INTRODUCTION

Chilli plant is one of the horticultural commodities that has economical value in Indonesia. The demand for chilli is increasing, encouraging farmers to cultivate chilli plants. One of the chillies that are cultivated by farmers in the Tapin area of South Kalimantan namely Hiyung cayenne pepper. The year 2012 Hiyung chili pepper is listed on the Center for Crop varieties Protection and Agriculture Licensing Ministry of Agriculture Republic of Indonesia No. 09/PLV/2012 dated 12 April 2012 as a local variety with the name of Hiyung cayenne pepper (Balai Penelitian dan Pengembangan Pertanian, 2018), and in June 2012 the Ministry of Agriculture of the Republic of Indonesia established Hiyung cayenne pepper as a national variety. This cayenne pepper can support national chili production because of its high productivity with good market prospects, growing both on swampy and dry land. Compared to the three commercial varieties (Sonar, Bara, and Santika) Hiyung cayenne pepper has the highest dry weight, and higher productivity with a longer harvest duration, with the highest capsaicin levels about 699 ppm (Pramudyani, Sabran, & Noor, 2019).

Plant disease is one of the factors of failure in harvesting, including in chili plants (Islam, Schreinemachers, & Kumar, 2020). Species of fungi that can infect chili plants include Colletotrichum scovillei (Caires et al., 2014), Aspergillus spp.,

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ABSTRACT

Chili farming faces several constraints, one of which is the pathogenic fungus Colletotrichum capsici. To overcome it can be used indigenous endophytic fungus and liquid smoke wood Ulin (Eusideroxylon zwageri Teijsm. & Binn.) which has the potential as antimicrobial can be used. This research aimed to quantify and measure the effectiveness of an antimicrobial liquid smoke, endophytic filtrate, and the combination to suppress C. capsici growth. Subsequently, the research was conducted to apply the liquid smoke, endophytic fungi, and the two combinations of treatments on the growth of C. capsici. Thus, the results of this research showed that liquid smoke with a concentration of 0.085-1.75% can inhibit 3.56-62.17% in range. Meanwhile, the endophytic fungi filtrate, of 2% concentration can inhibit 91.69% C. capsici. Two of the combination liquid smoke in a concentration of 0.68%, 1.36% and the endophytic fungi filtrate in 2% have shown to inhibit the growth of C. capsici with the highest inhibition into 88.08%. Based on the analysis results, liquid smoke, endophytic fungi filtrate, and a combination of both showed significantly different inhibitory effects between treatments. This indicates that all those three treatments have antimicrobial potential.

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Fusarium spp., Colletotrichum spp. (Frimpong et al., 2019) 8 genera, and 17 species were identified on the basis of morphology, culture characteristics, and DNA sequencing of the internal transcribed spacer (ITS, and many more fungi have been reported. Fungi that cause disease in plants often cause structural and physical damage (Marques, Soares, & Appezzato-Da-Gloria, 2013). Safe and environmentally friendly control of pathogenic fungi can be done by utilizing microorganisms derived from the plant itself, such as endophyte microbes (Köhl, Postma, Nicot, Ruocco, & Blum, 2011).

Endophytic microbes can derive from bacteria and fungi groups that have the ability to create a colony, part or all of its life cycle on a plant network without harming its host (Köhl, Postma, Nicot, Ruocco, & Blum, 2011; Selim, El-Beih, AbdEl-Rahman, & El-Diwany, 2012). Endophytic fungi are found in the plant tissue system, including flowers, twigs, leaves, and plant roots. These microorganisms grow and take food from the plant, and can infect healthy crops in certain tissues as well as able to produce mycotoxin, enzymes, and antibiotics (Stone, Polishook, & White Jr, 2004). Endophyte fungi also can inhibit pathogenic microbes that cause plant disease through mechanisms i.e. space and nutrition competitions and producing bioactive compounds such as antibacterial and antifungal (Gao, Dai, & Liu, 2010). Fungi also have the ability to produce plant growth hormones such as auxin (Imaningsih, Kadarsah, & Rusmannurrachmad, 2019) and herbicidal activity (de Souza et al., 2017).

