Phenazine and 1-Undecene Producing *Pseudomonas chlororaphis* subsp. *aurantiaca* Strain KNU17Pc1 for Growth Promotion and Disease Suppression in Korean Maize Cultivars

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In this study, strain KNU17Pc1 was tested for its antifungal activity against *Rhizoctonia solani* AG-1(IA), which causes banded leaf and sheath blight (BLSB) of maize. KNU17Pc1 was tested further for its broad-spectrum antifungal activity and in vitro plant growth promoting (PGP) traits. In addition, the in vivo effects of KNU17Pc1 on reduction of BLSB severity and seedling growth promotion of two maize cultivars under greenhouse conditions were investigated. On the basis of multilocus sequence analysis (MLSA), KNU17Pc1 was confirmed as *P. chlororaphis* subsp. *aurantiaca*. The study revealed that KNU17Pc1 had strong in vitro antifungal activity and was effective toward all in vitro PGP traits except phosphate solubilization. In this study, for the first time, a strain of *P. chlororaphis* against *Colletotrichum dematium*, *Colletotrichum gloeosporioides*, *Fusarium oxysporum* f.sp. *melonis*, *Fusarium subglutinans* and *Stemphylium lycopersici* has been reported. Further biochemical studies showed that KNU17Pc1 was able to produce both types of phenazine derivatives, PCA and 2-OH-PCA. In addition, solid phase microextraction-gas chromatography–mass spectrometry (SPME-GC-MS) analysis identified 13 volatile organic compounds (VOCs) in the TSB culture of KNU17Pc1, 1-undecene being the most abundant volatile. Moreover, for the first time, Octamethylcyclotetrasiloxan (D4), dimethyl disulfide, 2-methyl-1,3-butadiene and 1-undecene were detected in *P. chlororaphis*. Furthermore, this study reported for the first time the effectiveness of *P. chlororaphis* to control BLSB of maize. Hence, further studies are necessary to test the effectiveness of KNU17Pc1 under different environmental conditions so that it can be exploited further for biocontrol and plant growth promotion.

**Keywords:** Biocontrol, PGPR, *Pseudomonas chlororaphis*, SPME-GC-MS, *Zea mays*

**Introduction**

Maize (*Zea mays*) is one of the most economically important food and feed crops on the planet and it is also widely used in industrial products, including starch, biofuels, beverages and oil [1]. However, several factors (both biotic and abiotic) limit the yield of maize worldwide [2, 3].

*Rhizoctonia solani* Kühn, [teleomorph *Thanatephorus cucumeris* (Frank) Donk], is one of the major soil-borne pathogens of maize and it causes banded leaf and sheath blight (BLSB) disease. This plant pathogen can cause severe economic loss with up to 100% crop failure [4]. Moreover, the nature of the pathogen to infect a broad range of hosts and to survive as sclerotia under difficult environmental conditions makes the pathogen difficult to control [4, 5].

Biological control continues to grow in popularity and it is becoming a good alternative in reducing the use of chemicals in sustainable agriculture [6]. Plant-associated *Pseudomonas* spp. have been reported to have plant growth promotion, biological control activities and great potential in agro-biotechnological applications [7, 8]. Therefore, this study aims to investigate the in vitro antagonistic activities of strain KNU17Pc1, which is isolated from maize rhizosphere, against *R. solani* AG-1(IA) as well as other...
economically important plant fungal pathogens. The effect of volatile organic compound (VOC) emissions from strain KNU17Pc1 on the selected plant fungal pathogens was studied and the VOC profiles of strain KNU17Pc1 were further examined to investigate the chemical properties of the VOCs responsible for the observed biological activities. Furthermore, under greenhouse conditions, this study was aimed at evaluating strain KNU17Pc1 for its in vivo PGP activities and for its potential to control BLSB on two different Korean maize cultivars (Mibeak-2 and Miheugchal).

