Biocontrol Activity and Plant Growth Promotion Exerted by *Aureobasidium pullulans* Strains

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Abstract

The most common leguminous plants’ diseases are caused by soil-borne pathogens leading to important economic losses worldwide. Strains L1 and L8, belonging to *Aureobasidium pullulans* species, were tested in vitro and in vivo as biocontrol agents (BCAs) against *Rhizoctonia solani* (AG-4) and as plant growth promoters (PGPs). The non-volatile metabolites produced by L1 and L8 strains inhibited the pathogen mycelial growth by 87.9% on average, with no significant differences between the two strains. The lower pathogen diametric growth inhibition was displayed by both yeasts’ volatile metabolites (VOCs) that significantly reduced the colony growth of *R. solani*, and similarly to the control, with an average of 10.5%. By in vivo assay, L1 and L8 strains showed the ability to control the pathogen virulence probably through the biofilm formation around the bean and soybean plant roots, as confirmed by scanning electron microscope (SEM) analysis. The spectroscopic analysis highlighted the composition of non-volatile compounds: complex carbohydrates (pullulan), degrading enzymes, siderophores and antifungals (aureobasidins). Moreover, the ability of L1 and L8 strains to stimulate the bean and soybean plant roots, stems, and leaves growth was investigated, showing that these yeasts could have an application not only as BCAs but also as plant growth biostimulator.

Keywords Yeasts · Roots · Biofilm · Growth promoter · Metabolites

Introduction

*Rhizoctonia solani* represents one of the principal soil-borne pathogens that can attack a wide range of plant species, mainly leguminous plants, causing seed decay, damping-off, hypocotyl rot, and web blight (Thiessen and Woodward 2012; Müller Schmidh et al. 2018). In particular, *R. solani* belonging to anastomosis group AG-4 represents the causal agent of root rot of plants under greenhouse and field conditions causing great qualitative and economical damages (Ahed and Kamil 2013). Synthetic fungicides are the main systems for controlling plant diseases; however, there is a great concern for the wide use of chemical substances, due to the harmful effects on the environment and human health, and to the risk of pathogen resistance emergence. For these reasons, there is an increasing necessity to find alternative control strategies to overcome these disadvantages (Akladious et al. 2019). Alternative strategies, such as the use of antagonistic microorganisms, offer the potential to provide sustainable and long-term protection of crops (Kotze et al. 2011). The application of biological control agents (BCAs) is considered a safe and environmentally friendly alternative (Zhang et al. 2010) to synthetic fungicides. Among recent studies, the yeast *Aureobasidium pullulans* strains L1 and L8 (Di Francesco et al. 2018a) showed high efficacy against fruit postharvest pathogens (Di Francesco et al. 2017, 2018b, 2020a) such as *Monilinia* spp. and *Botrytis cinerea*, and also against endophytic pathogen such as *Neofusicoccum parvum* (Rusin et al. 2019).
Among soil microorganisms, yeasts have received little attention as biocontrol agents of soil-borne fungal plant pathogens in comparison to bacterial, actinomycetes, and filamentous fungal antagonists (El-Tarabily and Sivasithamparam 2006). Yeasts show similar mechanisms of action against soil-borne fungal plant pathogens and plant’s aerial parts pathogens (El-Tarabily and Sivasithamparam 2006). Their ability to multiply rapidly, to produce antifungal diffusible metabolites, and cell wall-degrading enzymes (CWDEs) (Kohl et al. 2019), is a hallmark for potential application as soil BCAs and plant growth promoters (PGPs).

In the present study, the efficacy of volatile and non-volatile metabolites, produced by A. pullulans strains L1 and L8 in inhibiting growth and enzymatic activity of the soil-borne pathogen R. solani of bean and soybean plants, was assayed in vitro. Moreover, the composition of the non-volatile compounds was analyzed for the first time through Fourier transform infrared spectroscopy (FT-IR) and Raman spectroscopies. In particular, the biofilm formation (polysaccharide matrices) of both strains was confirmed by scanning electronic microscopy (SEM). Subsequently, the antagonistic activity of both strains was evaluated by in vivo assays against the pathogen in bean and soybean plants. This paper describes the first application of both strains as BCAs of a soil-borne fungal plant pathogen and as PGP.