Control of pathogenic fungi as a plant destruction organism can also be done by utilizing liquid smoke, a vapor condensate of pyrolysis of wood containing the main compounds of acids, phenols, and carbonyl (Lee et al., 2011). The constituent components of the liquid smoke compound have the ability as antimicrobial. Endophytic molds and liquid smoke have the same ability to act as antimicrobials, but their potential is unknown when combined between the two. This research was intended to study the ability of endophytic fungi and liquid smoke to inhibit the growth of pathogenic fungi and examine the ability of endophytic fungi added by a combination of liquid smoke at different concentrations to the growth of pathogenic fungi.

**MATERIALS AND METHODS**

**Isolation and Purification of Endophytic Fungi**

This research was conducted from November 2018 to April 2019 at the Microbiology Laboratory of the Faculty of Mathematics and Natural Sciences, Lambung Mangkurat University, Banjarbaru, South Kalimantan. Samples of chili cayenne pepper plants were taken from the village of Hiyung, Tapin District, South Kalimantan in November 2018. The chili plants that were taken were put in polybags and coded as sampling information. Isolation and purification of endophytic fungus were carried out to obtain pure isolates by direct planting methods. The procedure is performed based on the method used by Septiana, Sukarno, Sukarno, & Simanjuntak (2017) with modification.

**Identification and Screening of Endophytic Fungi**

The fungi Identification includes macroscopic and microscopic observation. The macroscopic observation of pure isolates was done by observing the shape, color, and diameter of the colony for 7 days, the color and presence of the Hypha region determined on the 7th day. Microscopic observation was done by the method of slide culture (Rosana, Matsuzawa, Gonoi, & Karuniawati, 2014). Isolation identification refers to the Fungi identification book: Illustrated Genera of Imperfect Fungi (Barnett & Hunter, 1998) and The Genera of Hyphomycetes (Seifert & Gams, 2011).

Screening of endophytic fungi is conducted through the test of pathogenicity and antagonism. The pathogenicity test was carried out against the endophyte isolates that were derived from the roots of healthy chili plants based on the methods done...
Percent germination is obtained by dividing the number of germinated seeds (normal/abnormal/not growing) by the total number of seeds multiplied by 100%. The antagonism test refers to the dual culture method in the PDA medium (Tomah, Abd Alamer, Li, & Zhang, 2020). The inhibitory percentage was obtained by dividing the difference from the diameter of the control pathogen colony and the diameter of the treatment pathogen colony by the diameter of the control pathogen multiplied by 100% (Kunova et al., 2016).

**Production of Endophytic Fungus Filtrate**

The endophytic fungus was prepared using the method of Qiao, Ling, Yu, Huang, & Wang (2017) that has been modified. The endophytic fungus was inoculated in a slanted PDA medium incubated for 7 days, then harvested by adding 9 ml of sterile distilled water to a tube containing endophytic fungi. The culture was homogeneous using a soft brush to obtain a spore suspension, then the suspension was transferred into another sterile test tube. Fungus suspension is centrifuged at a speed of 3000 rpm for 20 minutes to get the filtrate supernatant. The results of the process were separated using a 0.45 µm syringe filter and used as a crude biocontrol agent for further testing.

**Inhibition Ability Test of Liquid Smoke Endophytic Fungi on C. capsici**

Test the ability to inhibit liquid smoke is carried out by the agar dilution method (Balouiri, Sadiki, & Ibnsouda, 2016). The liquid smoke used in this study came from Ulin wood (Eusideroxylon zwageri Teijsm. & Binn.) obtained from the result of condensation of Ulin charcoal smoke production in Ranggang Village, Takisung District, Tanah Laut Regency, South Kalimantan. This method is carried out by mixing the media with liquid smoke, and the pathogen C. capsici is inoculated on the media that has been mixed with liquid smoke. The concentration used in this test was 0.00%; 0.085%; 0.17%; 0.34%; 0.68%; 1.36%; 1.75%. Observations were made from day 1 to day 7 after inoculation. The minimum inhibitory concentration was determined by the dilution method, using the method of Venkateswarulu et al. (2018) that has been modified. The concentration of endophytic fungus used in this test was 0%; 2%; 4%; 6%; 8%; 10% Positive control treatment using ketoconazole 200 mg at a concentration of 2%. Pathogenic mold pieces 6 mm inoculated in the middle of the PDA medium positive control treatment and concentration treatment of endophytic fungus filtrate, then incubated at 28°C for 7 days, and observations were made every day. The minimum inhibitory concentration is determined from the presence or absence of pathogenic fungus growth at the lowest concentration of endophytic fungus filtrate.

