The Compound 2-Hexyl, 5-Propyl Resorcinol Has a Key Role in Biofilm Formation by the Biocontrol Rhizobacterium Pseudomonas chlororaphis PCL1606

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The production of the compound 2-hexyl-5-propyl resorcinol (HPR) by the biocontrol rhizobacterium Pseudomonas chlororaphis PCL1606 (PcPCL1606) is crucial for fungal antagonism and biocontrol activity that protects plants against the phytopathogenic fungus Rosellinia necatrix. The production of HPR is also involved in avocado root colonization during the biocontrol process. This pleiotrophic response prompted us to study the potential role of HPR production in biofilm formation. The swimming motility of PcPLL1606 is enhanced by the disruption of HPR production. Mutants impaired in HPR production, revealed that adhesion, colony morphology, and typical air–liquid interphase pellicles were all dependent on HPR production. The role of HPR production in biofilm architecture was also analyzed in flow chamber experiments. These experiments revealed that the HPR mutant cells had less tight unions than those producing HPR, suggesting an involvement of HPR in the production of the biofilm matrix.

Keywords: antifungal, biocontrol, biofilm, motility, adhesion, confocal laser scanning microscopy

INTRODUCTION

Members of the genus Pseudomonas possess a substantial amount of metabolic diversity, and many of them are able to colonize a wide range of niches (Madigan and Martinko, 2015). Pseudomonas spp. can produce a variety of metabolites, many of them inhibitory to other microorganism and are also involved in the biological control of plant pathogens (Morrissey et al., 2004; Gross and Loper, 2009; Raaijmakers and Mazzola, 2012). The biocontrol mechanisms that Pseudomonas spp. display against soilborne plant pathogens include the production of antibiotics, hydrolytic enzymes, and volatile organic compounds, competition for space and nutrients, and the induction of systematic resistance in plants (Raza et al., 2013;
Highly in depth research has been performed on the exploration and identification of the antibiotics produced by different Pseudomonas spp., biocontrol pseudomonads frequently produce more than one antimicrobial compound (Haas and Keel, 2003). The most well-known antibiotics produced by Pseudomonas spp. include 2,4-diacytethylphloroglucinol (2,4-DAPG), phenazines (PHZs), pyrolnitrin (PRN), pyoluteorin (PLT), hydrogen cyanide (HCN), and 2-hexyl-5-propyl resorcinol (HPR; Cazorla et al., 2006; Gross and Loper, 2009). These antibiotics could be directly involved in other different phenotypes related to biocontrol ability in addition to antagonism, such as plant growth promotion and niche competition (Weller, 2007; Ramette et al., 2011; Wang et al., 2015; Raio et al., 2017).

Pseudomonas chlororaphis PCL1606 (PcPCL1606) is a biocontrol agent able to suppress plant diseases caused by different soilborne phytopathogenic fungi (Cazorla et al., 2006). Previous studies revealed that this rhizobacterium produces the three antifungal compounds HPR, PRN, and HCN; however, only HPR has been demonstrated to be directly involved in the antagonism and the biocontrol ability of this strain (Calderón et al., 2015). HPR is a small molecule, which belongs to the group of alkyresorcinols, produced by different bacteria. This compound is liberated from the cell to the environment, where display some antimicrobial activity (Nowak-Thompson et al., 2003). The genes responsible for HPR production, the Dar genes, which were previously identified in P. chlororaphis subsp. aurantiaca BL915 (Nowak-Thompson et al., 2003), have also been demonstrated to be present in strain PcPCL1606 (Calderón et al., 2013, 2014a). In PcPCL1606, the Dar genes are located in a cluster containing three biosynthetic genes (darA, darB, and darC), followed by two transcriptional regulatory genes darS and darR, both belonging to the transcriptional regulatory family AraC/XylS.

