Rice phyllosphere bacteria producing antifungal compounds as biological control agents of blast disease

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Abstract. Wiraswati SM, Nawangsih AA, Rusmana I, Wahyudi AT. 2020. Rice phyllosphere bacteria producing antifungal compounds as biological control agents of blast disease. Biodiversitas 21: 1273-1278. In vitro analysis of rice phyllosphere bacteria can be developed as biocontrol agents of blast disease using their antifungal activity. To prove the efficacy of bacterial isolates in reducing the severity of blast disease, a greenhouse experiment was conducted. Furthermore, antifungal compounds produced by potential bacterial isolate were also identified using Thin Layer Chromatography (TLC) and Liquid Chromatography-Mass Spectrometry (LC-MS/MS) analysis. The application of seven rice phyllosphere bacterial isolates significantly reduced the severity of blast disease in rice (var: Ciherang). Surprisingly, the isolate STGG 14 is the most effective with 70.83% of blast disease reduction. Isolate STGG 14 has been identified as Bacillus subtilis subsp. subtilis which is known as various bioactive compounds producer. It was confirmed through identification of antifungal compounds produced by isolate STGG 14. The bioautography test shows that crude bioactive compounds from isolate STGG 14 could inhibit P. oryzae race 173. On the further analysis using LC-MS/MS, four bioactive compounds isolated from isolate STGG 14 were identified as moracin C, psoralen, 4,6-dimethyl-3 (4’-hydroxyphenyl) coumarin and xanthotoxin. All identified compounds except moracin C, belong to the coumarin group. According to the previous studies, among all compounds, psoralen and xanthotoxin displayed antifungal activity against several fungal pathogens.

Keywords: Bacillus, bioactive compounds, bioautography, Pyricularia oryzae, TLC

INTRODUCTION

The phyllosphere is defined as aerial or above-ground parts of plant including stem, leaves, flowers, buds, and fruits, whereas surface area of leaves constitute phylloplane. This niche is considered as hostile environment due to large fluctuations of physical and nutritional conditions (Batool et al. 2016). Nevertheless, phyllosphere environment is inhabited by a large and diverse microorganism including bacteria, fungi, yeast and archaea which have commensal, pathogenic and mutualistic interactions with the host plant. Leaves are the most dominant aerial part which occupied by bacteria as the most abundant resident (Rastogi et al. 2013). Bacterial density in the leaf surface reaching as high as 10^6 - 10^9 cells per gram of plant material (Yadav et al. 2010) and majority of them are commensal (Ali et al. 2012). Phyllosphere bacteria are capable to protect host plants from pathogens through indirect mechanisms such as production of antimicrobial compounds and competition of resources. Antimicrobial compounds will be secreted by phyllosphere bacteria in order to create a competitive environment for inhibiting the growth of plant pathogens (Chaudhary et al. 2017). Bacteria producing antimicrobial compounds especially antifungal properties have been studied and applied to manage plant pathogens. Bacillus, Pseudomonas, and Pantoea are examples of bacteria that potentially produce antifungal compounds against fungal pathogens such as Pythium, Fusarium oxysporum, Rhizoctonia solani (Junaid et al. 2013) and Pyricularia oryzae (Sha et al. 2016). Generally, Bacillus species have been known as lipopeptides compounds producer with fungal cell wall degradation activity (Tserkovniak et al. 2009) such as fengycin and iturin. Both lipopeptides groups are able to inhibit the growth of filamentous fungi through destruction of sterol, phospholipid and oleic acid on the fungal cell membrane (Gond et al. 2015).

Rice fungal pathogen from Ascomycete group, Magnaporthe oryzae (anamorph: P. oryzae) is the most devastating pathogen in rice plant cultivation. This pathogen belongs to the top 10 fungal diseases that endanger universal food security (Dean et al. 2012). Pyricularia oryzae is the causing agent of blast disease in rice that attacks leaves, culms and panicles of rice plants as well as decreases photosynthesis effectivity and grain yield (Ribot et al. 2008). Among all parts of rice plants that are infected by P. oryzae, neck blast is the most destructive form of blast disease (Zhang et al. 2014). This disease spread in more than 85 countries and has caused great yield loss. At least 30% of the rice harvest is lost due to blast disease infection annually. This amount is equal with rice necessary to feed more than 60 million people for 1 year (Sha et al. 2016). To deal with it, the use of bacteria to reduce crop diseases particularly blast disease seems to be the most suitable way to replace synthetic fungicide (Junaid et al. 2013). The application of bacteria producing antifungal compounds can be effective if they are able to colonize the host plant properly. Likewise, in managing
blast disease, the biocontrol agents should be able to survive on the stems and leaves of rice plants. Therefore, the use of phyllosphere bacteria as biocontrol agents is predicted to be a reliable method for managing foliar diseases like rice blast.