An inhibitory test of the combination of endophytic fungus and liquid smoke was carried out based on the method carried out by Balouiri, Sadiki, & Ibnsouda (2016) the combination of endophytic fungus filtrate (EFF) and liquid smoke (LS) was 2% for filtrate and 0.68%, 1.36% for liquid smoke. So the first combination is 2% (EFF) + 0.68% (LS) and 2% (EFF) + 1.36% (EFF).

A 6 mm pathogenic fungus is inoculated in the middle of the PDA medium in a Petri dish. Observation of inhibition of pathogenic fungus was carried out from day 1 to day 7 after inoculation. The inhibitory effect was obtained by dividing the difference between the diameter of the control pathogen colony and the diameter of the treatment pathogen colony by the diameter of the control pathogen multiplied by 100% (Kunova et al., 2016).

**Data Analysis**

Analysis of the data using One-Way ANOVA at alpha 0.05 was followed by Duncan’s test, or when the data is not homogeneous, the Kruskal Wallis test is used. All data were analyzed using IBM SPSS Statistic Version 22, 2013.

**RESULTS AND DISCUSSION**

**Diversity of Endophytic Fungi from Roots of The Hiyung Cayenne Pepper**

Isolation and purification of endophytic fungi derived from chili cayenne root were obtained by 8 isolates and given a code name using a sample source that is chili root (ACH). Isolates obtained from chili cayenne roots were characterized by macroscopically and microscopically. Pure isolates obtained were Trichoderma sp. ACH1.1, Trichoderma sp. ACH1.6, Botrytis sp. ACH2.2, Botrytis sp. ACH2.3, Gliocladium sp. ACH2.4, Harmoniella sp. ACH2.5, Humincola sp. ACH2.6, Cunninghamamella sp. ACH2.7.

The factors affecting the presence of endophytic fungi include the environment and the plant tissue used (Maheswari & Rajagopal, 2013). The Habitat of plant origin is one of the environmental factors affecting the structure and type of microbes that colonize the plant tissue such as roots, stems,
leaves, and branches (Araújo et al., 2002). Sieber & Grünig (2006) mentions that affecting the diversity of endophytes derived from plants is environmental factors, vegetation, and interactions with other types of microbes.

**Pathogenicity and Antagonism of Endophytic Fungi from the Roots of Hiyung Cayenne Pepper**

Screening of endophytic fungi was carried out by pathogenicity and antagonistic tests. The pathogenicity test was carried out on pure isolates resulting from the isolation of endophytic fungi. The percentage rate of pepper seed germination on the 7 and 14th day after inoculation (DAI) in the pathogenicity test is presented in Table 1. The fungus penetrates its host through mechanical and enzymatic mechanisms (Ashry & Mohamed, 2012). Fungi penetrate the epidermis, the cuticles, and cell walls (Underwood & Somerville, 2008). Enzymes that act as degenerating cell walls are pectinase and cellulase, where these enzymes are used for fungus for the process of penetration and colonization of host plants (Ashry & Mohamed, 2012; Kikot, Hours, & Alconada, 2009).

The antagonistic test was carried out on 3 selected isolates from the results of the pathogenicity test using the dual culture method. The results of antagonistic tests based on diameter measurements of pathogenic fungi colonies and inhibitory effect can be seen in Table 2. This antagonistic nature is consistent with the statements of De la Cruz-Quiroz, Roussos, Rodríguez-Herrera, Hernandez-Castillo, & Aguilar (2018) that fungus is grown side by side and has the ability to grow faster, then these fungi are able to occupy space and suppress the growth of their opponent’s fungus. This antagonistic nature occurs because of the same needs as each fungus, nutrition, and growing needs.

Endophytic screening results showed isolates of Cunninghamella sp. ACH2.7 was selected as an isolate used for testing endophytic fungus filtrate and in combination using liquid smoke. This was obtained from the Kruskal Wallis test where the isolate had the highest normal growing percentage of sprouts, the lowest abnormal sprouts, and the lowest ungrown sprouts as well as the highest percent inhibition of pathogens (Table 1 and Table 2).

