17 Mei 2024 (Hasil Review ke-1)

1 Juli 2024 (Hasil Review ke-2)

2 Juli 2024 (Email LOA)

HASIL REVIEW ARTIKEL KE-1 17 MEI 2024

BIO Web of Conferences xx, xx ICoBEAF 2024

Antagonistic Activity of Endophytic Fungi from Maize Plants (*Zea mays*) Against *Fusarium oxysporum*

Oktira Roka Aji1*, and Dilla Rofiyanti1

¹Biology Department, Faculty of Applied Science and Technology, Universitas Ahmad Dahlan, Bantul, Yogyakarta 55191, Indonesia

Abstract. Endophytic fungi are microorganisms that can reside within plant tissues without harming their host plants. These fungi can act as biological agents against harmful fungi by utilizing mechanisms like competing for nutrients, parasitism, and antibiosis. This research aims to isolate and identify the genus groups of endophytic fungi from maize plants (Zea mays), understand the antagonistic mechanisms between endophytic fungi and Fusarium oxysporum through dual culture tests, determine the highest inhibitory percentage of endophytic fungi against Fusarium oxysporum in filtrate culture tests and volatile compound tests. Determining antagonistic activity against pathogenic fungi is carried out in vitro through dual culture methods, filtrate culture methods, and volatile compound production. Data obtained from the antagonistic activity of endophytic fungi were analyzed based on the calculation of inhibition percentages. The dual culture method revealed eight isolates effectively restraining the growth of Fusarium oxysporum. Among the filtrate culture experiments, isolate B1.1.1 exhibited the most promising outcomes with an inhibition rate of 39.83%. Notably, isolate D2.2.1 displayed the highest inhibition rate of 44.38% in the antagonism test employing the volatile compound production method. Identification was conducted through ITS gene amplification, confirming that isolate D2.2.1 corresponds to Trichoderma harzianum.

1 Introduction

Plants are vulnerable to numerous diseases, and fusarium wilt is one of them. It typically affects the roots or the base of the plant stem. The presence of fusarium wilt disease can be identified by the symptoms that appear on the top of the plant [1]. The affected plants exhibit symptoms such as rotten roots and are prone to falling over, making them easy to uproot [2]. Fusarium wilt disease is caused by a pathogenic fungus called *Fusarium oxysporum* Schltdl. This soil-borne pathogen can survive in unfavorable conditions and even form chlamydospores without host plants [3]. The fungus is known for attacking various plants such as corn, sugarcane, rice, and sorghum. It causes rot in the stems, cobs, and seeds of corn [4].

Commented [Tz1]: This research aimed to..... (cek lagi mohon abstrak bisa dibuat past tense)

Commented [Tz2]: Mohon bisa ditampahkan kata kunci yang berkaitan dengan SDG's e short titles of the 17 SDGs are: No poverty (SDG 1) Zero hunger (SDG 2) Good health and well-being (SDG 3) Quality education (SDG 4) Gender equality (SDG 5) Clean water and sanitation (SDG 6) Affordable and clean energy (SDG 7) Decent work and economic growth (SDG 8) Industry, innovation and infrastructure (SDG 9) Reduced inequalities (SDG 10), Sustainable cities and communities (SDG 11) Responsible consumption and production (SDG 12) Climate action (SDG 13) Life below water (SDG 14) Life on land (SDG 15) Peace, justice, and strong institutions (SDG 16) Partnerships for the goals (SDG 17).

Commented [Tz3]: Mohon bisa ditambahkan kaitannya dengan SDG's e short titles of the 17 SDGs are: No poverty (SDG 1) Zero hunger (SDG 2) Good health and well-being (SDG 3) Quality education (SDG 4) Gender equality (SDG 5) Clean water and sanitation (SDG 6) Affordable and clean energy (SDG 7) Decent work and economic growth (SDG 8) Industry, innovation and infrastructure (SDG 9) Reduced inequalities (SDG 10), Sustainable cities and communities (SDG 11) Responsible consumption and production (SDG 12) Climate action (SDG 13) Life below water (SDG 14) Life on land (SDG 15) Peace, justice, and strong institutions (SDG 16) Partnerships for the goals (SDG 17).