In addition, it has been demonstrated that the biosynthetic genes are positively regulated by the DarSR transcriptional regulators, and all the Dar genes are under the control of GacS (Calderón et al., 2014a). In addition, some alkyresorcinols (to which the compound HPR belongs) can be proposed to be possible signal molecules in the genus Photorhabdus (Brameyer et al., 2015).

It was demonstrated that HPR production is involved in the multitrophic avocado root-Rosellinia necatrix-PcPCL1606 interaction. HPR production by PcPCL1606 was shown to play a key role in the persistence and colonization ability on the avocado roots, while non-HPR-producing mutants show lower colonization levels on the avocado roots and on R. necatrix CH53 hyphae (Calderón et al., 2014b). Other recent studies suggested additional roles of the antibiotics produced by a P. chlororaphis strain. Thus, the role of PHZ production by P. chlororaphis in biofilm formation has been extensively studied, confirming its involvement in biofilm formation but reducing its role during biocontrol (Maddula et al., 2008; Selin et al., 2010, 2012).

The objective of this study was to elucidate the role of HPR production in the ability of PcPCL1606 to form biofilms and determine the presence of this antifungal antibiotic affects the development and biofilm structures.

**MATERIALS AND METHODS**

**Bacterial Strains and Culture Conditions**

The wild-type strain PcPCL1606 and the different derivative strains used in this study (Table 1) were grown on tryptone-peptone-glycerol (TPG) medium (Calderón et al., 2013). The bacterial strains were stored at –80°C in LB with 10% dimethyl sulfoxide. The media was supplemented with kanamycin (50 μg/ml) and gentamicin (30 μg/ml), when necessary.

A collection of insertional mutants in each of the Dar genes was available from earlier studies (Calderón et al., 2013; Table 1). However, to use genetically clean mutants of the biosynthetic Dar genes, a deletional mutant in the Dar biosynthetic gene (ΔDarB) was constructed as previously described (Matas et al., 2014). Briefly, upstream and downstream fragments of the Dar region to be deleted were cloned into the pGEM-T Easy Vector® as described by Matas et al. (2014). Later, the npII kanamycin resistance gene obtained from pGEM-T-KmFRT-HindIII was introduced into the plasmid, yielding pGEM-T-ΔDarB-Km. Finally, each plasmid was transformed by electroporation into PcPCL1606 for marker exchange mutagenesis. Screening and verification of the mutants was conducted as previously described (Matas et al., 2014).

For visualization using confocal laser scanning microscopy (CLSM), the bacterial strains listed in Table 1 were transformed with the plasmid pBAH8 (Huber et al., 2002), which expresses the gfp gene (green) as previously described (Calderón et al., 2014b). The phenotypic characteristics of each transformed bacterial strain were analyzed (growth on minimal and rich medium, antagonism, and HPR production) using the procedures described below.

**HPR Production and Antagonism**

To check the proper phenotypes, HPR production and fungal antagonism were tested for the wild-type and different derivative strains as previously described (Cazorla et al., 2006). Briefly, for HPR production, cell-free supernatants from 5-day-old liquid KB cultures of the test strain were extracted using chloroform/methanol (2:1, v/v). The organic fractions were dried and resuspended in 100 μL acetonitrile. Fifty microliters of the extractions were fractionated by thin layer chromatography (TLC) using silica RP-18F254s TLC plates (Merck AG, Germany) in chloroform/acetone (9:1, v/v). After drying, the chromatogram was visualized under UV light at 254 nm, and the Rf values were calculated. Antibiotic production was also determined by spraying these TLC plates with diazotized sulfanilic acid and watching for a characteristic color change (Whistler et al., 2000). Spots with a Rf value of approximately 0.9–0.95 that were brown to dark green in color were considered positive for HPR. The wild-type strain PcPCL1606 was used as a reference for antibiotic production (Cazorla et al., 2006). The HPR production was quantified as previously described (Calderón et al., 2014a).