Materials and Methods

Antagonists

Aureobasidium pullulans strains L1 and L8, molecularly characterized by Di Francesco et al. (2018a), were grown on nutrient yeast dextrose agar (NYDA: 8 g of nutrient broth, 5 g of yeast extract, 10 g of dextrose, and 25 g of technical agar, Oxoid Basingstoke—UK) in 1 L of distilled water. Each antagonist was grown on NYDA at 25 °C for 48 h before the beginning of the experiments, and yeast cells were collected in sterile distilled water containing 0.05% (v/v) Tween 80 (Sigma-Aldrich, St. Louis, MO, USA) then adjusted with a hemocytometer to a final concentration of 10⁶ cells/mL.

Pathogen

Rhizoctonia solani isolate (Rs1) used in the present experiments was obtained from symptomatic bean plants (Phaseolus vulgaris L.) grown in an experimental field located in Cadriano (Bologna, Italy), molecularly identified, and deposited to Criof mycological collection. The isolate was chosen for its highly virulent behavior. For in vitro experiments, the isolate was maintained on potato dextrose agar (PDA, 39 g in 1 L of distilled water) (Oxoid, Basingstoke, UK) at 25 °C for 5 days until use.

For in vivo experiments, the inoculum for infecting plants was prepared by cutting 6 mm diameter plugs taken from the margin of 5 days-old fungal colony and was transferred to flasks with 50 mL of potato dextrose broth (PDB, 39 g in 1 L of distilled water) (Oxoid, Basingstoke, UK) and incubated at 25 °C in a rotary shaker (250 rpm) for 5 days. The culture was then centrifuged at 5000xg for 20 min at 4 °C, and the mycelium was washed with distilled water, filtered through two layers of miracloth, finally homogenized by using a mixer (Imetec Ch4, Italy) and immediately used for the inoculation of plant pots.

Plants

Bean (Phaseolus vulgaris L.) and soybean (Glycine max L.) plants were obtained from seeds cv ‘Borlotto’ and cv ‘Demetra’ respectively (Syngenta, Switzerland) seeded in 50 mL pots filled with moss, until the 2 expanded leaves stage (15 days), under daylight conditions at 25 ± 2 °C and 60% of relative humidity (RH).

In Vitro Assays: Yeasts Metabolites Effect on Rhizoctonia solani Mycelial Growth

Metabolites produced by both strains L1 and L8 were assayed on R. solani mycelial growth by three in vitro experiments. The yeast-pathogen direct interaction was firstly determined in co-culturing conditions, and then the effect of volatile and non-volatile compounds produced by both strains on pathogen growth was verified. The direct interaction assay between yeasts and the pathogen was assayed on PDA medium in 90 mm diameter Petri plates. Pathogen mycelial plug (6 mm diameter) cut from the margin of a 5 days-old colony was inoculated in the center of the plate and L1 and L8 cells, taken from 48 h cultures grown on NYDA, were streaked on the plate 22.5 mm from both sides of the pathogen plug through a sterile loop. In control plates, yeasts cells were not inoculated.

Plates were incubated at 25 °C for 5 days and the diameter of the pathogen colony was measured on a line passing along the center of the plug and perpendicular to yeast streaks, after 3, 4, and 5 days of incubation. For each strain and the control, 10 agar plates were used, and the assay was repeated twice. To verify the effect of volatile compounds produced by L1 and L8 strains on R. solani mycelium growth the double Petri dishes assay was used as previously described by Di Francesco et al. (2015). For this purpose, NYDA plates were inoculated by spreading 100 μL of antagonist cells suspension (10⁶ cells/mL). The lid of the plate was replaced, after 48 h of incubation at 25 °C, by a base plate of PDA inoculated with a 6 mm diameter pathogen plug cut.
from the margin of a 5 days-old colony. The two base plates were sealed immediately with a double layer of Parafilm and incubated at 25 °C for 5 days. Control was represented by plates inoculated only with the pathogen. To verify the further effect of yeasts metabolites on pathogen mycelium growth, at the end of the incubation time, the weight of each dish was measured. The sample unit was represented by ten plates (replicates) for the control and each antagonist. The assay was repeated twice. To verify the effect of L1 and L8 non-volatile metabolites, 100 μL of antagonist cells suspension (10^8 cells/mL) were spread on a sterile cellophane disc (Courtauld Films: 90 mm diameter) laying down on PDA medium plate and incubated at 25 °C for 48 h. The cellophane with the yeast was then removed and, a 6 mm diameter pathogen plug, cut from the margin of a 5 days-old colony and incubated at 25 °C for 5 days, was positioned at the center of the plate. One hundred microliters of sterile water were streaked on cellophane representing the control. The sample unit was represented by ten plates (replicates) for the control and each antagonist. The assay was repeated twice.

