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1	How siderophore production can influence the biocontrol activity of Aureobasidium pullulans
2	against <i>Monilinia laxa</i> <mark>on peaches</mark>
3	

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5

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8

9 Abstract

10 In the present study Aureobasidium pullulans strains L1 and L8 were shown both in vitro and in vivo to compete for iron with Monilinia laxa through the secretion of siderophores (1.2 and 1.4 mg 11 ml-1, respectively) and to prevent postharvest fruit decay of peaches. The two strains reduced 12 mycelial growth and conidial germination of M. laxa specially in presence of lower iron 13 concentrations (5 and 10 µg l⁻¹ FeCl₃), confirming a better efficacy when nutrients are scarce. In in 14 vivo assay, A. pullulans L1 and L8 strains inhibited pathogen virulence, reducing by 83.5% and 15 84.4% on average respectively the peach lesion diameter for each tested iron solutions (5, 10, 20 μ g 16 1⁻¹ FeCl₃,). The highest iron solution slowed down the antagonists' action and conversely increase 17 the pathogen aggressiveness. Results shows that A. pullulans L1 and L8 strains compete with M. 18 *laxa* for iron, so revealing new biocontrol aspects. Both strains showed the capability to decrease 19 the accumulation of iron competing with some fungal pathogens and reducing their virulence. These 20 results provide new perspectives for the use of biocontrol agents in agriculture. 21 22

- 23 Keywords: Yeast Iron Stone fruit Monilinia laxa Siderophore
- 24
- 25 **1. Introduction**

Monilinia laxa is the most common species in European stone fruit orchards, and able to cause
postharvest losses that reach high values (59%) (Larena et al., 2005). Monilinia control depends on
an integrated strategy based on fungicide use and cultural practices. However, to overcome the
issues related to the use of fungicides, alternative pathogen control strategies have been
investigated, such as the use of biocontrol agents (BCAs) (Di Francesco et al., 2016). *Aureobasidium pullulans* L1 and L8 strains showed a high capability to control *Monilinia* spp. on
stone fruit as previously reported by Mari et al. (2012).

Among the various modes of action involved in the control of postharvest diseases by antagonistic yeasts, competition for nutrients and space is considered one of the most important, particularly to control wound pathogens for which the availability of exogenous nutrients is crucial during the early stage of infection (Elad and Chet 1987, Mekbib et al., 2011, Bautista-Rosales et al., 2013, Di Francesco et al., 2016, 2017, 2018a).

In fact, previous studies reported how the antifungal ability of *A. pullulans* (Janisiewicz et al., 2000; Bencheqroun et al., 2006), *Pichia caribbica* (Xu et al., 2013), and *Pichia guilliermondii* (Chanchaichaovivat et al., 2008) against *Penicillium expansum*, *Rhyzopus stolonifer*, and *Colletotrichum capsici* respectively, can be influenced by the presence or the absence of exogenous nutrients (Druvefors et al., 2005, Bencheqroun et al., 2006, Liu et al., 2010).

Besides the role of competing for nutrients, iron may also play a role in the Biocontrol Agents
(BCAs) modes of action (Raaska and Mattila-Sandholm 1995) through the production of chelating
molecules (siderophores).

These are molecules able to chelate low weight ferric compounds enhancing the effectiveness of BCAs by subtracting iron to pathogen inhibiting its growth and metabolic activity (Riquelme, 1996). Microorganisms can produce a variety of chelating agents, which solubilize ferric iron and transport it into the cell (Liu et al., 2013, Calvente et al., 2001a, Sanz Ferramola et al., 2013). Under competitive conditions when nutrient availability is a limiting factor, the production of siderophores can represent a distinctive feature for a microorganism. It is known that yeasts produce only a type

52	of siderophore: hydroxamate (Riquelme, 1996). The siderophores production could play an
53	important role in the biocontrol of pathogens because by iron sequestering their growth and
54	metabolic activity were inhibited or slowed down (Riquelme 1996). Calvente et al. (2001a)
55	demonstrated that rhodotorulic acid, a siderophore produced by Rhodotorula glutinis improved the
56	biological control of blue rot of apples caused by P. expansum. Also, Bacillus subtilis produced
57	siderophores playing an important role in the control of <i>Fusarium oxysporum</i> (Yu et al., 2011).