**Table 1.** The percentage rate of pepper seed germination on the 7 and 14th day after inoculation (DAI) in the pathogenicity test

| Endophytic fungi       | Germination (%)* | 7th-DAI | 14th-DAI |
|------------------------|------------------|---------|---------|
|                        | Normal | Abnormal | Not grow | Normal | Abnormal | Not grow |
| Without endophytic fungi addition | 100±0c   | 0±0a     | 0±0a     | 100±0c  | 0±0a     | 0±0a     |
| *Trichoderma* sp. ACH1.1 | 0±0ab  | 0±0a      | 100±0ab  | 30±51.96ab  | 6.67±5.77a | 63.33±55.07ab |
| *Trichoderma* sp. ACH1.6 | 0±0ab  | 0±0a      | 100±0ab  | 33.33±41.63ab  | 10±10a     | 56.67±45.09ab |
| *Trichoderma* sp. ACH2.2 | 0±0a    | 23.33±20.82ab  | 76.66±20.81ab  | 0±0a   | 60±30ab  | 40±30ab  |
| *Botrytis* sp. ACH2.3 | 0±0a    | 16.67±28.86ab  | 83.33±28.86ab  | 0±0a   | 30±26.46ab  | 70±26.45ab  |
| *Gliocladium* sp. ACH2.4 | 0±0a    | 0±0a      | 100±0b   | 0±0a   | 26.67±30.55a  | 73.33±30.55b  |
| *Harmoniella* sp.ACH2.5 | 0±0abc  | 3.33±5.77a  | 96.66±5.77ab  | 90±0abc  | 3.33±5.77a  | 6.67±5.77ab  |
| *Humicola* sp. ACH2.6 | 10±17.32abc  | 0±0a      | 90±17.32ab  | 73.33±23.1abc  | 13.33±11.55a | 13.33±11.54ab  |
| *Cunninghamella* sp. ACH2.7 | 66.67±25.16bc  | 0±0a      | 33.33±25.16ab  | 90±10bc  | 0±0a | 10±10ab  |

Remarks: * The number followed by the same letter is not significantly different based on Duncan (α=0.05)
The Ability of Liquid Smoke and Endophytic Fungi Inhibits C. capsici

Based on the test results of liquid smoke ability inhibit the growth of C. capsici, obtained at all concentrations of liquid smoke is able to inhibit growth. The concentration of ironwood liquid smoke from 0.085-1.75% can inhibit C. capsici by 3.56-62.17%. But only at concentrations of 0.34%, 0.68%, 1.36%, and 1.75% showed significant inhibition compared to control (0.00% liquid smoke concentration). Inhibitory effect (%) of liquid smoke against C. capsici on the 1-7th day after inoculation is shown in Fig. 1. The inhibition effect was significantly different between treatments (F=11.053, P=0.000). Ironwood liquid smoke inhibits C. capsici causing changes in the diameter of the colony. The increasing concentration of liquid smoke the smaller the colony that forms (Fig. 2). The condition occurs because the content of liquid smoke affects the growth of fungi. Phenol and acid compounds in liquid smoke can damage the structure of the fungus. This is in line with research of Suresh et al. (2019) which uses pyroligneous acid from a mixture of several kinds of wood that has the ability to inhibit the growth of fungi. The acid content of liquid smoke causes acid conditions in the cytoplasm, therefore causing damage to membrane surface tension and loss of active transport, resulting in unstable function and structure of cell components (Hassan, Sand, & El-Kadi, 2012).

Table 2. Pathogenic fungi colony diameter and inhibitory effect of endophytic fungi against C. capsici 7th day after inoculation (antagonisms test result)

| Endophytic fungi       | Colletotrichum capsici Diameter (mm) | Inhibitory effect (%)* |
|------------------------|--------------------------------------|------------------------|
| Harmoniella sp. ACH2.5 | 43.33±2.82                           | 10.71±4.79 b           |
| Humicola sp. ACH2.6    | 46.50±2.55                           | 5.11±0.18 c            |
| Cunninghamella sp. ACH2.7 | 32.23±1.50                           | 33.26±2.54 a           |

Remarks: * The number followed by the same letter is not significantly different based on the Kruskal Wallis Test (α=0.05)
The best concentration of liquid smoke is then selected for the combination test. Endophytic fungal filtrate tests selected from previous tests were also conducted to determine the minimum inhibitory concentration (MIC) to be used to test the combination of endophytic fungal filtrate and liquid smoke. Based on the test results, endophyte fungi filtrate can inhibit the growth of *C. capsici*. The results of the inhibition of endophilic fungus filtrate can be seen in Fig. 3. Inhibitory effects differ significantly between treatments (*F* = 41.634, *P* = 0.000). All concentration treatments differ significantly from control (ketoconazole 2%). Even at a concentration of 2%, *Cunninghamella* sp. ACH2.7 filtrate is already able to inhibit *C. capsici* (91.69%), even better when compared to ketoconazole 2% 200 mg as a positive control.