**Synthetic Volatile Compounds Effect on Rhizoctonia solani Mycelial Growth**

Di Francesco et al. (2015) identified through headspace solid-phase microextraction (HS-SPME-GCMS) the volatile compounds produced by L1 and L8. For this purpose, pure standards of 3-methyl-1-butanol, 2-methyl-1-butanol, 2-methyl-1-propanol and 2-phenylethanol (Sigma–Aldrich, St. Louis, MO) were tested on *R. solani* colony growth. Different aliquots of pure compounds 25, 50 and 100 μL were placed with a microsyringe on a filter paper (Whatman No. 1, 90 mm diameter) positioned inside the cover of a PDA dish previously inoculated with a 6 mm diameter pathogen plug cut from the margin of a 5 days-old colony. The aliquots of pure compounds introduced in the Petri dishes corresponded to 2.25, 1.12 and 0.56 μL/mL headspace. The dishes were quickly closed, sealed with Parafilm and incubated at 20 °C. The activity of each pure compound against mycelial growth was evaluated after 5 days of incubation. In the controls, pure compounds were substituted by equivalent amounts of distilled water. The sample unit was represented by 5 plates for each volatile compound concentration. EC_{50} values were calculated as the headspace concentrations (μL/mL) that inhibited mycelial growth by 50% compared to the control. The experiment was performed twice.

**Volatile Compounds Effect on Rhizoctonia solani Pathogenesis Enzymes Activity**

To verify the effect of volatile compounds produced by L1 and L8 strains on *R. solani* pathogenesis enzymes, the fungal mycelium derived from the double Petri dishes assay exposed or not to yeasts VOCs, was used to perform cellulase, xylanase, polygalacturonase, and pectinase assays. The enzymes agar composition was previously reported in Di Francesco et al. (2018c, 2020b). Assays were performed in Petri dishes containing specific agar medium. Twenty mycelial agar plugs (6 mm diameter) derived from each yeast VOCs treatment and the control were placed equidistant (3 cm) on each enzyme agar plate (30 × 30 × 2 cm. L × W × H) and incubated at 25 °C for 24 h. Enzymes activities were determined with specific colorimetric methods, described by Di Francesco et al. (2018c, 2020b), which have allowed visualizing a clear zone of substrate degradation around each plug. The diameter of degradation halos (mm) was measured with a ruler. Each enzyme assay was repeated twice.

**Non-volatile Metabolites Analysis**

Flasks containing 100 mL of nutrient yeast dextrose broth (NYDB) medium were inoculated with 100 μL of antagonists cells suspensions (10^8 cells/mL) and then incubated at 25 °C in a rotary shaker (250 rpm) for 1, 24 and 48 h. After each incubation time, the liquid cultures were centrifuged at 5000×g for 20 min at 15 °C and the supernatant was collected in sterile tubes (2 mL) and stored at −20 °C. The control consisted of growth medium without yeast cells inoculation. The supernatant was analyzed by two different spectoscopic techniques (FT-IR and Raman) to obtain a rapid and non-destructive characterization of their main molecular components. Infrared spectra were recorded with a Bruker ALPHA series FT-IR spectrophotometer (Bruker, Ettlingen, Germany) equipped with an apparatus for attenuated total reflectance (Diamond crystal). The spectra were collected from 4000 to 400 cm−1 and averaged over 100 scans (resolution = 4 cm−1): 4 spectra were measured for each sample for each sampling time. Raman spectra were recorded on a Bruker MultiRam FT-Raman spectrometer equipped with a cooled Ge-diode detector. The excitation source was a neodymium-doped yttrium aluminum garnet (Nd^{3+}-YAG) laser (1064 nm) in the backscattering configuration. The power delivered from the focused laser to sample has been measured to be ~200 mW. The spectra were collected from 4000 to 200 cm−1 and averaged over 10,000 scans (resolution = 4 cm−1).