**Materials and Methods**

**Bacterial and Fungal Materials**

In this study, bacterial isolates from the rhizosphere soil of maize were isolated as described by Egamberdieva et al. [9] in 2017 at Gangwon Province (37°.86'98.83''N, 127°.75'07.82''E), South Korea. Among morphologically different bacteria strains, strain KNU17Pc1 showing potent in vitro activity against *Rhizoctonia solani* AG-1(IA) was chosen. Strain KNU17Pc1 was stored at −80°C in tryptic soy broth (TSB) with 20% (v/v) glycerol for long-term use. Fungal plant pathogens used in the present study were acquired from Korean Agricultural Culture Collection (KACC), Korea. The fungal plant pathogens were: *Alternaria alternata* (KACC43921), *Colletotrichum dematium* (KACC40013), *Colletotrichum gloeosporioides* (KACC40003), *Fusarium graminearum* (KACC47499), *Fusarium oxysporum* f.sp. *melonis* (KACC47669), *Phytophthora capsici* (KACC40157), *Rhizoctonia solani* AG-1(IA) (KACC40102) and *Stemphylium lycopersici* (KACC40967). *Fusarium subglutinans* was provided by Prof. Kim Kyoung Su, Kangwon National University, South Korea.

**In Vitro Anti fungal Activity Assay**

The antifungal potential of strain KNU17Pc1 against *R. solani* AG-1(IA) was investigated using dual-culture technique on PDA medium. The Petri plates were incubated at 28 ± 2°C for 7 days and the diameter of the clear zone around the bacterial colony was measured.

**In Vitro Assay for Antifungal Traits**

Siderophore production potential of the strain KNU17Pc1 was determined following the method of Schwyn and Neilands [11]. The potential of KNU17Pc1 for protease activity was determined by spot inoculating overnight TSB culture of KNU17Pc1 on skim milk agar and a clear zone around the bacterial colony was considered as positive [12]. The method of Ghodsalaví et al. [13] was employed to determine the lipase activity of KNU17Pc1.

**In Vitro Assay for Plant Growth Promoting (PGP) Attributes**

Strain KNU17Pc1 was determined for its zinc solubilization potential on PKV medium amended with an insoluble zinc source (ZnO, 1.244 g/l) equivalent to 0.1% zinc. The phosphate solubilizing ability of strain KNU17Pc1 was evaluated on NBRIP (National Botanical Research Institute’s phosphate growth) medium supplemented with tricalcium phosphate as a sole phosphate source. The Petri plates were incubated at 28 ± 2°C for 7 days and the diameter of the clear zone around the bacterial colony was measured.

The potential of strain KNU17Pc1 to produce ammonia was determined following the method of Cappuccino and Sherman [14]. The amount of ammonia formed was determined spectrophotometrically (UV–1800, Shimadzu Corporation, Japan) using the standard curve of ammonium sulphate ranging from 0–10 μmol/ml. The method of Gordon and Weber [15] was employed to determine the potential of strain KNU17Pc1 to produce indole acetic acid (IAA). The amount of IAA concentration in the culture medium was measured using an IAA standard curve prepared by diluting pure IAA (Sigma–Aldrich, USA) in Luria-Bertani (LB) broth (Tryptone 10 g, yeast extract 5 g, and NaCl 10 g in 1 l of water) at different concentrations in the range of 0.05 to 2 mg/ml.

**Quantification of Phenazine**

Phenazines viz., phenazine-1-carboxylic acid (PCA) and 2-hydroxy-phenazine-1-carboxylic acid (2-OH-PCA) were extracted...
from strain KNU17Pc1 grown in seven different media. The media were: AB minimal medium [16] amended with 2% Casamino acids (Difco, Franklin Lakes, USA) (AB+CAA), King’s B medium (KB) [17], LB, minimal salts medium (MSM) [18], nutrient broth (Difco Laboratories, USA), pigment production medium (PPMD) [19] and TSB. Strain KNU17Pc1 was incubated at 28 ± 1°C with agitation (180 rpm) and the bacterial culture of each medium was collected at the same O.D. The method of Maddula et al. [20] was employed to extract phenazines from KNU17Pc1. Briefly, each culture (5 ml) was centrifuged (2,600 × g for 15 min), and the cell-free supernatant was acidified (pH <2) with a concentrated HCl. An equal volume of benzene (v/v) was added and total phenazines were extracted for 1 h. The mixture was centrifuged and the benzene phase was separated and evaporated to dryness. The crude extract was resuspended in 0.1 N NaOH and was quantified using UV-visible spectroscopy at 367 nm (PCA) and 484 nm (2-OH-PCA). A 0.1 N NaOH was used as a blank. The relative amounts of PCA and 2-OH-PCA were determined using standard extinction coefficients: εPCA (367 nm) of 3,019 M/cm and ε2-OH-PCA (468 nm) of 7,943 M/cm [20].