In the previous study, we successfully isolated seven rice phyllosphere bacteria with antifungal activity against *P. oryzae* race 173 in vitro analysis. Among the seven bacterial isolates, five isolates were *Bacillus* species (isolates STGG 3, STGG 7, STGG 8 and STGG 14), one isolate was *Enterobacter* species (isolate STGV 8) and the other isolate was *Brachybacterium* species (isolate SKBG 78). These isolates have been proved to have the ability to produce bioactive compounds (Wiraswati et al. 2019). Furthermore, to confirm the efficacy of these isolates in reducing the severity of blast disease in rice plants, a greenhouse experiment was conducted in this study. Effect of crude bioactive compounds toward morphological alteration of *P. oryzae* race 173 was observed under SEM. Besides, thin layer chromatography and LC-MS/MS analysis were also carried out to identify the antifungal compounds produced by phyllosphere bacteria.

**MATERIALS AND METHODS**

**Procedures**

**Bacterial strains**

The seven bacterial isolates used in this research were obtained from the previous study by Wiraswati et al. (2019). The bacterial strains were isolated from rice phyllosphere environment which is taken from Cikembar sub-district, Sukabumi and Situgede sub-district, Bogor, West Java, Indonesia. The bacterial isolates were previously identified as *Bacillus* sp. STGG 3 (GenBank Accession No: MN240450), *B. subtilis* STGG 7, *B. amyloliquefaciens* STGG 8, *B. subtilis* subsp. *subtilis* STGG 14 (GenBank Accession No: MN240448), *B. subtilis* SKBV 1 (GenBank Accession No: MN240449), *Enterobacter cloacae* STGV 8 (GenBank Accession No: MN240451) and *Brachybacterium paraconglomeratum* SKBG 78 (GenBank Accession No: MN240452).

**Greenhouse experiment: efficacy of bacteria to control blast disease**

**Plant material and biocontrol agents.** This experiment used rice variety of Ciherang as plant material which susceptible to rice blast disease. The rice seeds were sown on soil for 14 days and then 5 rice plants were transferred to pots containing the mixture of paddy field soil and organic fertilizer (±300 g). For bacterial application, seven rice phyllosphere bacteria (STGG 3, STGG 7, STGG 8, STGG 14, SKBV 1, STGV 8 and SKBG 78) were inoculated on 200 mL Luria Bertani (LB) broth medium (1% NaCl, 1% Tryptone, 0.5% Yeast extract, 2% agar in 1 L distilled water) for 4 days or until reached ±10^9 cfu mL^-1 of bacterial density. Afterward, these bacterial cultures were applied twice, i.e 7 and 14 days after planting (dap) by spraying a total of 20 mL of bacterial culture per pot. This experiment was run with complete randomized design using 10 treatments which consisted of 7 bacterial isolates, commercial fungicide with isopropanol as active ingredient (Fujiwan 400EC), pathogen (*P. oryzae* race 173) and sterile distilled water. Each treatment was replicated 3 times and each replication consisted of 5 rice plants. The pathogen treatment without bacteria (PT) was used as positive control of disease, while sterile distilled water treatment without pathogen and bacteria (WT) was used as negative control.