^{*} Corresponding author: oktira.aji@bio.uad.ac.id

Farmers typically rely on fungicides to manage plant diseases. However, the recurrent use of these chemicals can negatively impact human health and the environment due to the residual effects they leave behind. Consequently, there is a need to explore alternative eco-friendly approaches. One such strategy involves using endophytic fungi as biological agents to control diseases caused by plant pathogens [5]. This approach may offer a promising solution to the problem of fungicide overuse and its harmful consequences.

2 Material and Methods

2.1 Sample preparation

Maize plant samples were obtained from Pathuk District, Gunung Kidul. The selected plants were approximately 80 days old, healthy, and devoid of deformities. Samples were taken from the roots, stems, and leaves to isolate endophytic fungi.

2.2 Endophytic fungal isolation

The isolation of endophytic fungi was done using a direct seed-planting method. Maize plants' leaves, roots, and stems were washed with running water until they were utterly soil-free. Corn plant organ samples were sterilized by placing them in 70% (v/v) alcohol for 1 minute, then putting in a 0.5% (v/v) NaClO solution for 3 minutes, then placing them in 70% (v/v) alcohol for 30 seconds, and rinsing them with sterile distilled water twice and drying them on sterile Petri dishes [6,7].

Organ samples were cut using a sterile scalpel into 1-2 cm sizes as eptically. The organ samples were placed on PDA media and added with 100 μ g/mL chloramphenicol as eptically. A total of 0.1 mL of sterile distilled water from the last rinse was poured on the PDA media by the spreading method to ensure that the surface sterilization process on the organ samples was successful. The culture was incubated at room temperature (25 - 28°C) for seven days [6,7].

The successfully isolated fungi were then subcultured on PDA media and incubated for three days at room temperature (25 - 28°C). Macroscopic observations were made by observing the morphology of the fungal mycelium, which included the color of the upper surface colony, the color of the reverse colony, and the texture of the colony surface. Microscopic observations were carried out with a pure culture of endophytic fungi isolated aseptically using a sterile ose, transferred to the surface of a glass object, and then dripped with Lactophenol cotton blue. The preparations were observed with a light microscope. Observations include the structure of hyphae (concentrated or non-concentrated) and the structure of spores [8].

2.3 Dual Culture Assay

To carry out an antagonism test, small pieces of pure culture of the fungus *Fusarium* oxysporum Schltdl. (\pm 5 mm) and each endophytic fungal isolate (\pm 5 mm) was inoculated in one petri dish containing PDA media at a distance of 3 cm. As a negative control, small pieces of *F. oxysporum*. (\pm 5 mm) was planted in the center of the Petri dish containing PDA media. Isolates were incubated for seven days [9].

Commented [Tz4]: Bisa ditambahkan kenapa mengambil sampel jagung dari lokasi tersebut dan kenapa 80 hari ?

2.4 Antifungal non-volatile compounds test

Pure cultures of endophytic fungi were inoculated in 10 mL of PDB media. The endophytic fungal culture was incubated for ten days at room temperature (25-28°C). The fungal culture was filtered using filter paper, and the filtrate was filtered again using a 0.22 μ m sterile syringe filter. The cell-free filtrate was mixed with sterile PDA media in petri dishes. Pieces of fungal culture of *Fusarium oxysporum* Schltdl. (± 5 mm) were inoculated at three points of the petri dish. As a negative control, the fungus *F. oxysporum*. It was inoculated on a petri dish containing PDA medium only. The isolates were incubated for five days [10,11].

2.5 Antifungal volatile compounds test

Pure culture of *Fusarium oxysporum* Schltdl. (\pm 5 mm) and endophytic fungi (\pm 5 mm) were inoculated in the center of a petri dish containing PDA media separately. Both Petri dishes were cupped facing each other (*F. oxysporum*. fungus on top, endophytic fungus on bottom). As a negative control, the petri dish containing the fungus *Fusarium oxysporum* Schltdl. It was cupped with a petri dish containing PDA medium only. The fungus was incubated for six days [12].

2.6 Fungal identification

Molecular identification of fungal strains was completed by DNA amplification and sequencing of internal transcribed spacer (ITS) regions. Fungal mycelia (20 mg) were frozen overnight. Fungal DNA isolation was performed according to the manufacturer's protocol using the Wizard® Genomic DNA Purification Kit Protocol (Promega). Then, the isolated DNA was amplified by polymerase chain reaction (PCR). PCR was performed using GoTaq® Master Mix (Promega). The primers used, namely ITS 1 (TCCGTAGGTGAACCTGCGG) and ITS4 (TCCTCCGCTTATTGATGC) (references?), were mixed with the Master Mix Kit and DNA template with a total volume of 50 μ L. The mixture was then added into a thermal cycler using programmed PCR (BioRad, USA). The amplified fungal DNA (PCR products) was then sent to a commercial service for sequencing, and base sequences were compared using the BLAST Algorithm with publicly accessible databases, including GenBank.