Analysis of Volatile Organic Compounds (VOCs)

Volatile organic compounds (VOCs) from TSB liquid medium inoculated with strain KNU17Pc1 were analyzed using solid phase microextraction-gas chromatography-mass spectrometry (SPME-GC-MS) with SPME fiber assembly (CAR/PDMS) (Supelco, Inc., Bellefonte, USA). The strain KNU17Pc1 was grown in TSB liquid medium for 48 h in dark at 28 ± 2°C on a rotary shaker at 180 rpm. To prevent escaping of VOCs, a rubber stopper containing active charcoal was plugged into the test tubes. A non-inoculated TSB liquid medium was served as a control. The GC (7890A gas chromatograph, Agilent Technologies, USA) was programmed at an initial temperature of 60°C for 5 min, and then gradually increased by 5°C/min to a temperature of 325°C held for 50 sec. The GC transfer line was maintained at 280°C, and detector temperature at 250°C. Inlet pressure was 67 kPa; He, 20 ml/min; filament voltage, 70 eV ionization energy; transfer line 280°C. The VOCs were identified by comparing the spectra mass obtained from the KNU17Pc1 sample with those from reference spectra in the Wiley 9th edition spectral libraries and National Institute of Standards and Technology 2014 V2.20 (NIST, USA, http://www.nist.gov).

Greenhouse Experiments

The potential of strain KNU17Pc1 to control BLSB of maize on two commonly grown Korean maize cultivars (Mibaek-2 and Miheugchal) were carried out under greenhouse conditions. The experiment was conducted in 16 cm diameter pots filled with sterilized soil. The soil was collected from the top 30 cm of topsoil in Gangwon Province, South Korea. Three replications were maintained with 10 plants per replication.

Preparation of Fungal and Bacterial Inoculums

The inoculum of R. solani AG-1(IA) was cultured on PDA plates at 28 ± 2°C for 7 days. The soil where the seeds were to be sown was removed and then two colonized PDA plugs (8 mm in diameter) were replaced seven days before planting. To prevent drying of the colonized disks, they were covered with soil [21]. For bacterial inoculum preparation, the strain was grown in TSB liquid medium for 48 h in dark at 28°C on a shaker incubator at 150 xg. The liquid culture was centrifuged at 6,000 xg for 5 min and the harvested bacterial cells were washed several times with a solution of phosphate-buffered saline (PBS; 5 mM K,HPO4, 150 mM NaCl, pH 7.0). Then, the bacterial inoculum was adjusted to a concentration of 10^8 cells/ml^2. Seeds of both cultivars were surface-sterilized in 2% sodium hypochlorite for 2 min, followed by 70% ethanol for 2 min, and finally rinsed several times in sterile distilled water. Maize seeds (10) of each cultivar were soaked separately into the bacterial suspensions (10 ml) and were incubated at 28 ± 2°C for 12 h in a rotary shaker at 150 xg. Subsequently, seeds of each cultivar were air-dried in a laminar air-flow at ambient temperature and one seed per pot was planted. Non-treated plants (neither R. solani AG-1(IA), nor strain KNU17Pc1) served as positive control and plants inoculated only with R. solani AG-1(IA) were served as negative control.

Disease Assessment

The potential of strain KNU17Pc1 on reduction of disease severity (DS) of BLSB of maize was assessed at 30 days after planting and the DS was recorded using a 1-5 scale [22]. For analysis, the scale was converted into a percentage severity index (PSI) [23].

\[ PSI = \frac{\sum \text{of all numerical ratings} \times 100}{\text{Total number of observations} \times \text{maximum score on scale}} \]

Plant Growth Promotion Assessment

The physical plant growth characteristics such as plant height and stem circumference of maize plants were measured. In addition, data regarding total chlorophyll content (SPAD unit) of the CS leaf of maize were measured using chlorophyll-meter SPAD 502 (Konica Minolta, Japan) 30 days after planting (DAP). Maize seedlings were collected at 30 DAP and morphological data viz., dry weights of shoot and root (65°C, 72 h electric oven) and leaf number were taken. Three replications were maintained with 10 plants per replication.