**Conidial production.** In this study, we used *P. oryzae* race 173 from Indonesian Center for Rice Research (ICRR) collection. The pathogen mycelial block agar was transferred to Petri dishes containing oatmeal agar medium (3% oatmeal, 2% agar in 1 L of distilled water) and incubated for 10 days at room temperature (±25°C). The fungal cultural was then aseptically scrapped with small brush and exposed with neon lamps for 2 days. Afterward, the conidial were harvested by adding sterilized water containing 0.1% tween 80 and conidial concentration was quantified using haemocytometer and adjusted to final concentration 2 x 10^6 conidial mL^-1. The conidial inoculum was applied at 21 dap by spraying it in to the rice plants. Immediately after pathogen inoculation, the rice plants were transferred to a moist room which maintained at 25 ± 2°C of temperature with relative humidity of 90 ± 5%. The rice plants were then submitted to an initial 48 h dark period for penetration of conidia and blast disease development (Harsonowati et al. 2017). Afterward, the rice plants were transferred to a greenhouse and kept for a long experimental period. The temperature and humidity inside the greenhouse were maintained at 25 ± 2 °C (day) to 20 ± 2 °C (night) and 90 ± 5%, respectively. The non-inoculated pathogens of rice plants were sprayed with sterile distilled water as a negative control, while the positive control of blast disease was sprayed by pathogen inoculum and sterile distilled water.

**Extraction of crude bioactive compounds from isolate STGG 14**

Isolate STGG 14 was grown in 1 L LB broth medium and incubated at 150 rpm for 4 days. Crude bioactive compounds were obtained by using ethyl acetate as solvent (Islam et al. 2012). The bacterial culture was mixed with ethyl acetate with a ratio of 1:1 (v:v) and vigorously shaken for 1 hour. Afterward, the organic layer was collected and evaporated by using vacuum evaporator at 40°C of temperature. The resulted crude extract was stored at 5°C for further analysis.

**Effect of crude extract toward Pyricularia oryzae race 173**

The effect of crude extract from isolate STGG 14 toward the alteration of fungal mycelium was determined using Scanning Electron Microscope (SEM) S-5310LV (Jeol Ltd., Japan). Antifungal activity assay of crude extract was conducted using agar diffusion method. The Potato Dextrose Agar (PDA) medium (3% PDA (Himedia); 1 L distilled water) which contained 500 ppm of crude extract was solidified in Petri dishes and then 5 mm of fungal mycelium block agar was grown above the PDA medium. In addition, fungal pathogen which grown in the
PDA medium without crude extract was used as a negative control. After 10 days of incubation, the fungal mycelium was prepared for SEM observation. Fungal mycelium was excised and prepared by soaking in cacodylate buffer for 2 hours and agitated in ultrasonic cleaner for 5 min. Afterward, sample prefixation was done by soaking in 2.5% glutaraldehyde for 2 days and fixation sample by soaking in 2% tannic acid for several days. Furthermore, sample was cleaned with cacodylate buffer for 5 min; 4 times. Dehydration step was conducted by soaking the sample in a graded alcohol concentration (50%, 70%, 85%, 95%, and 100%) for 20 min for each concentration. The last step was draining the sample by soaking it in tert butanol for 2 x 10 min, freezing it in the freezer and transferring it to the freeze-drying instrument. Before observation step, the dry sample was coated with aurum (Au) using ion coater.

Thin layer chromatography analysis

The crude extract from isolate STGG 14 was separated using TLC plates (TLC silica gels 60 F\textsubscript{254}, 20x20 cm, layer thickness 0.25 mm, Merck, Darmstadt, Germany). The crude extract was dissolved using methanol to final concentration of 200 mg mL\textsuperscript{-1}. Afterward, the dissolved extract was injected into TLC plates using CAMAG Linomat 5 and developed using eluent system: acetone: chloroform (v/v = 1/1) as optimum eluent. Eluent residue was removed by incubating the TLC plates at room temperature for 15 minutes. The separated compounds were visualized under UV light with 254 nm and 366 nm wavelength using CAMAG TLC Visualizer 2.

TLC direct bioautography analysis

Ten microliters of methanol dissolved extract (200 mg mL\textsuperscript{-1}) were loaded to TLC plates (TLC silica gels 60 F\textsubscript{254}, 2x10 cm, layer thickness 0.25 mm, Merck, Darmstadt, Germany) and separated using eluent system: acetone: chloroform (v/v = 1:1) in the TLC chamber. The TLC-direct bioautography test was conducted using agar diffusion method. Mycelia of \textit{P. oryzae} race 173 was harvested from 7 days old of fungal culture (10\textsuperscript{6} cfu mL\textsuperscript{-1}) which previously grew on PDB medium and mixed with 15 mL of PDA medium (48°C). Furthermore, the developed TLC plates were placed aseptically in the Petri dishes and covered with PDA broth medium that contained fungal mycelia. After 10 days of incubation, antifungal activity of crude extract was observed. The active fraction with antifungal activity was characterized by clear zone around the developed TLC plate.