The antagonistic activity was tested in vitro as previously described (Geels and Schippers, 1983; Cazorla et al., 2006) using the fungal pathogens R. necatrix CH53 and Fusarium oxysporum.
f. sp. radicis-lycopersici ZUM2407 (Table 1). Experiments were performed on KB and PDA plates as follows: a 0.6-cm-diameter mycelial disk from a 5-day-old fungal culture was placed in the center of a Petri dish at 24°C, and the bacterial strain was inoculated at a distance of approximately 3 cm from the fungus. The antagonistic strain PcPCL1606 was used as a control. The fungal inhibition was recorded after 5 days of growth.

**Swimming Motility Assay**

For the swimming analysis, the bacteria were inoculated with a sterile toothpick into the center of a 0.3% agar plate with TPG diluted 1/20 in MilliQ water similarly to a method previously described (Dekkers et al., 1998). The plates were analyzed after 24 h of incubation at 25°C. Measurements of the motility circle radius enabled the calculation of the motility area. Five independent experiments were performed.

**Biofilm Adhesion Assay**

To test the early phases of biofilm formation, adhesion to an abiotic surface was performed. Biofilm formation was assayed by the ability of the cells to adhere to the wells of 96-well microtiter dishes comprised of polyvinylchloride plastic (PVC, Tissue culture plate 96-well, round bottom suspension cells, Sarstedt) as previously described with modifications (O’Toole and Kolter, 1998). An exponential culture of the bacterial strains (TPG media, 10 h at 25°C) was adjusted to an O.D. of 0.08 (10^8 cfu/mL) with sterile TPG medium. The different test bacterial strains were distributed (100 µL of bacterial suspension) into each well (at least six wells inoculated per strain). After inoculation, the plates were incubated at 25°C for 3 days without movement. A total of 120 µL of a 1% crystal violet solution was added to each well to stain the cells, and the plates were incubated at room temperature for 30 min and rinsed thoroughly and repeatedly with water. Finally, 120 µL of 50% methanol was added to each well to solubilize the crystal violet at room temperature for 20 min. The amount of crystal violet present in each well was determined by absorbance at 595 nm to quantify the biofilm.

**Interface Pellicle Air–Liquid Formation**

Pellicle biofilm formation on the surface of a liquid culture was analyzed using liquid TPG media (Calderón et al., 2013). For pre-cultures, each test strain was exponentially grown in TPG medium during 10 h. The bacterial concentration was adjusted with TPG medium to an O.D. of 0.8 (10^8 cfu/mL) with TPG media, and 10 µL of the adjusted suspension was used to inoculate 1 mL of TPG in 24-well plates to a final O.D. of 0.08. The inoculated plates were incubated without agitation at 25°C for 6 days, and the presence of the characteristic pellicle was then reported (Friedman and Kolter, 2004).

For chemical complementation assays, PcPCL1606 cultures were grown in a 200 mL TPG medium for 28 h at 25°C under orbital shaking (150 rpm), and the production and accumulation of HPR were confirmed as previously described (Calderón et al., 2014a). The cultures were centrifuged, sterilized by filtration (0.22-µm filter), and the cell-free supernatant was collected.

### Table 1: Bacterial strains used in this study.

| Strain                     | Relevant characteristics | Reference                          |
|----------------------------|--------------------------|------------------------------------|
| **Bacterial strains**      |                          |                                    |
| *Pseudomonas chlororaphis* |                          |                                    |
| PCL1606                   | Wild-type, isolated from Spanish avocado rhizosphere, HPR ++++ | Cazorla et al., 2006               |
| darA-                     | PCL1606 derivative insertional mutant in darA gene, HPR -, Km' | Calderón et al., 2013              |
| darB-                     | PCL1606 derivative insertional mutant in darB gene, HPR -, Km' | Calderón et al., 2013              |
| darC-                     | PCL1606 derivative insertional mutant in darC gene, HPR + , Km' | Calderón et al., 2013              |
| darS-                     | PCL1606 derivative insertional mutant in darS gene, HPR +++, Km’ | Calderón et al., 2013              |
| darR-                     | PCL1606 derivative insertional mutant in darR gene, HPR +++, Km’ | Calderón et al., 2013              |
| GacS-                     | PCL1606 derivative insertional mutant in gacS gene, HPR -, Km' | Cazorla et al., 2006               |
| ΔdarB                     | PCL1606 derivative deletional mutant in darB gene, HPR - | This study                          |
| ComB                      | ΔdarB transformed with the plasmid pCOMB, HPR +++, Gm' and Km' | This study                          |