3 Results and Discussion

The results of the isolation of endophytic fungi from corn plants (*Zea mays* L.) obtained 28 endophytic fungi, with 16 isolates from leaf organ parts, one isolate from stem organ parts, and 11 isolates from root organ parts. More endophytic fungi were found in the leaf organ. The isolation results of endophytic fungi were found more in the leaf organs [13]. The corn leaves used were old leaves, so the results of endophytic fungi isolation were more widely obtained in leaf organs. More endophytic fungal isolates were found in leaf organs, especially in older leaves than young leaves [14]. Fewer endophytic fungi were found in the stem organ than in the leaf organ because the epidermis layer of corn stems has a dense structure that functions as a protector called silica cells, making it difficult for endophytic fungi to penetrate plant tissue. The epidermis has a dense structure and can thicken because it contains silica to strengthen plant tissue structure and function as a protector [15].



Fig. 1. Isolation results of endophytic fungi in maize plants (Zea mays L.).

Eight-eight isolates are known to inhibit the growth of the pathogenic fungus *Fusarium* oxysporum Schltdl. Through the mechanism of nutrient competition, the diameter of the growth of endophytic fungi was more significant than the diameter of the fungus *Fusarium* oxysporum Schltdl. So that the growth of endophytic fungi can fill the media and suppress the growth of *Fusarium oxysporum* Schltdl fungus. In addition, there is an isolate with the code A3. 2. 1 isolated from the root organ of corn plants (*Zea mays* L.) has an antibiotic mechanism. Although the results showed that the pathogenic fungus was more significant in diameter than the endophytic fungus, it could still suppress the growth of pathogenic fungi grow slowly and do not fill the space. A clear zone is formed from the results of the inhibition. The mechanism of nutrient competition occurs due to the growth of pathogenic fungi is suppressed in the media space. In contrast, the antibiosis mechanism occurs due to the presence of antibiotic substances produced by endophytic fungi, which is characterized by the inhibition zone formed [16]—the formation of a precise zone results from forming an inhibition zone for pathogenic fungi.



Fig. 2. Antagonistic activity of the endophytic fungus from maize plants (Zea mays) against Fusarium oxysporum.

Seven isolates were Isolates that can inhibit pathogen growth by producing non-volatile compounds. In comparison, six other isolates cannot impede the development of F. oxysporum, in this case, proving that endophytic fungi have the effectiveness to inhibit the growth of pathogenic fungi by producing a specific metabolite. The efficacy of the inhibitory power of endophytic fungi against other pathogens is thought to be that the isolate produces metabolite compounds that have activity as antifungal [17]. The isolate with the highest percentage of inhibition is isolate B1. 1. 1, which amounted to 39.83% inhibition.



Fig. 3. Antifungal non-volatile compounds test- make the title more informative.

Antagonism activity test using the volatile compound production method aims to see whether endophytic fungi can produce a volatile compound that can inhibit the growth of pathogenic fungi. The results of the volatile compound production test showed that endophytic fungi could produce a volatile compound (metabolite) that could inhibit the growth of pathogenic fungi. Endophytic fungal activity inhibits the growth of pathogenic microbes through competition for space and nutrients and antibiosis by producing volatile and non-volatile compounds [18]. The volatile compound production test results measured the pathogen growth's diameter. Then, they calculated the percentage of inhibition resulting from the antagonism activity test between endophytic fungi and the pathogenic fungus *Fusarium oxysporum* Schltdl. Identification was conducted through ITS gene amplification, confirming that isolate D2.2.1 corresponds to *Trichoderma harzianum*.



Fig. 4. Antifungal volatile compounds test.

The endophytic fungus genus *Trichoderma harzianum* can inhibit the growth of the pathogenic fungus *Fusarium oxysporum* Schltdl. Because it antagonistically inhibits other pathogenic fungi by producing a volatile metabolite. T. Trichoderma harzianum can produce a volatile compound that can inhibit or kill other pathogenic fungi, so T. Trichoderma harzianum fungi include endophytic fungi with high antagonism [19]. T. Trichoderma harzianum produces volatile compounds such as 6-pentyl pyrrole, viri dins, harzianum A, haziarnic acid, gliotoxin, kininginins, cytosperone, and Trichoderma [20].