Molecular Characterization

Identification of strain KNU17Pc1. The genomic DNA of strain KNU17Pc1 was isolated using the HiYield Genomic DNA Mini Kit (Real Biotech Corporation, Taiwan) following the manufacturer’s instructions. PCR amplification of the 16S rRNA gene was performed using universal primers 27F (5'-AGAGTTTGATCCTTGTTACGACT-3') and 1492R (5'-TACGGYTACCTTGTTACGACTT-3'). Sequencing was performed at Macrogen Inc. (Korea) using a 3730XL DNA sequencer (Applied BioSystems, USA). Furthermore, Multilocus Sequence Analysis (MLSA) was employed to charac-
Characterization of Multi-Trait Strain KNU17Pc1

Pairwise sequence analysis indicated that of the 16S rRNA nucleotide sequence of strain KNU17Pc1 shared high similarity (99.0–100%) with reference species of *Pseudomonas*. The phylogenetic relationship using the 16S rRNA gene sequence placed the strain KNU17Pc1 within the *P. chlororaphis* cluster in our phylogenetic tree (Fig. S1). More importantly, a multilocus sequence analysis (MLSA) was employed for further identification. The phylogenetic analysis based on the concatenated sequences of seven

**Results**

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**Detection of phzE Gene**

The strain KNU17Pc1 was subjected to PCR targeting the phenazine (phzE) gene, which is involved in antifungal activity [8, 25]. The pair of primers and PCR conditions for phzE were performed as previously described [26] and a Zymoclean Gel DNA Recovery Kit (Zymo Research) was used to gel purify the PCR products. Sequence analysis of the PCR products was performed with their respective designed primers at Macrogen Inc. (Korea) using a 3730XL DNA sequencer (Applied BioSystems).

**Genbank Accession Numbers**

The 16S rRNA gene sequences of the strain KNU17Pc1 have been deposited in the NCBI’s GenBank under accession number MH82498. The sequences of MLSA loci of the strain have been deposited in the GenBank under accession numbers MH568738 (ileS), MH568736 (glnS), MH568737 (gyrB), MH568739 (nuoD), MH568740 (recA), MH568741 (rpoB) and MH568742 (rpoD). The sequence data of the phzE gene of strain KNU17Pc1 was submitted to Genbank and can be found under accession number, MH388419.

**Data Analysis**

The experimental data were subjected to analysis of variance (ANOVA) using SAS software version 9.2 [27]. All experiments were replicated at least three times and the results were expressed as mean ± standard error. Means were separated using Duncan’s Multiple Range Test (DMRT) at \( p \leq 0.05 \).
housekeeping genes revealed that strain KNU17Pc1 belongs to *P. chlororaphis* subsp. *aurantiaca* and it was diverged from the other closely related species of the *P. chlororaphis* cluster (Fig. 1).

In Vitro Antagonism and Sclerotial Germination Assay

In the dual-culture assay, strain KNU17Pc1 was capable to inhibit the growth of *R. solani* AG-1(IA) (Table 1 and Fig. 2). The strain was also able to completely curb the germination of sclerotia of *R. solani* AG-1(IA) in a sclerotial germination test even after a long period of incubation (data not shown). The strain was further tested in a dual culture assay for its activity against 8 economically important phytopathogens. The results showed that the strain had strong antifungal activity against all the tested fungi (Table 1 and Fig. 2). The strain KNU17Pc1 showed the highest growth inhibition effect against *A. alternate* and *C. dematium*. In the volatile metabolite assays (assays on divided Petri plates), strain KNU17Pc1 showed high inhibitory effect against *R. solani* AG-1(IA) and other important plant fungal pathogens (Table 1 and Fig. S2).

**Table 1.** Antifungal activity of *Pseudomonas chlororaphis* strain KNU17Pc1.

<table>
<thead>
<tr>
<th>Target pathogens</th>
<th>Percent inhibition zone (mm) (Mean±SE)</th>
<th>Diffusible</th>
<th>Volatile</th>
</tr>
</thead>
<tbody>
<tr>
<td><em>A. alternate</em></td>
<td>45.4 ± 2.4⁴ 48.7 ± 1.2⁴</td>
<td>45.4 ± 1.3³ 45.8 ± 1.6³</td>
<td></td>
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<tr>
<td><em>C. dematium</em></td>
<td>43.9 ± 1.0⁵ 19.1 ± 0.9⁵</td>
<td>41.0 ± 0.8⁶ 46.8 ± 2.1⁶</td>
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<tr>
<td><em>F. oxysporum f.sp. melonis</em></td>
<td>37.8 ± 1.3⁶ 51.9 ± 0.8⁶</td>
<td>39.1 ± 2.2⁶ 76.2 ± 3⁶</td>
<td></td>
</tr>
<tr>
<td><em>S. lycopersici</em></td>
<td>35.9 ± 0.9⁷ 89.2 ± 4.2⁷</td>
<td>35.5 ± 1.2⁷ 33.9 ± 0.9⁷</td>
<td></td>
</tr>
<tr>
<td><em>F. subglutinans</em></td>
<td>35.1 ± 0.7⁷ 29.6 ± 0.7⁷</td>
<td>35.1 ± 0.7⁷ 29.6 ± 0.7⁷</td>
<td></td>
</tr>
</tbody>
</table>

⁴Values are means of three replications.
³The antifungal activity determined using dual-culture assay.
⁵The antifungal activity determined using volatile metabolite assays (assays on divided Petri plates).