Other studies reported the siderophores ability to inhibit the growth of certain pathogenic fungi, like *Pythium ultimum* and *Sclerotinia sclerotiorum* (Hamdan et al., 1991, McLoughlin et al., 1992), establishing that these molecules could be considered as a potential mechanism of action against several fungal pathogens. Parafati et al. (2015) hypothesized that competition for iron was as one of the main mechanisms of action for *Metschnikowia pulcherrima* against *Botrytis cinerea*.

In a previous study, *A. pullulans* L1 and L8 strains yet showed competition for nutrients like amino
acids and space (Di Francesco et al., 2017, 2018a) against *Monilinia laxa* and *B. cinerea*.

In the present work, we investigated for the first time the involvement of iron in the biocontrol activity of both strains against the brown rot agent of peaches. Specifically, the aims of this study were to determine the ability of L1 and L8 to produce siderophores through i) the chrome azurol S (CAS) assay, ii) the spectrophotometer quantification, and iii) the *in vitro* and *in vivo* competition for iron with *M. laxa* by testing three different concentrations of FeCl₃ (5, 10 and 20 μ g ml⁻¹).

70

71 2. Materials and methods

- 72
- 73 2.1 Fruit

Peaches (*Prunus persica* (L.) Batsch) cv 'Redhaven' were harvested at commercial maturity in orchards located in Cadriano (Bologna, Italy). Fruit-were stored at 0 °C and used within 5 days from harvest and wounded by a sterile nail (3x3x3 mm) on opposite sides of the equatorial area.

78 2.2 Pathogen and antagonists

Monilinia laxa (ML4 strain from CRIOF-DipSA collection) (Di Francesco et al., 2017) was chosen 79 for the present study for its aggressiveness and grown as by Martini et al. (2016). Conidia from 80 pathogen colonies 10 days-old, grown on tomato agar (20 g of Agar Technical, Oxoid Basingstoke, 81 Hampshire, UK; 750 ml distilled water to which 250 ml tomato sauce was added after sterilization) 82 at 25 °C, were collected and suspended in sterile distilled water containing 0.05 %-(v/v) Tween 80. 83 The concentration of each conidia suspension was quantified with haematocytometer and adjusted 84 to a concentration of 10⁵ conidia ml⁻¹ with sterile distilled water. Aureobasidium pullulans L1 and 85 L8 strains (Di Francesco et al., 2018b), were maintained on nutrient yeast dextrose agar (NYDA: 8 86 g of nutrient broth, 5 g of yeast extract, 10 g of dextrose and 15 g of agar in 1 L of distilled water) at 87 4 °C until use. Two days before trials, each antagonist was inoculated on NYDA and incubated at 88 25 °C for 2 days. The yeast cells were collected in sterile distilled water and adjusted with 89 haematocytometer to a concentration of 10⁸ cells ml⁻¹. 90

91

92 2.3 CAS assay

CAS-blue agar was prepared according to Schwyn and Neilands (1987). Dishes were prepared with 93 30 ml of culturing medium for each antagonist and fungus: NYDA for L1 and L8 and PDA for M. 94 laxa. Then the solidified growth media was cut in two halves, one of which was replaced by CAS-95 blue agar. The dishes were inoculated placing in the centre of the borderline, between the two 96 media, a 6-mm-plug of L1, L8 or *M. laxa* derived from a colony in active growth. The plates were 97 incubated in the dark at 25 °C until the agar colour-change. The CAS reaction was determined by 98 measuring the advance of the colour-change in the CAS-blue agar from blue to different shades of 99 red. The control plates were not inoculated but incubated under the same conditions described 100 above. The sample unit was represented by 10 dishes and the experiment was repeated twice. 101

- 102
- 103 2.4 Spectrophotometer siderophore assay

For the siderophore production by yeasts and *M. laxa*, one loop of L1 and L8 cells and one of *M*. 104 laxa conidia were transferred in a siderophore solution as described in Wang et al. (2009) and 105 incubated at 25 °C at 200 rpm for 24 h. In addition, L1 or L8 cells and M. laxa were co-cultured in 106 the siderophore culture solution and incubated in the same conditions above mentioned. Aliquots of 107 5 ml of each sample's suspension were inoculated into 45 ml of new siderophore solution and 108 suddenly incubated for 120 h at 25 °C at 200 rpm. The supernatant of the centrifuged (5.000 rpm 109 for 20 min) cultures (500 µl) was added to 2.5 ml of a solution containing 5 mM FeCl₃ and used for 110 quantitative determination of siderophore by a spectrophotometer at 440 nm (Infinite® 200 PRO-111 Tecan) after 30 min of incubation at 25 °C. A standard curve (0, 0.05, 0.1, 0.25, 0.5, 0.75 and 1.0 112 mg ml⁻¹) was prepared with deferoxamine mesylate (Sigma-Aldrich) (one type of hydroxamates) as 113 chelating agent standard (Calvente et al., 2001). The siderophore amount was extrapolated from the 114 deferoxamine mesylate standard curve. The sample unit was represented by three flasks (replicates) 115 for each condition. The experiment was repeated twice. 116