**Fungal strains**

| **Fusarium oxysporum f. sp. radicis-lycopersici** | | |
| ZUM2407 | Causes crown and foot rot of tomato | IPO-DLO Wageningen, The Netherlands |

**Plasmids**

| Name                        | Description                                                                 | Reference       |
|-----------------------------|-----------------------------------------------------------------------------|-----------------|
| pCOMB                       | darB gene cloned into pBRR1MCS-5 used for complementing mutation on strain ΔdarB, Gm' | Calderón et al., 2013 |
| pBAH                        | pBRR1MCS-5-containing PA1/04/03-gfp mut3-To-T1; Gm'                       | Huber et al., 2002 |
| pOEM®-T Easy Vector         | Linearized vector with single 3'-terminal thymidine at both ends            | Promega         |

*HPR production level of the wild-type strain *P. chlororaphis* PCL1606; +++ = HPR production; ++ = 1/4 of HPR production; + = 1/8 of HPR production; = no production (Calderón et al., 2014a). Antibiotic resistance: Km' = Kanamycin, Gm' = Gentamycin. IPO-DLO: Institute for Plant Protection – Agriculture Research Department.
One hundred milliliters of PcPCL1606 cell-free supernatant was added to 100 mL of the TPG medium as previously described (Mori et al., 2017). The mixture was distributed in 24-well plates. Finally, 10 µL of a cell suspension (0.8 at O.D.600nm) of ΔdarB, COMB, or PcPCL1606, was inoculated in the plate wells containing the cell-free supernatant/TPG mixture, and interface pellicle air–liquid formation assays were conducted. This same experiment was repeated using the wild-type PcPCL1606 and ΔdarB GFP-tagged to visualize the pellicle structure using scanning confocal laser microscopy. After 6 days of growth, the pellicle was separated from the well and carefully removed for microscopic visualization. As a negative control, this experiment was also performed using the supernatant of the ΔdarB mutant. To avoid the possibility of other compounds that may play a role in regulating biofilm formation, the culture supernatant of the ΔdarB mutant on biofilm formation in the wild-type strain was tested.

The same experiment of chemical complementation was performed using gfp-tagged strains (wild-type and ΔdarB-derivative strain), in order to visualize the pellicle using CLSM. The obtained pellicle was carefully removed and mounted on a glass slide for observation. When no pellicle was observed, the culture was sampled to confirm the presence of planktonic gfp-tagged strains. All the experiments were performed in biological triplicate.

Colony Morphology Assay
The wild-type strain and the derivative mutants were plated on 1% TPG agar plates to obtain single isolated colonies to observe the colony morphology and polysaccharide production, measured as Congo red binding ability of the colony. One percent TPG agar medium was supplemented with 40 µg/mL Congo red (CR; Sigma) and 20 µg/mL Coomassie brilliant blue (CB; Sigma). Ten microliters of cell suspension with an O.D.600nm of 0.8 (10⁶ cfu/mL) was spotted onto the different agar plates and grown at 25°C during 5 days (Ramos et al., 2010). Images were captured using a fluorescence stereomicroscope AZ-100. All the assays were repeated three times with independent bacterial cultures.