Isolation of antifungal compounds from TLC plate

A total 600 \mu L of crude extract (200 mg mL\textsuperscript{-1} in methanol) was injected into TLC plates (TLC silica gels 60G F\textsubscript{254}, Glass Plates 20x20 cm, layer thickness 0.25 mm, Merck, Darmstadt, Germany) and developed in acetone: chloroform (v/v = 1/1) as mobile phase eluent. Afterward, the developed TLC plate was visualized under UV light and the active fraction with antifungal activity was marked. The active fraction with high antifungal activity was scrapped out from the developed TLC plates and mixed with methanol as solvent. The organic layer was then separated by centrifugation in 5,000 x g and evaporated at 40°C of temperature for 2 days.

Identification of antifungal compounds

The isolated antifungal compounds were identified using LC-MS/MS analysis with Acquity UPLC\textsuperscript{®}H-Class System dan Xevo G2-S QTof (Waters, USA). This analyzer was equipped with capillary column Acquity UPLC\textsuperscript{®}HSS C18 (1.8 \mu m 2.1x100 mm). The LC column and room temperature were programmed at 50°C and 25°C. The mixture of water + 5 mM ammonium formic and acetonitrile + 0.05 % formic acid was used as mobile phase with sample flow rate of 0.2 ml/min. Meanwhile, the desolvation temperature of MS analysis was programmed at 350°C with desolvation gas flow rate and collision energy of 793 L/hour and 4 volts, respectively. Furthermore, the chromatogram and mass spectra were analyzed and interpreted using MassLynx V.4.1 program. The m/z peaks of antifungal compounds were compared with the mass spectrum database in Chemsider (www.chemspider.com) and Massbank website (www.massbank.jp).

Data analysis

\textit{Blast disease assessment and statistical analysis}

The severity of blast symptoms on the experimental rice plant was observed every 3 days until 21 days after inoculation (dai) of pathogen using a standard evaluation system for rice blast with 6 ratings: 0, no lesions; 1, not more than five lesions, and no longer than 10 mm; 3, 6-10 lesions, longer than 10 mm; 5, 11-25 lesions, covering between 10 and 25% of total leaf area; 7, more than 26 lesions, covering between 26 and 50% of total leaf area; 9, lesion area greater than 50% of total leaf area (Yang et al. 2007). At the end of observation, blast disease severity was calculated using the following formula:

\[ \text{Disease severity} = \frac{\sum (n \times v)}{N \times Z} \times 100\% \]

Where: \(n\) is the number of leaves infected by blast disease, \(v\) is value score of blast disease, \(N\) is the total of leaves observed, and \(Z\) is the value of the highest score blast disease. Afterward, means of blast disease severity and disease reduction were calculated and statistically analyzed using a one-way analysis of variance (ANOVA) and Tukey’s honestly significant difference test at 5% probability level using SPSS version 20.0 (SPSS, IBM, Armonk, NY, USA).

RESULTS AND DISCUSSION

\textit{Antifungal activity of phyllosphere bacteria in greenhouse experiment}

The seven bacterial isolates are significantly capable in reducing rice blast disease in greenhouse compared to control of pathogen (PT). The least blast symptoms were discovered in rice plants treated with isolate STGG 14,
compared to rice plants treated with isolate STGV 8 (Figure 1). Furthermore, isolate STGG 14 also significantly reduced blast disease at 70.83% reduction with relative inhibition that was higher than fungicide treatment (Figure 2). Thus, isolate STGG 14 can be developed as biocontrol agents of blast disease dan selected for further analysis in this study.

**Morphological alteration of pathogen-induced by crude extract**

Crude bioactive compounds were extracted from isolate STGG 14 with ethyl acetate as solvent. From 7 L of bacterial culture, we obtained 0.944 g of crude bioactive compounds extract. The effect of crude extract toward *P. oryzae* was displayed by cracked hyphae (Figure 3.B), curly shape and smaller size of hyphae (Figure 3.C) under scanning electron microscope observation. Meanwhile, the hyphal morphology in the control treatment grew normally with compact shaped (Figure 3.A). Furthermore, the growth of *P. oryzae* in the PDA medium with crude extract was also slower than the control experiment.