4 Conclusion

Endophytic fungi have been successfully isolated from maize plants. These fungi can inhibit the growth of pathogenic fungus *Fusarium oxysporum* through multiple mechanisms. Among all the fungi studied, *Trichoderma harzianum* exhibits the most potent inhibition and has the

Commented [Tz5]: ?

potential to be used as a biological control agent against the fungal pathogen *Fusarium* oxysporum.

Acknowledgment

The authors thanks to LPPM Universitas Ahmad Dahlan for supporting this research.

References

- Asniah, Lestari, D., Mariadi, dan Darlian, L. 2014. Potensi cendawan endofit nonpatogen asal akar tanaman cabai (Capsicum annuum L.) sebagai biofungisida patogen *Fusarium* oxysporum. AGRIPLUS, 24(2), pp.177-183.
- Soenartinigsih, Aqil, M., dan Andayani, N. N, 2016. Strategi pengendalian cendawan Fusarium sp. dan kontaminasi mikotoksin pada jagung. Iptek tanaman pangan, 11(1), pp.85-93.
- 3. Sudantha, I. M. dan Abadi, A., L, 2011. Uji efektivitas beberapa jenis jamur endofit Trichoderma spp. isolat lokal NTB terhadap jamur *Fusarium oxysporum* f. sp. vanillae penyebab penyakit busuk batang pada bibit vanili. Crop Agro, 4(2), pp.64-73.
- 4. Suriani dan Muis, A, 2016. Fusarium pada tanaman jagung dan pengendaliannya dengan memanfaatkan mikroba endofit. Iptek Tanaman Pangan, 11(2), pp.133-141.
- 5. Harni, R. dan Munif, A, 2012. Pemanfaatan agens hayati endofit untuk mengendalikan penyakit kuning pada tanaman lada. Buletin RISTRI, 3(3), pp.201-206.
- 6. Yurnaliza, Aryantha, I. N. P.A., Esyanti, R. R., and Susanto, A, 2014. Antagonistic Activity of Fungal Endophytes from Oil Palm Tissues Against Ganoderma boninense Pat. Plant Pathology Journal, 13(4), pp.257-267.
- 7. Nuraini, F. R., Setyaningsih, R, and Susilowati, A, 2017. Screening and characterization of endophytic fungi as antagonistic agents toward *Fusarium oxysporum* on eggplant (Solanum melongena). Biodiversitas, 18(4), pp.1377-1384.
- 8. Cappuccino, J. G. and Welsh, C. T, 2020. Microbiology: A Laboratory Manual. Twelfth Edition, Pearson Education: English.
- Landum, M. C., Felix, M. D. R., Alho, J., Garcia, R., Cabrita, M. J., Rei, F., and Varanda, C.M. R, 2016. Antagonistic activity of fungi of Olea europaea L. against Colletotrichum acutatum. Elseiver, 183, pp.100-108.
- Rabha, A. J., Naglot, A., Sharma, G. D., Gogoi, H. K., and Veer, V., 2014. In vitro, evaluation of antagonism of endophytic Colletotrichum gloeosporioides against potent fungal pathogens of Camellia sinensis. Indian J. Microbiol, 54(3), pp.302-309.
- 11. Hamzah, T. N. T., Lee, S. Y., Hidayat, A., Terhem, R., Hanum, I. R., and Mohamed, R, 2018. Diversity and characterization of endophytic fungi isolated from the tropical mangrove species, Rhizophora mucronata, and identification of potential antagonists against the soil-borne fungus, Fusarium solani. Frontiers In Microbiology, 9, pp.1-17.
- Katoch, M. and Pull, S, 2017. Endophytic fungi are associated with Monarda citriodora, an aromatic and medicinal plant, and their biocontrol potential. Pharmaceutical Biology, 55(1), pp.1528-1535.
- Zafitra, Elfina, Y., dan Ali, M. 2017. Uji antagonis jamur Trichoderma, Verticillium dan Torulomyces terhadap Ganoderma boninense Pat. Secara in vitro. JOM FAPERTA, 4(1): 1-6.