Means followed by the same letter (s) within a column are not significantly different (*p* ≤ 0.05), DMRT.

SEM Analysis

The effect of *P. chlororaphis* strain KNU17Pc1 on the mycelial morphology of *R. solani* AG-1(IA) and the selected plant fungal pathogens were examined under scanning electron microscope by observing the mycelia found along the edges of the inhibitory halo zone. Microscopic examination showed that KNU17Pc1 caused significant changes in the hypha morphology of all the selected plant pathogens compared to the untreated control. The strain caused the hyphae of *R. solani* AG-1(IA) to lyse and deform (Fig. 3). In addition, the strain was capable of causing substantial change in hyphal morphology of *A. alternate*, *F. graminearum* and *F. oxysporum f.sp. melonis* (Fig. 3).

Characterization of In Vitro Antifungal and PGP Traits of KNU17Pc1

Strain KNU17Pc1 was found to be positive for zinc.

![Fig. 2. Antifungal activity of *Pseudomonas chlororaphis* strain KNU17Pc1 against different plant pathogens from dual culture assay (diffusible metabolites): (A) *A. alternate*, (B) *C. dematium*, (C) *C. gloeosporioides*, (D) *F. graminearum*, (E) *F. oxysporum f.sp. melonis*, (F) *F. subglutinans*, (G) *P. capsici*, (H) *R. solani* AG-1(IA), (I) *S. lycopersici.*](image-url)
solubilization, siderophore production and extracellular activities. However, KNU17Pc1 had no phosphate solubilization or amylase activity (data not shown). In the ammonia production test, KNU17Pc1 showed positive results for ammonia production after 72 h of incubation (data not shown). The amount of ammonia produced in peptone water was 9 µmol/ml. The ability of KNU17Pc1 to produce IAA was tested in LB medium amended with L-tryptophan at various concentrations ranging from 0.05 to 2.0 mg/ml. The study showed that KNU17Pc1 produced 11 µg/ml of IAA at higher L-tryptophan concentration (2 mg/ml) after 72 h of incubation (Fig. S3).

Quantification of PCA and 2-OH-PCA

The result of the present study revealed that strain KNU17Pc1 was capable of producing both types of phenazine derivatives, PCA and 2-OH-PCA (Figs. 4A and 4B). PCA and 2-OH-PCA production by KNU17Pc1 was tested when the bacterial culture of each tested media reached the same O.D. The maximum amount of PCA and 2-OH-PCA was recorded in PPMD broth (101.8 µg/ml and 9.5 µg/ml, respectively) followed by TSB broth (87.4 µg/ml and 5.6 µg/ml, respectively). On the other hand, a lower amount of PCA and 2-OH-PCA was recorded in King’s medium B (5.8 µg/ml and 0.5 µg/ml, respectively).

Fig. 3. Scanning electron micrographs of (A and E) A. alternate, (B and F) F. graminearum, (C and G) F. oxysporum f.sp. melonis and (D and H) R. solani AG-1(IA) (top = treated with KNU17Pc1, bottom = control), (scale bars: 10 µm). A. white arrows indicate deformed and/or lysed fungal structures.

Fig. 4. Production of (A) PCA and (B) 2-OH-PCA of strain KNU17Pc1 grown in seven different liquid media.
Detection of Phenazine Biosynthesis (phzE) Gene by PCR Amplification

In this study, we tested if our strain KNU17Pc1 carries phzE, which catalyzes the first step in the biosynthesis of PCA and other phenazines [24, 25]. The results of PCR using specific primers revealed the gene that codes for phzE (phenazine) was successfully amplified from strain KNU17Pc1 with the expected band size of 450 bp (Fig. S4). Furthermore, pairwise sequence analysis showed that nucleotide sequences of phzE (Acc. No. MH388419) from KNU17Pc1 showed high sequence similarities (99%) to the gene of \textit{P. chlororaphis} subsp. \textit{aurantiaca} involved in phenazine (Acc. No. AB794886.1). The deduced amino-acid sequences of phzE from KNU17Pc1 also revealed very high sequence identity (98%) with previously reported phenazine phzE amino-acid sequences of \textit{P. chlororaphis} (Acc. No. BAM94425.1), with only a two-amino-acid residue difference (Fig. 5).