**TLC analysis of antifungal compounds from isolate STGG 14**

Optimation of mobile phase was trialed using 9 solvent during TLC analysis and resulted in the mixture of acetone/chloroform (v/v = 1/1) as the most optimum eluent for crude extract separation from isolate STGG 14. TLC fractionation of crude extract resulted in 10 fractions on the TLC plate (Figure 4A). The bioautography test showed that fraction 8 (Rf = 0.83) and 9 (Rf = 0.86 ) displayed antifungal activity with no mycelium growth around the spot (Figure 4.B).

**Isolated antifungal compounds**

Two active fractions corresponding to fungal inhibition were successfully purified from TLC plate using methanol as solvent. We obtained 0.074 g and 0.033 g of extract from fractions 8 and 9. Each isolated compound from spot 8 and 9 was further dissolved in methanol for LC-MS/MS analysis.

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**Figure 1.** Blast disease severity on rice variety Ciherang treated with rice phyllosphere bacteria at 21 days after pathogen inoculation compared to control pathogen. Bars shared the same letter is not significantly different according to Tukey's honestly significant difference test ($p \leq 0.05$). WT is water treatment without pathogen and bacteria, PT is pathogen treatment without bacteria.

**Figure 2.** Reduction of blast disease in rice plant treated with rice phyllosphere bacteria and fungicide at 21 days after pathogen inoculation in greenhouse. Bars shared the same letter is not significantly different according to Tukey's honestly significant difference test ($p \leq 0.05$).

**Figure 3.** Morphology of *P. oryzae* race 173 under scanning electron microscope observation with 2000 times of magnification: A. Normal hyphae of *P. oryzae* on the PDA medium; B. cracked hyphae; C. Hyphae with curly shaped; b and c were grown in PDA medium with crude bioactive compounds from isolate STGG 14. Red arrows showed hyphal damage due to crude extract treatment.
Table 1. Identified bioactive compounds produced by isolate STGG 14 and their bioactivities

| Identified compounds | Formula       | Mass | Bioactivity            | References                                |
|----------------------|---------------|------|------------------------|------------------------------------------|
| Psoralen             | C\(_{12}\)H\(_{16}\)O\(_3\) | 186  | Antifungal             | Yu et al. (2017)                          |
| Xanthotoxin (8-methoxypsoralen) | C\(_{20}\)H\(_{10}\)O\(_4\) | 216  | Antifungal, antibacterial | Al-Barwani and Eltayeb (2004)            |
| Moracin C            | C\(_{10}\)H\(_{14}\)O\(_2\) | 298  | Antiinflammatory        | Yao et al. (2016)                        |
| 4,6-dimethyl-3 (4'-hydroxyphenyl) coumarin | C\(_{18}\)H\(_{16}\)O\(_3\) | 266  | Antibacterial           | Asif (2015)                              |

**Antifungal compounds from isolate STGG 14**

According to LC-MS/MS analysis, fraction 8 consisted of four dominant peaks, while fraction 9 consisted of two dominant peaks. The dominant peak 1 (Retention time= 8.65 min) and 2 (Rt= 9.47 min) from fraction 8 as well as dominant peak 1 (Rt= 8.63 min) and 2 (Rt= 9.49 min) from fraction 9 were identified as psoralen and xanthotoxin. Meanwhile dominant peak 3 (Rt= 10.08) and 4 (Rt= 10.86) from fraction 8 were successfully identified as moracin C and 4,6-dimethyl-3 (4'-hydroxyphenyl) coumarin. These identified compounds belong to furanocoumarin group except moracin C. The psoralen and xanthotoxin were reported to have antifungal activity against several fungal pathogens on the previous study (Table 1).