117

118 2.5 *In vitro* competition for iron

The competition for nutrient exerted by L1 and L8 A. pullulans strains on M. laxa mycelial growth 119 was investigated through the dual interaction between the pathogen and the yeast strains. The 120 121 experiment conducted with the two strains and the pathogen was assessed on Minimal Medium Agar (MMA: Na₂HPO₄ 6 g, KH₂PO₄ 3 g, NaCl 0.5 g, NH₄Cl 1 g, MgSO₄ 1M, CaCl₂ 0.1M, Sucrose 122 30 g, Agar technical 20 g) amended with 0, 5, 10 and 20 µg ml⁻¹ of FeCl₃ (Saravanakumar et al., 123 2008). Monilinia laxa mycelial plugs (6 mm of diameter) from 7-day-old colony were inoculated at 124 30 mm of distance from the plate edge. Aureobasidium pullulans L1 and L8 cells were taken from 125 48 h culture on NYDA by sterile loop and were streaked at 30 mm from the edge of the other side 126 of the plate. Plates were incubated at 25 °C for 5 days and the radial growth of pathogen mycelium 127 was measured. Plates inoculated only with *M. laxa* plug represented the control. The sample unit 128 was represented by five plates (replicates) for each condition. The experiment was performed twice. 129

130

2.6 Influence of iron concentrations on antagonists' biocontrol activity against *Monilinia laxa*germination and germ tube elongation

The competition activity for iron of L1 and L8 was assayed on the germination and germ tube 133 elongation of *M. laxa* conidia. Tissue culture plates (Costar, Corning Inc., Corning, NY) and culture 134 plate inserts Millicell-CM (Millipore Corp., Bedford, MA) were used as reported by Janisiewicz et 135 al., (2000) with some modifications (Di Francesco et al., 2017). Aliquots (120 µl) of minimal 136 medium broth (without agar) amended with 0, 5, 10 and 20 µg ml⁻¹ of FeCl₃ were dispensed in the 137 wells of culture plates with 40 μ l of the pathogen conidia suspension (10⁵ conidia ml⁻¹), while the 138 same aliquots of L1 or L8 cell suspensions (10⁸ cells ml⁻¹) were dispensed inside the cylinder 139 inserts, without physical contact between antagonist and pathogen. The plates were placed at 25 °C 140 on a rotary shaker at 50 rpm; after 6 h of incubation, cylinders were removed from the wells and 20 141 µl of the medium were transferred to a glass slide for microscope (Nikon Eclipse TE2000–E) 142 observations. The percentage of conidia germination and the germ tube elongation (µm) were 143 determined (90 conidia per treatment, 3 microscopic fields with 30 conidia each). Cells without iron 144 solutions addition were considered as a control. The experiment was performed twice. 145

146

147 2.7 In vivo competition for iron

Peach fruits (20) cv Redhaven' were wounded and inoculated with 20 µl of each yeast suspension 148 10⁸ cell ml⁻¹; after 1 h at room temperature fruit were inoculated with 20 µl of the pathogen conidia 149 suspension (10⁵ cells ml⁻¹) and right after its drying, 20 µl of iron solutions, at concentrations above 150 151 mentioned, were added. Fruit treated with sterile distilled water instead of yeast cell suspensions represented the control. Fruit were stored at 20 °C and 90% of relative humidity (RH) for 7 days 152 and the lesion diameters determined by the pathogen were recorded. The sample unit was 153 represented by 45 peaches (15 for replicate) for each antagonist, iron concentration, and control. 154 The experiment was repeated twice. 155