Identification of Volatile Compounds (VOCs) from Strain KNU17Pc1

In this study, VOCs emitted from strain KNU17Pc1 grown in TSB media were analyzed by SPME–GC–MS. The volatile compound profiles produced by the strain KNU17Pc1 were compared with the volatile compounds found in the bacteria in the non-inoculated medium (control). The results revealed that there was a very clear separation between control and strain KNU17Pc1 as

**Fig. 5.** Sequence analysis of phenazine biosynthesis protein (phzE gene) of \textit{P. chlororaphis} strain KNU17Pc1 (KNU17Pc1), \textit{P. chlororaphis} subsp. \textit{aurantiaca} (BAM94425.1), \textit{P. chlororaphis} (CDL74722.1), \textit{P. aeruginosa} (OZO28361.1), \textit{P. fluorescens} (WP_068967162.1), \textit{Streptomyces cyaneofuscatus} (AQN67857.1).

▼ Distinct amino acid pointing to \textit{P. chlororaphis} subsp. \textit{aurantiaca}. *Fully conserved residue. °Highly conserved column. Weakly conserved column. Gaps are shown as dashes.

**Fig. 6.** Chromatographic profiles of VOCs of (A) \textit{P. chlororaphis} strain KNU17Pc1 incubated for 24 h in TSB medium (B) non-inoculated TSB medium.

The VOCs were as follows: (1) 1,4-Bis(trimethylsilyl)benzene, (2) propanoate, (3) 2-methyl-1,3-butanediene, (4) 1-Dimethylisopropylsiloxy-3-methylbut-2-ene, (5) Acetaldehyde methyl hydrazone, (7) N-Methylthiourea, (8) Dimethyl disulfide, (9) 2-Amino-1H-benzimidazole, (11) 1,4-Bis(trimethylsilyl)benzene, (12) 5H-Naptholo[2,3-b]carbazole, (14) Octamethylcyclotetrasiloxane, (15) 1-undecene and (16) 1,1,3,3,5,5,7,7,9,9-decamethylpentasiloxane.
indicated in Fig. 7. A total of 13 different emitted VOCs of strain KNU17Pc1 origin, which were absent in control, were identified (Fig. 6). The 1-undecene having high abundant peak (45.7%) with RT 10.15 was the most abundant volatile in strain KNU17Pc1.

**Greenhouse Experiments**

**Effect of strain KNU17Pc1 on disease severity in maize seedlings.** Strain KNU17Pc1 was able to suppress the severity of BLSB on both Korean maize cultivars (Miheugchal and Mibaek-2). KNU17Pc1 was efficient (64.1% and 69% for Mhengchal and Mibaek-2, respectively) in reducing the disease compared to the control. To the contrary, seedlings inoculated only with *R. solani* AG-1(IA) (negative control) were highly blighted in both cultivars by the artificially inoculated *R. solani* AG-1(IA) (Fig. 7). As expected, non-treated plants (neither the pathogen (*R. solani* AG-1(IA)), nor KNU17Pc1 strain) served as positive control and plants inoculated only with *R. solani* AG-1(IA) served as negative control.

**Plant growth promotion activity of strain KNU17Pc1.**

The results of the greenhouse experiment revealed that KNU17Pc1 had promising results in promoting the growth of both maize cultivars (Figs. 8 and 9). Strain KNU17Pc1 significantly ($p \leq 0.05$) increased shoot and root dry
weight of Mibaek-2 and Miheugchal compared to negative control (plants inoculated only with \( \textit{R. solani AG-1(IA)} \), nor KNU17Pc1 strain) served as positive control and plants inoculated only with \( \textit{R. solani AG-1(IA)} \) served as negative control.

(weight of Mibaek-2 and Miheugchal compared to negative control (plants inoculated only with \( \textit{R. solani AG-1(IA)} \)) (Fig. 8). Regarding the number of leaves, the strain showed insigniﬁcant differences compared to non-bacterized controls in both cultivars, (data not shown). The total chlorophyll content in the un-bacterized negative control of both cultivars was reduced while bacterized plants of both cultivars were signiﬁcantly improved (Fig. 8). Furthermore, in both maize cultivars, seed inoculation with strain KNU17Pc1 led to a signiﬁcant increase in the plant height of the seedlings along with an increase in stem diameter when compared to their respective non-bacterized controls (Fig. 9).