**Discussion**

The seven rice phyllosphere bacteria from West Java, Indonesia previously proved to have antifungal activity against *P. oryzae* in vitro analysis and potentially developed as biocontrol agents of blast disease. Before releasing the potential bacteria to the field, we have to prove that their antifungal activities against *P. oryzae* are also effective in rice plants. Therefore, this present study aims to reveal the efficacy of these bacterial isolates in reducing blast disease severity in greenhouse. The *in planta* analysis of bacterial isolates shows that the isolate has an ability to decrease severity of blast disease in rice variety Ciherang. All phyllosphere bacterial treatments significantly reduce blast disease severity compared to pathogen treatment (PT) with various relative inhibition percentages. Isolates STGG 3, STGG 4, STGG 7, STGG 8 and SKBV 1 which are identified as *Bacillus* spp. show higher antifungal activity than isolates SKBG 78 and STGV 8 which are identified as *Brachybacterium paraconglomeratum* and *Enterobacter cloacae*. The ability of *Bacillus* spces to inhabit the various environments will facilitate the bacteria to survive and colonize the aerial part of rice plant. Beside, *Bacillus* species have a role in plant growth stimulation as studied by Sunera et al. (2020) that isolated *B. cereus* from rhizosphere of tomatoes plant. In addition, *Bacillus* spp. also produce a number of bioactive compounds that have important roles in decreasing rice blast severity. Meanwhile, *B. paraconglomeratum* and *E. cloacae* which are known as plants growth promoter are predicted to have lower ability in colonizing the rice leaves, so that they display low antifungal activity.

Among seven rice phyllosphere bacteria, isolate STGG 14 shows the highest activity in reducing blast disease with 70.83% of relative inhibition (disease severity: 23.11%). Interestingly, isolate STGG 14 also shows higher blast disease reduction than commercial fungicide, Fujiwan 400 EC (Figure 2) which is commonly used by farmers in Indonesia. Isolate STGG 14 which is identified as *B. subtilis* subsp. *subtilis* is commonly used as biocontrol agents of various plant pathogens (Zhang et al. 2012) such as *Fusarium oxysporum*, *Colletotrichum capsici*, *Rhizoctonia solani*, etc. On the other hand, isolate STGV 8 which is identified as *E. cloacae* displayed the lowest relative inhibition to blast disease, ie. 48.35% (disease severity: 41.07%). This finding indicated that *B. subtilis* subsp. *subtilis* has a better ability than *E. cloacae* (isolate STGV 8) in reducing rice blast disease severity in greenhouse.

In this study, *B. subtilis* subsp. *subtilis* STGG 14 displayed constant antifungal activity both in vitro and in planta analysis. The efficacy of *B. subtilis* subsp. *subtilis* STGG 14 in reducing blast disease is caused by its ability to produce antifungal compounds. Our previous study successfully amplified Adenilation (A) domain as well as Ketasynthase (KS) domain of cluster gene NRPS and PKS from *B. subtilis* subsp. *subtilis* STGG 14 (Wiraswati et al. 2019). A domain and KS domain have a role in antimicrobial compounds synthesis which belong to nonribosomal peptide and polyketide groups such as lipopeptide, macroide, and polyene compounds. Several previous studies mentioned that *B. subtilis* subsp. *subtilis* can produce antifungal compounds like iturin, fengycin, surfactin (Dimiki et al. 2017) and bacilysocin (Tamehiro et al. 2002).

**Figure 4.** Purification of antifungal compounds from isolate STGG 14: A. TLC profile of crude bioactive compounds resulted in 10 fractions; B. TLC directed bioautography test observed after 10 days of incubation
In addition, Islam et al. (2012) and Luo et al. (2015) reported that *B. subtilis* subsp. *subtilis* C9 and *B. subtilis* 916 produced acetyl butanediol, bacilomycin L and surfactin that suppressed the growth of *R. solani*. *Bacillus subtilis* strain Y-IVI was able to produce iturin A and fengycin which have a role in suppressing *F. oxysporum* in Muskmelon (Zhao et al. 2016). Meanwhile, Hanene et al. (2012) also reported that fengycin produced by *B. subtilis* SR146 could inhibit the growth of *F. oxysporum*, *F. melonis*, *F. equiseti*, and *F. solani*.