157 **2.8 Statistical analysis**

158	Data were statistically handled by one-way analysis of variance (ANOVA). Statistical comparison
159	of means was carried out to reveal the differences between treatments using Tukey's HSD Test
160	$(\alpha = 0.05)$. Data were reported as mean values \pm standard error (SE) of two experiments. Before
161	analysis of data, homogeneity of variance was tested by the Kruskal–Wallis test. All analyses were
162	performed with Statgraphics software (version centurion 15.0).
163	
164	3. Results
165	
166	3.1 Aureobasidium pullulans L1, L8 strains siderophore production

In order to verify the siderophore production by the antagonists and *M. laxa*, CAS-blue agar was used as screening assay for the target microorganisms. Our results showed that both L1 and L8 induced a colour change of CAS-blue agar from blue to dark orange indicating that both *A*.

pullulans strains are siderophore producers. However, despite the two strains required the same time (twelve days) to change the CAS-blue-agar from blue to dark orange, the colour change halo produced by both yeasts was different in size, being 20 mm for L1, and 27 mm for L8 (Table 1) (Fig. 1). In fact, L1 strain after 12 days of incubation did not determined a total colour change of CAS-blue agar half, as instead L8. Conversely, *M. laxa* produced a barrage between the two media without any colour change of the indicative agar. Control plates, without plug inoculation, not

changed colour after incubation conditions.

177

178 3.2 Spectrophotometer quantification of siderophores

To quantify the siderophore production by *A. pullulans* L1 and L8 strains and the influence of *M. laxa* on this activity, a spectrophotometer assay with deferoxamine mesylate standard curve was conducted. Both strains showed a considerable activity, producing 1.2 and 1.4 mg ml⁻¹ of

- siderophore by L1 and L8 respectively, while *M. laxa* produced only 0.64 mg ml⁻¹ of siderophore, a
- 183 quantity not detectable by CAS-blue-agar.
- 184 Siderophore quantity was not influenced by *M. laxa* showing the same amount with respect to the 185 yeasts grown alone (Fig. 2).
- 186
- 187 3.3 *In vitro* competition for iron
- Competition for iron was tested co-culturing in MM plates amended with $FeCl_3$ (0, 5, 10, or 20 µg ml⁻¹) L1 and L8 strains with *M. laxa* isolate. As shown by Table 2, both *A. pullulans* L1 and L8 strains reduced *M. laxa* mycelium diameter by 41.3% and 43.3%, respectively when iron
- 191 concentration was not present in MM.
- 192 L1 and L8 competition was mainly stimulated by the presence of 10 μ g ml⁻¹ and 20 μ g ml⁻¹ of FeCl₃,
- both leading to an inhibition of fungal pathogen mycelial growth corresponding to 45% and 47.6%
- and 43% and 45%, respectively.
- 195 *Aureobasidium pullulans* strain L8 was slightly more effective with respect to L1 in competing for
- ¹⁹⁶ iron, determining a higher inhibition of *M. laxa* mycelial growth on average by 44.7% and 42.9%,
- 197 respectively.
- 198
- 199 3.4 Influence of iron concentrations on *M. laxa* germination and germ tube elongation
- After 6 h of incubation at 25 °C in MM broth without iron and in MM broth with $\frac{5 \ \mu g \ ml^{-1} \ of \ FeCl_{3}}{1}$
- 201 *M. laxa* conidial germination was reduced by both L1 and L8 strain of 50%, as showed in Table 3.
- 202 The presence of 10 μ g ml⁻¹ of FeCl₃ in the culture medium induced a higher antagonistic activity by
- 203 both strains against *M. laxa* conidial germination with respect to the other iron concentrations (5
- and 20 μ g ml⁻¹ of FeCl₃). In effect, conidial germination of *M. laxa* was reduced by 62.5% and
- 205 68.7%, respectively by L1 and L8. Conversely, 20 µg ml⁻¹ of FeCl₃ inhibited the strains antagonistic
- activity, that showed a reduction of *M. laxa* conidia germination of \sim 35%.