To confirm the chemical complementation, cells-free supernatants were obtained as described above. To prepare the complementation plates, 100 mL of cell-free supernatant was added to 100 mL of the TPG agar media (with a double agar concentration), supplemented with CR and CB adjusted to a final concentration of 40 and 20 µg/mL, respectively. The wild-type strain PcPCL1606 and the ΔdarB mutant were assayed. Ten microliters of the test cultures (0.8 at O.D.600nm) was inoculated in the TPG and in complementation plates. The characteristics of the bacterial colonies were observed after 5 days of growth. All the experiments were performed in triplicate.

Biofilm Architecture
To determine if the architecture of a biofilm was finally altered at the early stages of maturation, a flow experiment was performed. Biofilms were grown in flow cells supplied with AB minimal media (Clark and Maaløe, 1967). The flow system was assembled and prepared as described previously (Christensen et al., 1999). Briefly, the flow channels were inoculated with different GFP-tagged P. chlororaphis cultures grown at a low cell density (OD600nm) in AB minimal media supplemented with 1 mM citrate as the carbon source. The medium flow was maintained at a constant rate of 0.2 mm/s using a Watson-Marlow 205S peristaltic pump. The incubation temperature was 25°C. Microscopic inspection and image acquisition were performed using a confocal laser scanning microscope (DM5500Q; Leica) equipped with a 40/1.3 or a 63/1.4 oil objective. The images captured were analyzed with the Leica Application Suite (Mannheim, Germany) and the Imaris software package (Bitplane, Switzerland). Images were prepared for publication using CorelDraw (Corel Corporation) and PowerPoint (Microsoft) software.

Statistical Methods
The data were statistically analyzed using an analysis of variance (Sokal and Rohlf, 1986), followed by Fisher’s least significant difference test (P = 0.05) using the SPSS 24 software (SPSS Inc., Chicago, IL, United States). All the experiments were performed at least three times independently.

RESULTS
Phenotypic Characterization of the Mutant Derivatives
The analysis of the derivative strains used in this study was performed using molecular and phenotypical assays, such as antagonism and the production of HPR by the TLC test. Our results showed that the darA-, ΔdarB, darB-, and gacS- mutants did not produce HPR. HPR production was detected in the darC-, darS-, and darR- mutants, but at a lower amount when compared with the wild-type strain. The complemented strain displayed HPR production at levels similar to that observed for the wild-type strain (Table 1). Similar results were obtained when antagonism was analyzed, with an absence of antagonistic activity for darA-, ΔdarB, darB-, and gacS-, moderate antagonism for darC-, darS-, and darR-, and strong antagonism for the complemented strain (data not shown).

Swimming Motility Assay
The swimming motility was quantified by measuring the swimming area in the TPG medium agar plate (Figure 1A). All the non-HPR-producing mutants used in this study (Table 1) exhibited increased motility when compared to the wild-type strain. The mutants showed a swimming area of 7.46–13.35 cm², while the wild-type strain covered an area of 5.25 cm² (Figure 1B).

The introduction of a plasmid containing the darB gene into the ΔdarB mutant (COMB strain) restored the HPR production and decreased motility (5.90 cm²), reaching motility levels similar to those displayed by the wild-type strain when compared to those motility values of the ΔdarB mutant (12.2 cm²) affected in HPR production.
The mutant in GacS displayed the highest motility of all the strains assayed.

**Adhesion Assays to PVC Microtiter Wells**

Mutants with defects in the biosynthetic genes (darABC), and the transcriptional regulators darS and GacS were found to be impaired in their adhesion to the PVC surfaces (Figure 2). Impairment in the HPR production strains, such as the darC- or darS- mutants, produces lower amounts of HPR than the wild-type strain, and displayed significantly lower adhesion levels. Only the darR- mutant (producing half of the normal amount of HPR) and the COMB strain displayed the same adhesion levels as the wild-type strain.
In those experiments, the derivative mutant in GacS showed almost no adhesion to PVC with the lower values of absorbance at O.D.$_{595\text{nm}}$.