Discussion

Plant-associated \textit{Pseudomonas} spp. have been reported to have great potential for plant growth promotion and plant disease management [7, 8]. Therefore, we isolated a multi-trait, plant growth-promoting rhizobacteria (PGPR), strain KNU17Pc1. Based on 16S RNA and MLSA analysis, the results revealed that KNU17Pc1 belongs to \textit{P. chlororaphis} subsp. \textit{aurantiaca}. In keeping with this finding, Andreani et al. [28] discussed that the MLSA approach has high discrimination power in all the taxa including \textit{Pseudomonas}. Strain KNU17Pc1 suppressed the mycelial growth and germination of sclerotia of \( \textit{R. solani AG-1(IA)} \). Management of BLSB is difficult due to sclerotia formation which can overwinter in very harsh environmental conditions and continue cycling infection [29, 30]. Hence, the potential of the KNU17Pc1 in inhibiting the germination of sclerotia of \( \textit{R. solani AG-1(IA)} \) has huge implications in the sustainable BLSB management of maize. The effect of plant growth-promoting bacteria in inhibiting sclerotial germination of \( \textit{R. solani AG-1(IA)} \) has previously been reported [31, 32], however, the potential of \textit{Pseudomonas chlororaphis} was not reported. In addition, the strain showed strong antifungal activity against all the tested fungi in the in vitro antagonism tests, dual culture and volatile metabolite assays. The zone of inhibition around tested plant pathogens by KNU17Pc1 may be linked to the strain’s ability to produce secondary metabolites like phenazine, ammonia, and proteolytic enzymes [26, 33, 34]. Interestingly, this is the first report of \textit{P. chlororaphis} against \( \textit{C. dematium, C. gloeosporioides, F. oxysporum f.sp. melonis, F. subglutinans and S. lycopersici.} \) \textit{F. subglutinans} is the potential cause of mycotoxin contamination [35] and also causes seedling blight and stalk rot of maize that poses a potential risk to maize production [36]. Furthermore, \textit{C. dematium, C. gloeosporioides, F. oxysporum f.sp. melonis and S. lycopersici} are among the most economically important pathogens of crops [37–39]. This connotes the essentiality for exploitation of KNU17Pc1 for the management of \textit{C. dematium, C. gloeosporioides, F. oxysporum f.sp. melonis, F. subglutinans and S. lycopersici}. Microscopic examination showed that KNU17Pc1 caused hyphae of \( \textit{R. solani AG-1(IA)}, \textit{A. alternata, F. graminearum and F. oxysporum f.sp. melonis} \) to lyse and deform. Similar to the present study, Huang et al. [40] reported abnormal morphological changes in \( \textit{R. solani AG-1(IA)} \) by different biocontrol agents. To the best of our knowledge, the injurious effects of \textit{P. chlororaphis on F. oxysporum f.sp. melonis} have not previously been investigated.

Production of pigments in a medium is an indication of bacteria producing metabolites [8]. The present study revealed the potential of strain KNU17Pc1 to produce both types of phenazine derivatives, PCA and 2-OH-PCA. In

![Fig. 8. Effect of strain KNU17PC1 on the seedling growth performance of two maize cultivars after 30 days grown under greenhouse conditions. (A and B) total chlorophyll content of Mibaek-2 and Miheugchal, (C and D) seedling dry weight of Mibaek-2 and Miheugchal, respectively. Non-inoculated plants (neither the pathogen (\( \textit{R. solani AG-1(IA)} \)), nor KNU17Pc1 strain) served as positive control and plants inoculated only with \( \textit{R. solani AG-1(IA)} \) served as negative control.](image-url)
agreement with this report, previous studies [19, 41–43] reported the potential of *P. chlororaphis* to produce phenazine derivatives PCA and 2-OH-PCA. The maximum PCA yield in PPMD and TSB broth may be attributed to the presence of glucose and soy peptone, respectively. Similar to our report, He et al. [44] reported that glucose and soy peptone as carbon and nitrogen source, respectively were the most important factors for PCA production. The very low amounts of PCA production by *P. chlororaphis* in King’s medium B has also been previously reported by Chin-A-Woeng et al. [45].