Commented [Tz6]: Penulisan referensi mengikuti template :

The authors, in the form: initials of the first names followed by last name (only the first letter capitalized with full stops after the initials),

- The title of the article
- The journal title (abbreviated),
- The volume number (bold type),
- The article number or the page numbers,
- The year of publication (in brackets),
- The DOI number (digital object identifier)

- Ramadhani, S. H., Samingan, dan Iswadi, 2017. Isolasi dan identifikasi jamur endofit pada daun jamblang (Syzygium cumini L.). Jurnal Ilmiah Mahasiswa Fakultas Keguruan dan Ilmu Pendidikan Unsyiah, 2(2), pp.77-90.
- 15. Malak, B, I, 2017. Identifikasi anatomi tumbuhan sirih hutan (Piper aduncum L.). Biolearing Journal, 8, pp.50-55.
- Izzatinnisa, Utami, U., dan Mujahidin, A, 2019. Uji antagonisme beberapa fungi endofit pada tanaman kentang terhadap *Fusarium oxysporum* secara in vitro. Jurnal Riset Biologi dan Aplikasinya, 2(1), pp.18-25.
- Sunariasih, N. P. L., Suada, I. K., dan Suniti, N. W, 2014. Identifikasi jamur endofit dari biji padi dan uji daya hambatnya terhadap Pyricularia oryzae Cav. Secara in vitro. E-Jurnal Agroekoteknologi Tropika, 3(2), pp.51-60.
- Vinale, F., Sivasithamparam, K., Ghisalberti, E. L., Woo, S. L., Nigro, M., Marra, R., ...Lorito, M. 2014. Trichoderma secondary metabolites are active on plants and fungal pathogens. The Open Mycology J., 8(Suppl-1, M5): 127–139.
- Dendang, B, 2015. Uji antagonisme Trichoderma spp. terhadap Ganoderma sp. yang menyerang tanaman sengon secara in-vitro. Jurnal Penelitian Kehutanan Wallacea. 4(2), pp.147-156.
- Harni, R., Amaria, W., Mahsunah, A. H., dan Lakani, I, 2019. Pengaruh metabolit sekunder Trichoderma spp. dan fungisida nabati untuk mengendalikan penyakit VSD pada tanaman kakao. Jurnal Tanaman Industri dan Penyegar, 6(3), pp.109-118.

HASIL REVIEW ARTIKEL KE-2 1 JULI 2024

COBERFInternational Conference of
Biological, Environment, Agriculture,
and Food (ICOBEAF) 2024



HOME ABOUT USER HOME SEARCH CURRENT ARCHIVES

Home > User > Author > Submissions > #15369 > **Review**

#15369 Review

SUMMARY REVIEW EDITING

Submission

Authors	Oktira Roka Aji, Dilla Rofiyanti 🕮
Title	Antagonistic Activity of Endophytic Fungi from Maize Plants (Zea mays L.) Against Fusarium oxysporum Schltdl.
Section	Articles
Editor	admin icobeaf 🖾

Peer Review

Round 1

Review Version	15369-36631-1-RV.DOCX 2024-05-17
Initiated	2024-07-01
Last modified	2024-07-01
Uploaded file	Reviewer A 15369-37723-1-RV.DOCX 2024-07-01

Editor Decision

Decision	Accept Submission 2024-07-02		
Notify Editor	🕮 🛛 Editor/Author Email Record 🔜 2024-07-02		
Editor Version	None		
Author Version	None		
Upload Author Version	Choose File No file chosen Upload		

Aims and Scope
Keynote Speaker
Editorial Team and Reviewer
Peer Review Process
Open Access Policy
Online Submission
Publisher, Partners and Indexing
Conference Fee
Contact

BIO Web of Conferences xx, xx ICoBEAF 2024

Antagonistic Activity of Endophytic Fungi from Maize Plants (*Zea mays* L.) Against *Fusarium oxysporum* Schltdl.