**Interface Liquid–Air Pellicle Formation**

All the HPR producing strains (wild-type PcPCL1606 strain and derivatives $\text{darC}-$, $\text{darS}-$, $\text{darR}-$, and COMB) formed pellicles in the liquid–air interface, while no pellicle formation was observed with the non-HPR producing strains, such as $\text{gacS}-$ and the $\text{darA}-$ and $\text{ΔdarB}$ mutants impaired in HPR production (Figure 3A). When the chemical complementation was assayed, the derivative strain $\text{ΔdarB}$ (non-pellicle former strain on TPG) was able to form pellicle when growing on TPG media amended with sterile spent TPG media of a PcPCL1606 culture that had been growing previously (Figure 3B), but no pellicle was observed when grown on TPG amended with sterile supernatants from a $\text{ΔdarB}$ culture. The wild-type strain forms the typical pellicle when growing on the culture supernatant of the $\text{ΔdarB}$ mutant, excluding the presence of other compounds that could play a role in biofilm formation.

Because chemical complementation experiments demonstrated the presence of pellicle when HPR production has taken place in the media, a similar assay was performed using gfp-tagged strains. The pellicle formed by the wild-type

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**FIGURE 3 | Interphase air–liquid pellicle formation.** (A) Pellicle formation of $P$. chlororaphis PCL1606 and its derivative mutants. Strains were inoculated (final O.D.$_{600\text{nm}}$ of 0.08) and grown on TPG media in 24-well plates without agitation at 25°C for 6 days. (B) A similar experiment but using cell-free supernatants of PcPCL1606 mixed with TPG as growth media in order to demonstrate the pellicle recovery in the liquid medium by the defective mutant $\text{ΔdarB}$. (C) The same experiment as B, using GFP-tagged strains to visualize the pellicle using confocal scanning laser microscopy. Size of the bars in the microwell pictures: 5 mm; bars of confocal laser scanning microscopy: 5 µm.
FIGURE 4 | Colony morphology assay and analysis of Congo red binding. *P. chlororaphis* PCL1606 and its derivative mutants when were grown on (A) 1% TPG agar plates and (B) 1% TPG agar plates containing 40 µg/mL Congo red and 20 µg/mL Coomassie brilliant blue. (C) a similar experiment but using TPG plate media amended with cell-free supernatants of PcPCL1606 (1:1) to demonstrate the restoration in colony morphology of the defective mutant ΔdarB. The plates were incubated at 25°C for 5 days. Size of the bar is 5 mm.
strain PcPCL1606-GFP (HPR producing strain) and ΔdarB-GFP (non-HPR producing strain) showed no differences when compared under CLSM with fluorescent cells tightly grouped when PcPCL1606-GFP is grown in TPG and when ΔdarB-GFP is grown in the liquid media amended with the cell-free supernatants from PcPCL1606 (Figure 3C). The mutant ΔdarB-GFP, which is defective for HPR production, showed no biofilm and only dispersed viable planktonic cells were observed.

Colony Morphology on Agar-Congo Red
In the colony morphology assay, cell suspensions are spotted on 1% TPG agar-solidified media supplemented with CR and CB (Friedman and Kolter, 2004; Madsen et al., 2015). In this media, the wild-type strain and the HPR producing strains are characterized by a wrinkled surface and a strong red color due to the Congo red binding, indicating the polysaccharide production. However, the derivative mutants affected in HPR production (darA- and ΔdarB) and in the transcriptional regulator GacS (gacS-) showed a smooth surface (on TPG media) and a lack of red colors on TPG media with CR and CB (Figures 4A,B).

In a similar assay, but using a solid agar plate supplemented with cell-free PcPCL1606 supernatant (50% v/v), the ΔdarB derivative mutant was able to accumulate some red color in the colony, partially complementing the mutant phenotype and exhibiting more similar to the wild-type colony (Figure 4C).