**Biofilm Formation**

Biofilm formation was evaluated following a procedure previously described by Parafati et al. (2015), partially modified. Yeast strains were grown overnight at 25 °C in liquid medium (NYDB) in a rotary shaker (250 rpm). Cells were then collected, washed twice with phosphate-buffered saline
(PBS, pH 7.2), and re-suspended in NYDB to $10^7$ cells/mL. Aliquots of cell suspensions (100 μL) were inoculated into the wells of a 96-well polystyrene plate and incubated at 25 °C for 3 h in a rotary shaker (100 rpm); the medium without yeasts cells was used as control. After this period, the wells were washed once with 150 μL of PBS, and 100 μL of NYDB were transferred into each washed well. The plates were incubated at 25 °C in agitation (100 rpm) for 3, 24, and 48 h. The medium was replenished daily by aspiration of the spent medium and the addition of the fresh medium. After each incubation time, wells were washed twice with PBS and stained with 100 μL of 0.4% aqueous crystal violet solution for 45 min. The wells were washed four times with sterile distilled water (SDW) and destained with 200 μL of 95% ethanol. After 45 min, 100 μL of the destaining solution was transferred to a new 96-well plate and the amount of crystal violet was measured at 590 nm with a spectrophotometer (Infinite® F50 Tecan, Männedorf, CH). The controls’ absorbance values were subtracted from the test well values to minimize background interference. The assay was repeated twice.

**In Vivo Assays**

Roots plants were immersed in 25 mL of yeasts cells suspensions ($10^7$ cells/mL) for 1 h. Plants were then transplanted in a new plot where *R. solani* mycelium was inoculated 24 h earlier (1 g per 220 g of peat moss) (1 plant per pot). The control consisted of roots plants treated with water only. The sample unit was represented by ten plants (replicates) for each yeast strain and the control. After the stem-end symptom appearance (21 days) the disease incidence was evaluated and a Koch postulate was conducted.

Hypocotyls with severity of *R. solani* of each plant were evaluated using a 0–4 scale developed by Cardoso and Echandi (1987) with some modifications, where 0 = no lesions, 1 = lesions < 2.5 mm, 2 = lesions 2.5–5 mm, 3 = lesions > 5 mm, 4 = lesions girdling plant and wilting visible on leaves. A disease severity index (DSI) was calculated for each treatment using the formula:

$$DSI = \frac{\text{percentage of incidence} \times \text{mean severity}}{5}$$

To evaluate the yeast strains growth promotion effect on bean and soybean plants (10 plants for each strain and the control) without pathogen mycelium inoculation, the length of the roots (mm), the stems diameter (mm) and the leaves areas (mm²) were measured after 21 days from the treatment of the roots by using a digital caliber (Dirtygal) and Tomato Analyzer Software (Brewer et al. 2006), respectively. Treatment effects were assayed in two independent experiments. The control consisted of roots plants treated with water only.

**Scanning Electron Microscopy Observation**

The biofilm formation on bean and soybean plants roots was investigated by SEM observations. Plant roots were treated with 25 mL of yeasts cells suspensions ($10^7$ cells/mL) and the biofilm assessments were carried out after 3, 24 and 48 h from the treatment. Little portions of roots tissues (3 × 3 × 3 mm) were excised, washed three times for 7 min in 0.1 M Phosphate buffer pH 7.2. Three samples from each plant were fixed with glutaraldehyde 3% in phosphate buffer solution at pH 7.4. Before observation, the fixed tissues were rinsed three times with phosphate buffer (pH 7.4), dehydrated in increasing ethanol solutions (20, 30, 50, 75, 95, 100%), and with the use of a conductive 3 M 465kp scotch tape were set on aluminum pin stubs. Samples were then coated with gold particles using Emitech K500 sputter at 40 mA for 3 min. Samples were observed under SEM (Philips SEM 515 microscope at 30 kV).