207	As regards to M. laxa germ tube length, the highest inhibition values exerted by L1 and L8 were
208	mainly showed with 5 and 10 μ g ml ⁻¹ of FeCl ₃ solution by 59.5%, 63.1% and by 65.9%, 67.6%,
209	respectively. Nevertheless, 10 μ g ml ⁻¹ of FeCl ₃ stimulated mostly pathogen germ tube elongation
210	(10.4%) with respect to the other concentrations (Table 4). Anyway, L8 strain was always slightly
211	more effective with respect to L1 in determining a higher inhibition of <i>M. laxa</i> conidial germination
212	and germ tube elongation on average by 52.1%, 50.2%, and by 57.7%, 57.1%, respectively.
213	
214	3.5 <i>In vivo</i> competition for iron
215	The influence of the addition of aliquots of iron solutions at different concentrations on fungal
216	virulence and yeasts antagonistic activity on peach fruit was measured. Monilinia laxa growth on
217	peach fruit in absence of antagonists was slightly enhanced by 6.8% only at FeCl ₃ concentrations of
218	10 or 20 μ g ml ⁻¹ whereas no influence was observed on the pathogen aggressiveness at 5 μ g ml ⁻¹ ,
219	the lowest FeCl ₃ concentrations (Fig. 3).
220	However, L1 and L8 A. pullulans strains showed a great capability to inhibit M. laxa
221	aggressiveness, reducing the peach lesion diameters by 83.5% and 84.4% on average respectively,
222	at each iron concentration. Furthermore, the highest concentration of iron (20 µg ml ⁻¹) seemed to
223	slow down the antagonists action and conversely increase the pathogen aggressiveness. In fact, both
224	L1 and L8 controlled <i>M</i> . <i>laxa</i> incidence in presence of 20 μ g ml ⁻¹ FeCl ₃ on average by 88% with
225	respect to 92%, in presence of the other iron concentrations (data not reported).
226	
227	4. Discussion
228	Aureobasidium pullulans (L1 and L8) are considered promising alternatives to fungicides in the

Aureobasidium pullulans (L1 and L8) are considered promising alternatives to fungicides in the reduction of the stone fruits postharvest diseases' incidence caused by *Monilinia* spp. (Mari et al., 2012; Di Francesco et al., 2017). In our work, competition for nutrients was studied and in particular the competition for iron, a nutrient that played a significant role in biocontrol interactions (Raaska and Mattila-Sandholm 1995).

As known, iron is an essential nutrient for all organisms (Wang et al., 2009) acting as a cofactor for 233 enzymes and regulatory proteins involved in many cellular processes (Miethke and Marahiel 2007; 234 Li et al., 2018) but not promptly available due to the low solubility in alkaline environments. One of 235 the main strategy used by microorganisms (such as yeasts) and by plants to obtain iron is the 236 secretion of siderophores (Li et al., 2018), which are low-molecular-weight compounds that chelate 237 iron in the extracellular phase re-entering it in the cells by specific membrane transporters (Hider 238 and Kong 2010). More than 500 siderophores have been discovered from microorganisms and 239 plants and their chemical diversity often renders a variety of biological functions beyond capture 240 iron (Ho et al., 2019). Yeasts produce only hydroxamate-type siderophore (Riquelme 1996) derived 241 from the amino acid ornithine and classified into four structural families: fusarines, coprogens, 242 ferrichromes and rhodotorulic acid (Johnson 2008). Wang et al., (2009) showed that A. pullulans 243 could produce 1.1 mg ml⁻¹ hydroxamate type siderophore. In our experiments, both L1 and L8 were 244 evaluated for their capability to produce siderophore by CAS screening assay (Schwyn and 245 Neilands 1987). This assay was used for the screening of siderophore producing microorganisms by 246 the induction of a colour change of the growth medium: from blue to orange, red, or purple. 247

Our results showed that L1 and L8 strains produced siderophore, displaying after twelve days of 248 incubation a colour change of the CAS agar, from blue to red. Moreover, L8 strain induced a more 249 250 intense agar colour change and extended halo with respect to L1. Conversely, M. laxa was not able to produce siderophores in solid medium but produced a clear barrage between the common fungal 251 culture medium (PDA) and the siderophore screening agar (CAS). In fact, CAS agar assay is mainly 252 useful for the identification of microorganisms capable to produce siderophores in high 253 concentrations and characterized by a high affinity for iron (III). Most likely, the assay was not so 254 sensitive to detect the low siderophores concentration produced by *M. laxa*. 255

In effect, these results were supported by the quantitative analysis conducted with the spectrophotometer. Both yeasts were found to produce high levels of chelating molecules (1.2 and 1.4 mg ml⁻¹, respectively) whit respect to the pathogen that seemed instead to be a poor siderophores

producer (0.64 mg ml⁻¹). Probably siderophores role vary among different fungal pathosystems (Chen et al., 2013) and according to the strain intrinsic characteristics. In fact, the importance of siderophores for fungal pathogenesis was firstly demonstrated for the pathogens such as *Cochliobulus heterostrophus* (Lee et al., 2005; Oide et al., 2006), *Alternaria* spp. (Chen et al., 2013), *Ustilago maydis* (Mei et al., 1993) and for the human pathogen *Aspergillus fumigatus* (Hissen et al., 2005; Schrettl et al., 2007; Chen et al., 2013).