The phenazine biosynthesis (phzE) gene, which codes for aminodeoxyisorhaminate synthase, is involved in the biosynthesis of biologically secondary metabolites and plays vital roles in biological control [46, 47]. In this study, nucleotide sequences of phzE (Acc. No. MH388419) and deduced amino-acid sequences of phzE open reading frame (ORF) from KNU17Pc1 showed high sequence similarities (99%) to the phenazine gene (Acc. No. AB79486.1) and phenazine phzE amino-acid sequences (Acc. No. BAM94425.1) of *P. chlororaphis* subsp. *aurantiaca*.

Previous studies [48, 49] reported that 1-undecene, having strong antifungal activity, was the main active compound emitted by *P. fluorescens*. Similarly, in our SPME–GC–MS analysis 1-undecene was the most abundant volatile in *P. chlororaphis* strain KNU17Pc1. Antifungal activity of 1-undecene against *R. solani* has previously been reported [50]. Nevertheless, this is the first report on emission of 1-undecene by *P. chlororaphis*. The Octamethylcyclotetrasiloxan (D4), which is volatile methyl siloxanes, has been reported to have antibacterial and antifungal activity [51] and was found in the present study. For the first time, 2-methyl-1,3-butadiene and dimethyl disulfide (DMDS) were detected in *Pseudomonas chlororaphis*. The 2-methyl-1,3-butadiene is an important biogenic hydrocarbon in reducing reactive oxygen species (ROS) including hydroxyl anions (OH⁻) and hydrogen peroxide (H₂O₂) in plant cells [52, 53]. The beneficial effect of DMDS for plant fungal disease suppression and growth promotion has been previously reported [40, 54, 55].
In the in vitro PGP traits assay, similar to previous reports [56, 57], strain KNU17Pc1 was positive for zinc solubilization and negative for phosphate solubilization. In agreement with our result, the potential of \textit{P. chlororaphis} to produce IAA has been previously reported [9] while other studies [8] noted that \textit{P. chlororaphis} did not produce IAA. In addition, strain KNU17Pc1 produced a high amount (9 \(\mu\)mol/ml) of ammonia in peptone water. Our result complies with previous reports [58] that several \textit{Pseudomonas} species produce ammonia. Previous studies [34, 59] discussed that inorganic volatiles like ammonia have been found to suppress the growth of plant fungal pathogens. In addition, ammonia-producing strains have a key role in the nutritional needs of plants by accumulation of nitrogen and offering ammonia to nearby plants by breaking up complex nitrogenous materials [60].

Similar to the result of our in vitro assay for antifungal traits, production of siderophores, protease and lipase by \textit{Pseudomonas chlororaphis} has been previously reported [8]. However, KNU17Pc1 was amylase negative [8]. Previous studies [41, 61] discussed that production of siderophores offers beneficial microbes a competitive advantage to suppress the rise of plant pathogens and also help the surrounding plants to obtain iron for optimal growth. These hydrolytic enzymes were found to play a key role in fungal cell walls degradation, which indirectly promotes plant growth [45].

The results of greenhouse experiments showed that strain KNU17Pc1 was able to protect both cultivars from BLSB on artificially infested soil with \textit{R. solani AG-1(IA)}. The previous study [62] reported that BLSB was effectively controlled by \textit{P. fluorescens}. Nevertheless, this is the first research report on the efficacy of \textit{P. chlororaphis} in suppressing the disease severity of BLSB on maize. This may be partly due to the strong antifungal activity of \textit{P. chlororaphis} against \textit{R. solani AG-1(IA)} in the soil. Similar to our study, previous reports [63, 64] indicated that treating maize seeds with plant growth-promoting bacteria \textit{Pseudomonas} species promoted plant growth. The beneficial effect of the strain KNU17Pc1 on plant growth promotion of maize cultivars may be attributed to the direct plant growth promoting effect of the strain [65] and may be indirectly due to the antagonistic nature of the strain in influencing the growth of mycelia and germination of sclerotia of \textit{R. solani} [66]. In addition, the significant growth reduction of maize seedlings in the negative control of Mibaek-2 and Miheugchal may be partly due to infection caused by \textit{R. solani} [67]. Therefore, further studies are needed to evaluate the wide-range biocontrol activities of strain KNU17Pc1 against economically important plant fungal pathogens under field conditions.

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**Conflict of Interest**

The authors have no financial conflicts of interest to declare.

**References**


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