Fifteen samples derived from 5 plants were examined for each plant species, sampling time and treatment.

**Statistical Analysis**

All experiments were carried out in a completely randomized design. Obtained data were analyzed by one-way ANOVA ($P < 0.05$). Separation of means was performed using the least significant difference (LDS) test, at $P < 0.05$. All analyses were performed with the software Statgraphic Plus Version 2.1 (Statistical Graphics Corp., USA 1996).

**Results**

**In Vitro Assays: Yeasts Metabolites Effect on *Rhizoctonia solani* Mycelial Growth**

The effects of L1 and L8 strains on *R. solani* mycelial growth through the direct interaction in the co-culturing condition and the exposure to non-volatile and volatile metabolites produced by the strains are reported in Table 1. In the yeast-pathogen co-culturing experiment, both L1 and L8 strains significantly reduced the pathogen colony growth compared to the control without a substantial difference between the two antagonists. However, the pathogen colony growth was inhibited by 57.3% by both strains. The non-volatile metabolites produced by L1 and L8 inhibited the pathogen mycelial growth of about 87.9%, with no significant differences among the two strains. The lower pathogen colony growth inhibition was displayed by both yeasts volatile metabolites and non-volatile metabolites. In fact, the colony growth of *R. solani* was significantly reduced similarly by both strains of about 10.5% compared to the control. Nevertheless, the volatile compounds produced by both strains reduced by 60% the
pathogen mycelial weight (data not shown). Non-volatile metabolites from both strains were significantly more effective than volatiles.

**Synthetic Volatile Compounds Effect on *Rhizoctonia solani* Mycelial Growth**

The effect of pure volatile compounds on *R. solani* colony growth is reported in Table 2. Among the four tested compounds, 2-methyl-1-butanol, 3-methyl-1-butanol, and 2-phenylethanol showed the highest efficacy in reducing pathogen growth, with EC$_{50}$ values of 0.18, 0.13, and 0.26 µL/mL, respectively. The least effective compound was 2-methyl-1-propanol with an EC$_{50}$ value of 1.50 µL/mL.

**Volatile Compounds Effect on *Rhizoctonia solani* Pathogenesis Enzymes Activity**

The effect of VOCs produced by L1 and L8 was tested on the pathogenesis enzymes activity of *R. solani*. Both VOCs strains constrained the pathogenesis activity of the fungus, showing a clear reduction of the diameter of the enzymatic halo (mm) (Fig. 1). Both strains inhibited, on average, the tested fungal pathogenesis enzymes activity by 17.9%, 7.3%, 8.4%, 36.1% respectively for cellulase, xylanase, polygalacturonase, and pectinase, displaying an indirect action against *R. solani* pathogenic activity. Furthermore, the xylanase enzyme resulted in the most produced by the pathogen and polygalacturonase the least.
Non-Volatile Compounds Analysis

The spectroscopic measurements were carried out to determine the main non-volatile metabolites production by *A. pullulans* strains L1 and L8 in the liquid culture medium after 1, 24, and 48 h of incubation at 25 °C. Figures 2 and 3 show the main peaks related to the metabolites groups detected after 1, 24, and 48 h of incubation by FT-IR and Raman spectroscopies, respectively. More in detail, Figs. 2a and 3a show the FT-IR and Raman spectra of L1 metabolites, Figs. 2b and 3b the FT-IR and Raman difference spectra between L1 and the control medium, and Figs. 2c and 3c the FT-IR and Raman difference spectra between L1 and L8 strains after 48 h. Spectroscopic peaks were attributed to five classes of compounds that are highlighted with different colors in the figures: nutrient medium (black peaks); complex carbohydrates (Bozoudi and Tsaltas 2018), as pullulan and other β-glucans (blue peaks); lytic enzymes (Bozoudi and Tsaltas 2018), as amylases, cellulases, lipases, xylanases, proteases, laccase and mannanases (red peaks); antifungal compounds (Bozoudi and Tsaltas 2018), as aureobasidins, derivatives of cyclic depsipeptides possessing, as typical amino acids, proline, leucine and phenylalanine (green peaks); siderophores (Bozoudi and Tsaltas 2018) (purple peaks).