In our study, the siderophore production by the yeasts was not affected by the presence of *M. laxa*, probably for the nutritional and biochemical preferences of the fungus. According to these results, we can suppose that *M. laxa* probably mainly compete for other nutrients rather than for iron; on the other side, *A. pullulans* strains L1 and L8 showed a high effective antagonistic action against *M. laxa* in presence of iron (10 μ g l⁻¹ FeCl₃), displaying a great capability to create critical conditions for the pathogen growth.

As other microorganisms (Droby et al., 1989, Bencheqroun et al., 2006) both L1 and L8, reduced their antagonistic efficacy with the addition of high concentrations of exogenous nutrients, e.g. with $273 \quad 20 \ \mu g \ l^{-1}$ of FeCl₃ both in *in vitro* and in *in vivo* assays.

In fact, the highest reduction of conidial germination, brown rot severity and incidence (data not shown) in fruit was revealed in presence of lower iron concentrations, confirming that both strains displayed their antagonistic efficacy when exogenous nutrients were scarce (Calvente et al., 1999, Calvo et al., 2008, Sanz Ferramola et al., 2013). Plante and Labbè (2019) showed how in *Schizosaccaromyces pombe* the siderophore synthetase expression remains upregulated under low iron conditions.

According with previous observations on nutritional requirements of *Monilinia* spp. germination (Byrde and Willetts 1977), in the present research, the germination rate of the pathogen conidia resulted low when grown in minimal medium. This confirmed that the target pathogen is nutrientdependent and needs an adequate amount of supplements for its growth: both conidial germination and hyphal development (Bencheqroun et al., 2006, Di Francesco et al., 2017). Moreover, the two

285	BCAs notably reduced <i>M. laxa</i> germ tube elongation: a fungal characteristic directly related to the
286	pathogen aggressiveness (Fujii 2004). In fact, one critical step of germination is the germ tube
287	elongation that can be critically blocked by iron deficiency (Plante and Labbè 2019), in this case
288	caused by the antagonistic action of both L1 and L8 to compete for nutrients.
289	In conclusion, A. pullulans L1 and L8 strains can compete with M. laxa for nutrients such as amino
290	acids as previously reported by Di Francesco et al. (2017, 2018a) and in the present study for iron, a
291	vital element for cell proliferation and survival for almost all organisms. Therefore, the ability of A.

292 *pullulans* strains L1 and L8 to acquire iron can counteract the expression of fungal pathogen genes

required for the production of siderophores and virulence in fruits (Chen et al., 2013).

This information can be very important to exploit L1 and L8 activity in a commercial formulation characterized by a right and efficient composition. More investigations are necessary to isolate and characterize siderophores produced by both yeast strains and possibly verify their environmental involvement in absorption and metabolization of metals. Nevertheless, in our study the strong efficacy exerted by L1 and L8 strains can be attributed most likely to a synergic activity of different mechanisms of action.

300

301 Conflict of Interest

302 The authors declare that they have no conflict of interests.

303

304 Authors contribution

Dr. Alessandra Di Francesco conducted all the experiments, interpreted data, and drafted the
 manuscript. Prof. Elena Baraldi coordinated the experimental plan, has corrected, and approved the
 final draft of the manuscript.