Biofilm Architecture
Biofilm architecture was analyzed at the initial stage of biofilm maturation using flow-through flow cells chambers (Figure 5). We found differences between the wild-type and the different derivatives strains tested in this experiment, in which the primary difference is related to HPR production. As shown in Figure 5, we observed that the wild-type strain, which produces a high amount of HPR (Calderón et al., 2014a), was dominated by large microcolonies forming the characteristics of mushroom-shaped structures and characterized by a completely covered surface, while the transcriptional regulator (darSR), which produced half of the HPR in comparison to the wild-type strain (Calderón et al., 2014a), only a few cells colonized the void space between the microcolonies. The primary difference with respect to the wild-type strain was observed with the non-HPR-producing mutants ΔdarB and gacS-. After 3 days of growth, the ΔdarB strain had formed flat and unstructured colonies with a low volume/area biofilm ratio, and with the gacS- mutant no colonies, but filamentous cells were observed.

DISCUSSION
Antibiotics are widely studied in plant beneficial Pseudomonas species, since they are considered to be essential for the biocontrol ability against different soilborne phytopathogenic fungi (Haas and Keel, 2003; Raaijmakers and Mazzola, 2012). However, there are also additional roles for the bacterial antibiotics, which can provide additional ways of participation in the bacterial interaction with the pathogen, such as serving as signals molecules regulating crucial phenotypes (Morohoshi et al., 2013; Brameyer et al., 2015).

Plant-associated pseudomonads have often been studied as model microorganisms for the microbial interaction with plant roots (Lugtenberg and Dekkers, 1999; Lugtenberg et al., 2001). In addition, some colonizing rhizobacteria have been found to form dense biofilm-like structures that occupy the rhizoplane, especially at the junctions between the epidermal root cells (Chin-A-Woeng et al., 1997; Normander et al., 1999; Cassidy et al., 2000; Ramos et al., 2000; Villacieros et al., 2003). However, biofilm formation is considered to be a complex process, and different strains could develop different biofilm architectures,
depending also depending on the environmental conditions (Petrova and Sauer, 2012).

PcPCL1606 synthesizes HPR in addition to other antifungal compounds. HPR is crucial in the biocontrol ability of this strain against different soilborne phytopathogenic fungi (Cazorla et al., 2006; Calderón et al., 2013). However, the antifungal activity is not the only role for HPR in PcPCL1606 biology. Previous results showed that non-HPR producing mutants of PcPCL1606 displayed lower root colonization levels (Calderón et al., 2014b), suggesting a possible role of HPR in biofilm formation, similar to what has been reported for the antifungal compounds PHZs. PHZs can participate in the interaction with the pathogen, participating in cell-to-cell communication as signal molecules with key roles in biofilm formation (Zhang and Pierson, 2001; Dietrich et al., 2006; Morohoshi et al., 2013). This phenotype has been explored in other studies, and it has been proposed that antibiotics can play multiple, concentration dependent roles. At lower concentrations, antibiotics could act as signaling molecules capable of modulating gene expression, while at higher concentrations, they function as inhibitors (Davies et al., 2006; Fajardo and Martínez, 2008).

Swimming motility is involved in biofilm formation (Watnick and Kolter, 2000). The swimming ability of PcPCL1606 could play an essential role during the first stage of biofilm formation through cell/cell and cell/surface interaction, as has been previously described in other *Pseudomonas* spp. (Li et al., 2015). Thus, swimming motility in the *dar* and *gacS*- derivative mutants of PcPCL1606 were analyzed. The *gacS*- mutant showed the expected phenotype of hypermotility, also observed for the *gacS*-derivatives mutant of *Pseudomonas fluorescens* F113, resulting in an increasing swimming motility (Martínez-Granero et al., 2006). Curiously, HPR-defective mutants also showed an increase in swimming motility, consistent with the hypothesis that HPR is involved in biofilm formation, since mobile and sessile are opposite biological states (Petrova and Sauer, 2012).