The peaks attributed to the nutrient medium were derived from the FT-IR and Raman spectra of pure NYDB (data not shown), while the attribution of carbohydrates, enzymes, siderophores and aureobasidins is based on the existing literature (Aghatabay et al. 2014; Bae et al. 2015; Cozar et al. 2006; Dahiya et al. 2016; Shingel 2002; Tu 1982; Um et al. 2013; Zhbankov et al. 2000). Different spectra between L1 and the medium control (Figs. 2b and 3b) were useful to discriminate the main metabolites produced by the strain (positive peaks) and the medium itself (negative peaks). In analogy, different spectra between L1 and L8 strains (Figs. 2c and 3c) allowed highlighting the production of the different metabolites by the two strains. Focusing the attention on these last two spectra (FT-IR spectrum: Fig. 2c and Raman spectrum: Fig. 3c), it is possible to state that L1 strain showed a higher production of complex carbohydrates and enzymes: the former showed negative peaks at 932 and 848 cm⁻¹ in the IR spectrum (Fig. 2c, blue peaks)
and 937 and 581 cm\(^{-1}\) in the Raman spectrum (Fig. 3c, blue peaks); the latter showed negative peaks at 1644, 1516 and 1395 cm\(^{-1}\) in the IR spectrum (Fig. 2c, red peaks) and 2918 and 1442 cm\(^{-1}\) in the Raman spectrum (Fig. 3c, red peaks). Interestingly, Fig. 2c also shows some positive peaks attributed to pullulan production at 1340, 1076 and 1046 cm\(^{-1}\); all these peaks are reported to be sensitive to the chain length and the molecular weight of polysaccharides (Singhel et al. 2002) and indicate a reduction of their polymeric chain. The behavior of aureobasidins and siderophores production is more complex since the difference spectra between the two strains showed both negative (1577, 1328 and 534 cm\(^{-1}\), Fig. 2c; 482 cm\(^{-1}\), Fig. 3c) and positive peaks (2969, 699 and 630 cm\(^{-1}\), Fig. 2c and 625 cm\(^{-1}\), Fig. 3c): this finding could suggest that the two strains produce different antifungal and siderophores compounds.

**Biofilm Formation**

Spectrophotometer assay tested the biofilm formation ability of L1 and L8 strains after 3, 24 and 48 h, as described in the previous section. *Aureobasidium pullulans* L1 strain after 3 h showed a great ability to adhere to the polystyrene plates after repeated washes (OD 1.00 ± 0.07), on the contrary L8 adheres less displaying an OD value of 0.26 ± 0.07. Experiments also demonstrated that both strains, L1 and L8, maintained high film-forming capacity until 48 h (OD 1.8 ± 0.18 and 1.3 ± 0.15, respectively). *Aureobasidium pullulans* strain L1 exhibited a biofilm formation phase peak at 24 h of growth (Fig. 4).

**In Vivo Assay**

The efficacy of both L1 and L8 strains against *R. solani* in in vivo experiments was of 80% in bean plants, and 80% and 50% in soybean plant respectively, compared to the control (Table 3). Moreover, the DSI in bean was reduced by 94% by both strains and in soybean by 94% and 71% by L1 and L8 respectively, compared to the control.

Moreover, strain L1 especially showed a stimulant effect on plant roots length, increasing by 20.4% and 24.4% bean and soybean roots length, respectively (Table 4) (Supplemental Fig. 1a). Also, the stem diameter and leaf area of both plants underwent a dimensional increase when grown with the yeasts compared to the control. For bean stem, an increase of 30% by L1 and 16.6% by L8 and for soybean
stem of 53.8% only by L1 was registered. For bean leaf, L1 and L8 more than doubled the areas. An increase of 48.3% and 17.8% by L1 and L8 respectively was registered for soybean leaf areas (Supplemental Fig. 1b).