308

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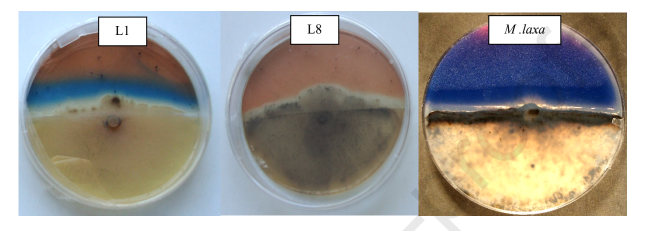
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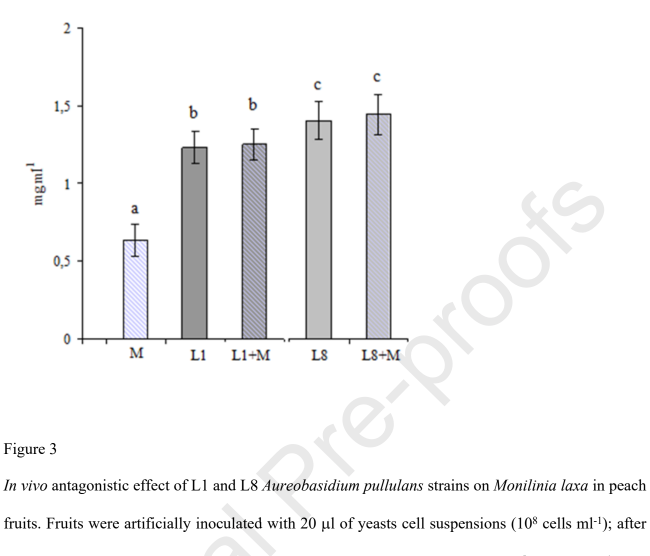
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- 475
- 476
- 477 Figure 1
- 478 CAS assay performed with Aureobasidium pullulans L1, L8 strains and Monilinia laxa (isolate
- 479 ML4).



- 480 481
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- 486 Figure 2

487 Quantitative (mg ml⁻¹) determination of siderophore at 440 nm by spectrophotometer, on 488 deferoxamine mesylate standard curve, of *Aureobasidium pullulans* L1 and L8 strains and 489 *Monilinia laxa* (M), alone and co-cultured. Data are the means of two independent 490 experiments \pm standard error, each consisting of three flasks per treatment. Different letters indicate 491 significant differences at α =0.05 according to *Tukey's* HSD Test.



air drying they were inoculated with 20 µl of *M. laxa* conidia suspension (10⁵ conidia ml⁻¹) and finally with an iron suspension (0, 5, 10 and 20 µg ml⁻¹ FeCl₃) (0, 5, 10, 20). Control consisted of peaches inoculated with water instead of yeasts suspensions. Data are the means of two independent experiments ± standard error, each consisting of 45 fruit per treatment. For each treatment group (Control, L1, and L8) different letters indicate significant differences at $\alpha = 0.05$ according to *Tukey's* HSD Test.

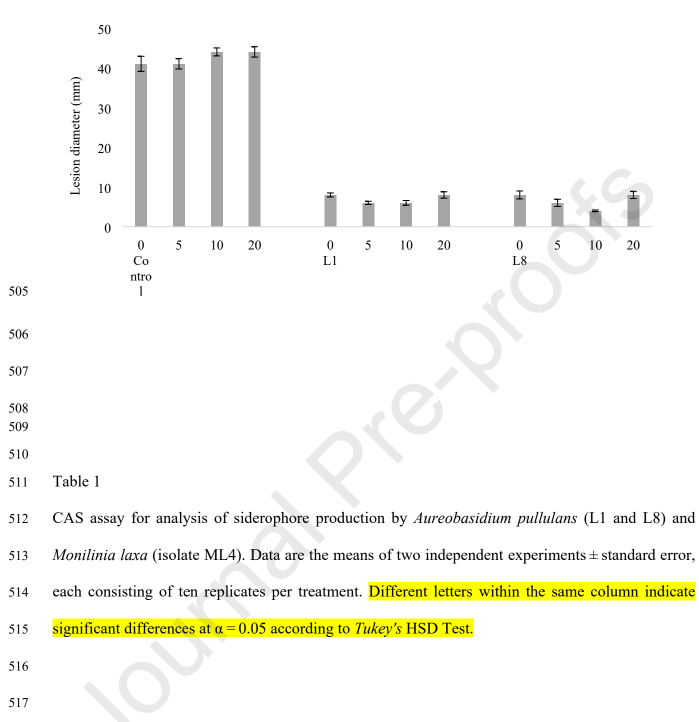
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Microorganism Growth tim		Color change	CAS reaction **
	(days)*		
Monilinia laxa	10±0.4b	-	0±0.0a
L1	12±0.0a	dark orange	20±1.9b
L8	12±0.0a	dark orange	27±1.5c