Thus, the opposite phenotype should be expected for these strains when the adhesion assay is performed. Adhesion is considered to be one of the first steps in biofilm development (O’Toole and Kolter, 1998). When the *dar* and *gacS* mutants were tested, non-HPR producers showed a reduced adhesion to the PVC surfaces with the exception of the *darR* mutant (Figure 2). Since this mutant is impaired in a regulatory gene, additional biological processes could be affected that may explain its similar phenotype to the wild-type strain (Calderón et al., 2014a). Similar results have also been observed for the production of antifungal PHZ production in other strains of *P. chlororaphis* and *P. aeruginosa* (Maddula et al., 2008; Selin et al., 2010).

When colony morphology analyses were performed, defective strains in HPR production and a defective mutant in GacS showed an extracellular component deficit. The lack of red dye in the bacterial colony when growing in the presence of CR revealed the lack of exopolysaccharides that usually dye red with CR (Friedman and Kolter, 2004). This lack of polysaccharides usually leads to inconsistency of the extracellular matrix (Madsen et al., 2015). This fact is reflected by the absence of pellicle formation in the air–liquid interphase. This pellicle is produced by the cultures of all HPR-producing strains. The non-pellicle former Δ*darB* mutant restores biofilm formation when it is genetically and chemically complemented at a level that is undistinguishable from the wild-type strain.

Finally, to unravel if the lack of HPR production can affect the development of a more mature biofilm, flow chamber experiments were conducted. When the biofilm architecture was observed on the wild-type strain and COMB derivative strain, the formation of a large and distinctive mushroom-like cluster with a high biofilm density around the colonies was observed. However, mutants unable to produce HPR, such as Δ*darB* or GacS-, showed the presence of a slightly disorganized layer of bacteria, probably due to a modification of the biofilm matrix, indicated by the absence of exopolysaccharide production on the CR plates, similar to the results previously observed for *Pseudomonas aeruginosa* (Colvin et al., 2011). The intermediate stages of HPR production, such as the *dar*- and *darR*- mutants, give rise to intermediate structures of biofilm, observing a direct relationship between HPR quantity and biofilm consistency. Similar results were described previously in *P. chlororaphis* strain 30-84, in which the biofilm formation is dependent on the type of PHZs produced. The production of only PCA or 2-OH-PCA resulted in a substantially thicker mushroom-shaped cluster with a higher cell density than the wild-type strain (Maddula et al., 2008).

Therefore, it is straightforward to conclude that HPR is a molecule that has a role in biofilm organization in PcPCL1606, and its lack resembles to the phenotype resulting from the inactivation of the regulatory system GacS-GacA (Choi et al., 2007). These facts reinforce the previously reported relation of the biosynthesis of HPR and its regulation by the *darSR* genes also regulated at a higher level by the two-component regulatory system gacS-gacA (Calderón et al., 2014a). The results presented in this study indicate the direct involvement of HPR in biofilm development by PcPCL1606 for the first time. In addition, the requirement for HPR has also been shown to be directly involved in the production of polysaccharide compounds in the biofilm matrix. Some *P. chlororaphis* strains have shown evidences that they of harbor an additional quorum-sensing system sharing genes with the PHZ regulation genes *phzI* and *phzR* (Zhang and Pierson, 2001), also required for biofilm formation (Maddula et al., 2008). However, PcPCL1606 is an atypical *P. chlororaphis*, and it does not produce any type of PHZs. HPR is the main antifungal molecule produced by PcPCL1606.

Finally, the fact that the dialkylresorcinols and cyclohexanediones could be a cell-to-cell communication signal molecule instead of AHL (Brameyer et al., 2015) strengthen the hypothesis that HPR acts as a regulatory signal for phenotypes, such as biofilm formation via exopolysaccharide production.

**AUTHOR CONTRIBUTIONS**

CC, LE, and FC designed the experiments. CC, ST, EA, ZH-P, and GC-O performed the experiments. CC, LE, and FC analyzed
the results and wrote the manuscript. All the authors read and approved the final manuscript.

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