**SEM Observations**

The development of biofilm structure by *A. pullulans* L1 and L8 strains on bean and soybean roots was assessed by observation under scanning electron microscope after 3, 24 and 48 h from the inoculation at room temperature, as reported in Fig. 5. Only images of L1 on bean are reported since no optical differences were founded between the two strains and the plant species. A great clustering of yeast cells formed all around the bean roots from the very first sampling time (3 h) (Fig. 5b).

Both strains colonized the surface of the roots by the biofilm formation preventing the pathogen attachment and colonization. Forty-eight hours after the inoculation, the surface of the roots was still totally covered by the yeast cells demonstrating how the biofilm structure is capable of protecting from the pathogen attack (Fig. 5d).

**Discussion**

In this study, *A. pullulans* L1 and L8 strains revealed the capability to control a soil-borne pathogen of bean and soybean plants, such as *R. solani*.

Yeasts have always received little attention as BCAs of soil pathogens compared to other microorganisms (El-Tarabily and Sivasithamparam 2006). The presence of yeasts in the soil principally depends on the amount of available nutrients and soil factors such as compaction and porosity, rainfall, cultivation, the presence of soil-inhabiting insects (Phaff et al. 1978). For these reasons, yeasts were mainly studied for the biocontrol of postharvest and foliar diseases probably because they are a major component of the leaves and other plants’ aerial parts surfaces (Wilson and Wisniewski 1989; McLaughlin et al. 1990; Filonow 1998). The most frequently
genera of soil yeasts are *Candida* spp., *Cryptococcus* spp., *Rhodotorula* spp., and *Aureobasidium* spp. (Spencer and Spencer 1997).

Our results also showed the potential of *A. pullulans* L1 and L8 strains to be involved in promoting leguminous plant growth. The two strains showed a different efficacy against the target pathogen, but both displayed the same main mechanisms of actions against *R. solani*, as suggested by the fact that both strains produced non-volatile metabolites and formed biofilm around leguminous roots.

The detected non-volatile organic compounds, mainly complex carbohydrates (such as pullulan or other β-glucans), degrading enzymes, cyclic depsipeptides and siderophores, inhibited the pathogen mycelial growth by 87.9%. More in detail, spectroscopic data (FT-IR and Raman techniques) revealed that L1 strain has higher production of lytic enzymes and complex carbohydrates. Pullulan and other polysaccharides may improve the biofilm formation and therefore adhesion, thus explaining the better performance of L1 on polystyrene plates. Moreover, FT-IR spectra showed that the polysaccharide production by L8 strain is characterized by a reduced chain length and average molecular weight of the polymer (Shingel et al. 2002), thus leading to a less efficient biofilm formation. Interestingly, the highest inhibition showed by L1 in in vivo test, can be related to a higher production of degrading enzymes (i.e., amylases, proteases or lipases) and to a different production of aureobasidins, a class of antifungal compounds typical of *A. pullulans* (Bozoudi and Tsaltas 2018) and siderophores. Aureobasidins are derivatives of cyclic depsipeptides, which were reported to be synthesized also by *Streptomyces* (Um et al. 2013; Bae et al. 2015) and by marine sponges (Dahiya et al. 2016) and possessing antifungal properties. Different strains of *A. pullulans* were reported to produce a different mix of these compounds which contain peculiar amino acids such as proline, leucine and phenylalanine (Takesako et al. 1991).

Wang et al. 2009, first characterized a hydroxamate siderophore produced by a strain of *A. pullulans*, demonstrating a strong inhibition towards the pathogenic *Vibrio anguillarum*. Both FT-IR and Raman spectra confirmed the presence of hydroxamate derivatives by several diagnostic bands (Cozar et al. 2006) (1564, 1330, 630 and 530 cm$^{-1}$ in the IR spectra and 1292, 880, 640, 620 and 480 cm$^{-1}$ in the Raman spectra). The production of the siderophores in L1 and L8 strains and their chemical nature is under investigation and will be assessed in a future study. Following our results, the synthesis of antibiotics compounds can be considered as an important system for the inhibition of soil-borne pathogen growth (Rothrock and Gottlieb 1981; Punja and Utkhede 2003).