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519	*Days required for the fungal mycelium to cover the non-CAS half of the plate.
520	** mm of advance of colour change front in the CAS blue agar after three weeks of incubation.
521 522	
523	Table 2
524	Mycelium diameter (mm) of Monilinia laxa growth in dual culture with Aureobasidium pullulans
525	L1 and L8 strains in minimal medium agar plates amended with 0, 5, 10 and 20 μ g ml ⁻¹ FeCl ₃ and
526	incubated at 25 °C for 5 days. Control plates were inoculated only with pathogen plug (6 mm).
527	Data are the means of two independent experiments \pm standard error, each consisting of five
528	replicates per treatment. Within the same FeCl ₃ treatment (column), the same lower-case letters
529	represent no significant differences at $\alpha = 0.05$ according to <i>Tukey's</i> HSD Test. Within the same
530	yeast strains or pathogen isolate (control) (row), the same upper-case letters represent no significant
531	differences at $\alpha = 0.05$ according to <i>Tukey's</i> HSD Test.

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	\sim	Treatment (FeCl ₃)			
	0 μg ml ⁻¹	5 µg ml ⁻¹	10 µg ml ⁻¹	20 µg ml-1	
Control	60.0±0.6cA	60.0±b1.2bA	63.0c±1.0cB	61.0±1.5cAB	
L1	35.2±1.9bB	35.0a±0.7aB	34.7b±1.3bA	34.9±1.1bA	
L8	34.0±1.1aB	34.4a±1.0aB	33.0a±0.8aA	33.6±0.8aAB	
T 11 0					

538 Table 3

Effect of *Aureobasidium pullulans* (L1 and L8) on conidia germination (%) of *Monilinia laxa* grown in culture medium (minimal medium) amended with 5, 10 or 20 μ g ml⁻¹ FeCl₃ and incubated

at 25°C for 6 hours. Data are the means of two independent experiments \pm standard error, each consisting of 90 conidia per treatment. Within the same FeCl₃ treatment (column), the same lower-

- 543 case letters represent no significant differences $a_t \alpha = 0.05$ according to *Tukey's* HSD Test.
- 544 Within the pathogen isolate (control) and yeast strains (row), the same upper-case letters represent
- 545 no significant differences at $\alpha = 0.05$ according to *Tukey's* HSD Test.
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		Treatment (FeCl ₃)				
	0μg ml ⁻¹	5 μg ml ⁻¹	10 µg ml ⁻¹	20 µg ml ⁻¹		
Control	40±1.3cA	41±1.8bA	48±0.8cC	43±1.5bB		
L1	20±2.1bB	19±1.7aB	18±1.4bA	28±2.1aC		
L8	19±0.9aB	19±1.5aB	15±1.2aA	29±1.8aC		

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552 Table 4

Effect of Aureobasidium pullulans (L1 and L8) on Monilinia laxa conidia germ tube elongation 553 (µm) grown in liquid culture medium (minimal medium) amended with 0, 5, 10 and 20 µg ml⁻¹ 554 FeCl₃ at 25 °C for 6 hours. Data are the means of two independent experiments ± standard error, 555 each consisting of 90 conidia per treatment. Within the same FeCl₃ treatment (column), the same 556 lower-case letters represent no significant differences significant differences at $\alpha = 0.05$ according 557 to *Tukey's* HSD Test. Within the pathogen isolate (control) and yeast strains (row), the same upper-558 case letters represent no significant differences significant differences at $\alpha = 0.05$ according to 559 Tukey's HSD Test. 560

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		Treatment (FeCl ₃)		
	0μg ml ⁻¹	5 µg ml ⁻¹	10 µg ml ⁻¹	20 µg ml ⁻¹
Control	16.2±1.3cA	16.3±1.6cA	18.2±2.1cB	16.5±1.8bA
L1	7.7±2.1aB	6.6±1.8bA	6.2±2.3bA	8.1±2.1aC
L8	8.3±1.5bC	6.0±1.6aB	5.7±3.5aA	8.2±1.8aC

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564 **Conflict of Interest**

- 565 The authors declare that they have no conflict of interests.
- 566

567 Authors contribution

568 Dr. Alessandra Di Francesco conducted all the experiments, interpreted data, and drafted the 569 manuscript. Prof. Elena Baraldi coordinated the experimental plan, has corrected, and approved the 570 final draft of the manuscript.

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573 Highlights

- Competition for nutrients is one of the most important mechanism of action for BCAs
- Iron plays a role in BCAs modes of action through the production of siderophores
- 577 Siderophores can inhibit the growth of certain pathogenic fungi
- 578 Aureobasidium pullulans L1 and L8 strains compete for iron against Monilinia laxa