of *A. pullulans* PL5 with the pathogen *P. expansum* on different
498 concentrations of PDA. Also the co-culture experiment, where different concentrations of *A.*
499 *pullulans* PL5 had effects on the spore germination and germ tube elongation, supported a
500 major role of the competition for nutrients in the activity of *A. pullulans* PL5 against
501 postharvest pathogens. When the concentrations of *A. pullulans* PL5 were lowered, the spore
502 germination incidence significantly increased and the germ tubes were longer. Competition
503 for nutrients and space are considered among the main modes of action of yeast biocontrol
504 agents (Spadaro et al., 2002). In particular, competition for amino acids, sugars and Fe^{3+} plays
505 an important role in the mechanism of competition for nutrients of some antagonists. Sipiczki
506 (2006) and Saravanakumar et al. (2008) demonstrated that several strains of *Metschnikowia*
507 *pulcherrima* control *B. cinerea* and *P. expansum* in apples through competing for Fe^{3+} with
508 the pathogens. Bencheqroun et al. (2007) proposed that apple blue mould biocontrol by *A.*
509 *pullulans* was based on competition for amino acids. The main mode of action of *Pichia*
510 *guilliermondii* strain R13 in controlling anthracnose on sweet peppers after harvest was
511 competition for nitrogen sources and sugars (Chanchaichaovivat et al., 2008). However,
512 which nutrient sources are exactly involved in the competition of *A. pullulans* PL5 with the
513 pathogens is still under investigation. Summarizing the results above, production of β -1,3-
514 glucanase and chitinase together with out-competition for nutrients and space as well as other
515 undetermined mechanisms constitute important modes of action of *A. pullulans* PL5 against
516 postharvest pathogens of fruits.

517

518 **Acknowledgements**

519 This research was funded by the projects “CIPE – Production of stone fruit in Piedmont:

520 monitoring, prevention and control of pathogenic and mycotoxigenic fungi to guarantee food
521 safety” and “DRUMP – Drupacee minori in Piemonte: problemi fitopatologici e difesa post-
522 raccolta” granted by the Piedmont Region.

523

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642

643 **Figure captions**

644

645 **Figure 1.**

646 a) β -1,3-glucanase activity (EC 3.2.1.39) of *A. pullulans* PL5 grown in LBMS medium
647 supplemented with 2 mg ml⁻¹ CWP (cell wall preparation of each pathogen: *M. laxa*, *B.*
648 *cinerea*, or *P. expansum*) as sole carbon source for 96 h at 25°C. Bars represented standard
649 deviations of the means; b) Exo-chitinase activity (EC 3.2.1.52) of *A. pullulans* PL5 grown in
650 LBMS medium supplemented with 2 mg ml⁻¹ CWP (cell wall preparation of each pathogen:
651 *M. laxa*, *B. cinerea*, or *P. expansum*) as sole carbon source for 96 h at 25°C. Bars represented
652 standard deviations of the means; c) Endo-chitinase activity (EC 3.2.1.14) of *A. pullulans* PL5
653 grown in LBMS medium supplemented with 2 mg ml⁻¹ CWP (cell wall preparation of each
654 pathogen: *M. laxa*, *B. cinerea*, or *P. expansum*) as sole carbon source for 96 h at 25°C. Bars
655 represented standard deviations of the means.

656

657 **Figure 2.**

658 a) Interaction of (A) and *B. cinerea* (B) *in vitro* (magnification 150 × ; bar = 40 μm); b) *B.*
659 *cinerea* hyphae (B) close to *A. pullulans* PL5 *in vitro* (magnification 600 × ; bar = 10 μm); c)
660 apple pulp tissue (*Malus × domestica*; M) as a control (magnification 150 × ; bar = 40 μm); d)
661 Interaction of *A. pullulans* PL5 (A), apple pulp (M) tissue and *B. cinerea* (B) *in vivo*
662 (magnification 150 × ; bar = 40 μm).