VOCs displayed a reduced inhibitory effect on *R. solani* mycelium growth (10.5%) but at the same time significantly influenced the production of CWDEs of the soil pathogen.

![Fig. 5](image-url) Scanning electron micrographs (SEM) of bean roots (surface) treated with water (control) (a) and *Aureobasidium pullulans* (L1 strain) cells suspension (10$^7$ cells/mL). Root tissues were collected at different times from the treatment (3, 24, 48 h) (b–d, respectively). SEM shown, 100× magnification in all images and 10× in the details images. Only images of L1 on bean roots were reported as no significant optical differences were founded between the two strains and the plant species.
VOCs seemed to inhibit mainly the activity of pectinase and cellulase pathogenic enzymes secreted by *R. solani*. It can be assumed that VOCs could reduce the decomposition of plant tissues by activating antioxidant enzymes to reduce plant cell damage and activate disease resistance enzymes to enhance the resistance of bean or soybean plants to *R. solani* and inhibit its growth.

Conversely, previous studies presented a great inhibitory effect exerted by L1 and L8 VOCs against the main fruit postharvest pathogens (Di Francesco et al. 2015, 2020a). This aspect supports the hypothesis that each antagonist employs different mechanisms to suppress different pathogens (Whipps 2001; Botha 2011) and its efficacy varies among the different pathosystems.

Biofilm formation is considered an important and strong mechanism of postharvest antagonistic microorganisms during the colonization of the substrate and fruit wounds (Klein and Kupper 2018) to prevent fungal attack. In this study, L1 and L8 showed the ability to rapidly colonize plant roots, one of the principal requirements for a BCA of soil-borne fungal pathogens, probably enhanced by the root's exudates (Van Overbeek and Van Elsas 1995). Rhizosphere ability should be considered a prerequisite for successful biological control of root diseases, and lack of reliable biological control observed in many studies may be related to the failure to adequately colonize roots (Weller 1988).

Our results indicated how both strains began to produce cells biofilm starting from 3 h by roots contact. After only 48 h, the surface of the root was totally covered by cells biofilm. The strains might preemptively colonize the rhizosphere and root without directly competing for the same site with the pathogen.

El-Tarabily (2004) reported that *C. valida* and *T. asahii* colonized 95% of roots by 6 days after radical emergence, whereas *R. glutinis* colonized 90% of roots after 8 days, showing how roots could be colonized at different times by different yeast species.

Considerable attention must be given to the detected ability of L1 and L8 strains to promote bean and soybean plant growth, in particular of roots and leaves. Plant growth-promoter microorganisms (PGPs) can act directly or indirectly. Indirect effects are mainly related to the production of siderophores (Buysens et al. 1996) which increase plant growth by decreasing the activities of pathogens. Our strains previously displayed the ability to produce a considerable amount of siderophores (1.2 and 1.4 mg/mL by L1 and L8 respectively) (article under review) competing for peat nutrients with *Monilinia laxa*, brown rot agent of stone fruit. As previously mentioned, spectroscopic investigations have highlighted the presence of siderophores in both L1 and L8 strains. However, PGP can directly affect plant physiological conditions promoting the nitrogen fixation (Cocking 2003) or the production of auxins, gibberellins, and cytokinins (Bottini et al. 2004; Kuklinsky-Sobral et al. 2004).

Recently, there is great interest in the agricultural sector to use PGPs (e.g., microorganisms, plant extracts, polyamines) (Compant et al. 2005) independently from pathogen presence, and additionally for reducing the chemical inputs (Gerhardson 2002).

For these reasons, an investigation of the capability of both strains to induce resistance in the treated plants can be foreseen. Verification of the siderophore production and the non-volatile compounds involvement in the inhibition of the target soil pathogen could be planned. Similarly, as previously reported for apple wood (Rusin et al. 2019), the role of both strains as endophytic microorganisms in plant roots warrant examination.

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**Compliance with ethical standards**

**Conflict of interest** The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

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