As mentioned before, *Bacillus* species including *B. subtilis* are very beneficial for agricultural purposes due to its ability to produce bioactive compounds. To prove it, we extracted crude bioactive compounds from isolate STGG 14 which were previously known to have antifungal activity against *P. oryzae* race 173 in vitro analysis (Wiraswati et al. 2019). The results of this study can be seen as evidence that isolates STGG 14 could produce antifungal compounds, which contribute to the rice blast disease control in greenhouse. To determine the fungicidal effect of extracted bioactive compounds, morphological observation was conducted toward *P. oryzae* race 173. According to Scanning Electron Microscope (SEM) observation, the crude antifungal compounds triggered abnormal fungal mycelium of *P. oryzae* that characterized by cracked hyphae (Figure 3B), curly shaped and smaller size hyphae (Figure 3C) than control hyphae. Meanwhile, *P. oryzae* without antifungal compounds treatment has a compact shaped of mycelium and grew normally (Figure 3A). As reported by Semangu (2006), antifungal compounds from plants and microbial extracts displayed several inhibition mechanisms toward fungal pathogens such as cell membrane permeability alteration and cell wall degradation. Suriani et al. (2015) declared that the alteration of hyphae and spores size of *P. oryzae* caused by lysis of mycelium cell phenomenon. Additionally, Liao et al. (2016) also informed that the application of fengycin from *B. amylophilica* BPDL toward *P. oryzae* resulted in swollen and cracked mycelium. The inhibition of fungal growth by antifungal compounds from isolate STGG 14 informs us that the crude extract displays fungistatic mode of action, because the extracted antifungal compounds only inhibit the growth of *P. oryzae* race 173 without killing the pathogen.

The identification of antifungal properties from isolate STGG 14 is important to understand the specific mechanism of the bacteria in controlling blast disease. Crude extract was separated using thin-layer chromatography and then antifungal compounds were detected by TLC-direct bioautography. Eluent optimisation was resulted in a combination of chloroform and acetone (v/v =1/1) as the most effective eluent to separate the crude extract. We obtained 10 fractions from TLC separation and among them, 2 fractions displayed antifungal activity against *P. oryzae* through bioautography test. Antifungal activity was characterised by clear zone formation around the spot 8 (Rf = 0.83) and 9 (Rf = 0.86) (Figure 4b). According to LC-MS/MS analysis, fraction 8 consists of 4 bioactive compounds i.e moracin C, psoralen, 4,6-dimethyl-3 (4′-hydroxyphenyl) coumarin and xanthotoxin (8-methoxypsoralen), while fraction 9 consists of 2 bioactive compounds i.e psoralen and xanthotoxin (8-methoxypsoralen) (Table 1). These identified bioactive compounds have resulted from crude extract which isolated extracellularly from isolate STGG 14.

Interestingly, all identified compounds (except moracin C) belong to the furanocoumarins group which is generally responsible for antimicrobial activities (Aslam et al. 2010). In addition, among all identified compounds, only psoralen and xanthotoxin which displayed antifungal activity against some fungal pathogens according to several previous studies. Furanocoumarins are linear coumarins with benzopyrone compounds that belong to flavonoid groups that usually presented in several natural products and obtained by plant extraction or microbial synthesis. More than 1300 coumarins have been identified as secondary metabolites from plants, bacteria, and fungi such as *Aspergillus* sp., *Ganoderma lucidum*, *Penicillium* sp., *Streptomyces* strain and *Bacillus* strain (Costa et al. 2016). Coumarin derivatives such as xanthotoxin and bergapten demonstrated antifungal activity against *Alternaria* sp., *Bipolaris* sp., and *Fusarium* sp. (Al-Barwani and El-tayeb 2004). Meanwhile, psoralen, linear furanocoumarins, also displayed antifungal activity against *R. solani*, *Botrytis cinerea*, *Alternaria solani* and Gibberela zeae in vitro analysis (Yu et al. 2017). Despite their potential as antifungal compounds, some coumarins are also regarded as phytoalexins which are synthesized by plant tissues in response to the pathogens infection, thus triggering the plant resistance (Yang et al. 2017). Psoralen and xanthotoxin compounds which successfully identified from *B. subtilis* subsp. *subtilis* STGG 14 has not been reported before. Therefore, the research finding that shows *B. subtilis* subsp. *subtilis* as bacteria having ability to produce antifungal compounds from coumarins group except lipopeptides compounds become new information that never been reported before.

To conclude, isolate *B. subtilis* subsp. *subtilis* STGG 14 shows the highest activity in reducing blast disease severity in rice plant variety Ciherang. As the most potential bacteria, isolate *B. subtilis* subsp. *subtilis* STGG 14 is proved to have capability to produce 2 antifungal compounds, i.e psoralen, and xanthotoxin which belong to furanocoumarins group. This result reveals that antibiosis mechanism of isolate *B. subtilis* subsp. *subtilis* STGG 14 contributes to the reduction of blast disease severity in rice plants.

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