In addition to inhibiting growth, isolate *Cunninghamella* sp. ACH2.7, also causes the morphology of *C. capsici* to change. One of them is a colony that was originally blackish gray to be lighter gray-white on PDA medium. These discoloration and morphology are likely caused by the ability of compounds produced by endophytic fungi to damage the structure of *C. capsici*, this requires further research. De la Cruz-Quiroz, Roussos, Rodríguez-Herrera, Hernandez-Castillo, & Aguilar (2018) in research on the ability of *Trichoderma* to inhibit *P. capsici* states that the pathogen cell part is used as a source of nutrition for the growth of *Trichoderma*, the same is likely to happen in this study.

### The Synergistic of Endophytic Mold and Liquid Smoke Filtrate Inhibits *C. capsici*

Based on the previous test results, 2 concentrations of ironwood liquid smoke (LS) were used (0.68% and 1.36%) and MIC concentration of filtrate *Cunninghamella* sp. ACH27(2%) (EFF) to be combined so that the synergy can be known. The combination of 0.68% LS and 2% EFF, as well as 1.36% LS and 2% EFF is able to inhibit the growth of *C. capsici* differs significantly with 2% ketoconazole as control (*F*=14.676, *P*=0.000) (Fig. 4). The utilization of a combination of liquid smoke and endophytic fungus filtrate as an antimicrobial can be developed with the use of the right concentration because both of these ingredients have active compounds that can be utilized. Liquid smoke and endophyte fungi have the same benefits that act as antibacterial and antifungal. In line with Aisyah, Sinaga, Nawangsih, Giyanto, & Pari (2018) that liquid smoke has the ability as an antibacterial agent and stimulates plant growth, in research on liquid smoke from some wood to overcome the banana disease. Gunatilaka (2006) also stated that endophyte fungi can produce secondary metabolites and compounds that act as an antifungal and antibacterial agent.

The combination of liquid smoke and the endophytic filtrate is able to inhibit growth and cause the morphology of *C. capsici* to change. Morphological changes in *C. capsici* due to the presence of active compounds that can damage cell membranes. Phenols from liquid smoke and flavonoids from endophytic fungi can improve the permeability of cell membranes, as well as inhibit the activation of essential enzymes, and the functioning of genetic materials (Konaté et al., 2012).
The treatment of a combination of liquid smoke and endophytic fungi filtrate in vitro is able to inhibit the growth of *C. capsici*, so from these results, there is the potential that the combination of those two compounds can be antimicrobial agents that can be used to control disease-causing pathogens in plants. Utilization of a combination of liquid smoke and endophytic fungi filtrate as antimicrobial agents can be developed with the use of appropriate concentrations because both of these materials have active compounds.

**Fig. 3.** Inhibitory effect (%) of *Cunninghamella* sp. ACH2.7 filtrate against *C. capsici* on the 1-7th day after inoculation the bar indicates standard deviation between replicates. The concentration of crude extract followed by the same letter is not significantly different based on Duncan test (α=0.05)

**Fig. 4.** Inhibitory effect (%) of the combination of Liquid Smoke (LS) and *Cunninghamella* sp. ACH2.7 filtrate/Endophyte Fungi Filtrate (EEF) against *C. capsici* on the 1-7th day after inoculation the bar indicates standard deviation between replicates. The concentration of crude extract followed by the same letter is not significantly different based on Duncan test (α=0.05)
CONCLUSION

The combination of liquid smoke and endophytic fungi filtrate has the ability to inhibit C. capsici, and has the potential to be antimicrobial. The highest inhibitory power was generated in a combination of 1.36% liquid smoke and 2% endophytic fungi filtrate at 88.08%. The benefits of the combination of liquid smoke and endophytic fungi filtrate need further research, especially its potential as antimicrobial and appropriate concentration so that it can be utilized to control pathogens that can damage crops especially Hiyung cayenne pepper.

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