Abstract. Endophytic fungi were microorganisms that resided within plant tissues without harming their host plants. These fungi acted as biological agents against harmful fungi by utilizing mechanisms like competing for nutrients, parasitism, and antibiosis. This research aimed to isolate and identify the genus groups of endophytic fungi from maize plants (Zea mays L.) and understand the antagonistic mechanisms between endophytic fungi and Fusarium oxysporum Schltdl. Through dual culture tests, the highest inhibitory percentage of endophytic fungi against Fusarium oxysporum Schltdl is determined. in the filtrate and volatile compound tests. Antagonistic activity against pathogenic fungi was determined in vitro through dual culture methods, filtrate culture methods, and volatile compound production. Data obtained from the antagonistic activity of endophytic fungi were analyzed based on the calculation of inhibition percentages. The dual culture method revealed eight isolates effectively restraining the growth of Fusarium oxysporum Schltdl. Among the filtrate culture experiments, isolate B1.1.1 exhibited the most promising outcomes with an inhibition rate of 39.83%. Notably, isolate D2.2.1 displayed the highest inhibition rate of 44.38% in the antagonism test employing the volatile compound production method. Identification was conducted through ITS gene amplification, confirming that isolate D2.2.1 corresponded to Trichoderma harzianum. This research aimed to improve productivity and food security by managing plant diseases, thereby contributing to the achievement of SDG 2 (Zero Hunger).

1 Introduction

Plants are vulnerable to numerous diseases, and fusarium wilt is one of them. It typically affects the roots or the base of the plant stem. Fusarium wilt disease can be identified by the symptoms on the top of the plant [1]. The affected plants exhibit symptoms such as rotten roots and are prone to falling over, making them easy to uproot [2]. Fusarium wilt disease is caused by a pathogenic fungus called *Fusarium oxysporum* Schltdl. This soil-borne pathogen can survive in unfavorable conditions and even form chlamydospores without host plants [3]. The fungus is known for attacking various plants such as corn, sugarcane, rice, and sorghum. It causes rot in the stems, cobs, and seeds of corn [4].

Farmers typically rely on fungicides to manage plant diseases. However, the recurrent use of these chemicals can negatively impact human health and the environment due to the residual effects they leave behind. Consequently, there is a need to explore alternative eco-friendly approaches. One such strategy involves using endophytic fungi as biological agents to control diseases caused by plant pathogens [5]. This approach may offer a promising solution to the problem of fungicide overuse and its harmful consequences. By controlling

Commented [A1]: It is better to raise issues related to the SDGs at the beginning so that the role of this research on the SDGs is more visible

Commented [A2]: emphasize that this solution can replace the use of pesticides so that it does not give a double meaning (as if it were used to treat pesticides in plants) plant diseases, this research helps increase productivity and enhance food security, reducing crop losses and ensuring a more stable food supply. Consequently, this research is crucial for ensuring food security and mitigating global hunger, contributing to the achievement of SDG 2 (Zero Hunger).

2 Material and Methods

2.1 Sample preparation

Maize plant samples were collected from Pathuk District, Gunung Kidul. The samples were chosen due to the typical cultivation environment and relevance to the study's focus on endophytic fungi interactions with *Fusarium oxysporum*. The selected plants were approximately 80 days old, healthy, and devoid of deformities to ensure that the samples were at a developmental stage relevant to the study, free from potential confounding factors, and capable of providing reliable data for analysis. Samples were taken from the roots, stems, and leaves to isolate endophytic fungi.

2.2 Endophytic fungal isolation

The isolation of endophytic fungi was done using a direct seed-planting method. Maize plants' leaves, roots, and stems were washed with running water until they were utterly soil-free. Corn plant organ samples were sterilized by placing them in 70% (v/v) alcohol for 1 minute, then putting in a 0.5% (v/v) NaClO solution for 3 minutes, then placing them in 70% (v/v) alcohol for 30 seconds, and rinsing them with sterile distilled water twice and drying them on sterile Petri dishes [6,7].

Organ samples were cut using a sterile scalpel into 1-2 cm sizes as eptically. The organ samples were placed on PDA media and added with 100 µg/mL chloramphenicol as eptically. A total of 0.1 mL of sterile distilled water from the last rinse was poured on the PDA media by the spreading method to ensure that the surface sterilization process on the organ samples was successful. The culture was incubated at room temperature (25 - 28°C) for seven days [6,7].

The successfully isolated fungi were then subcultured on PDA media and incubated for three days at room temperature (25 - 28°C). Macroscopic observations were made by observing the morphology of the fungal mycelium, which included the color of the upper surface colony, the color of the reverse colony, and the texture of the colony surface. Microscopic observations were carried out with a pure culture of endophytic fungi isolated aseptically using a sterile ose, transferred to the surface of a glass object, and then dripped with Lactophenol cotton blue. The preparations were observed with a light microscope. Observations include the structure of hyphae (concentrated or non-concentrated) and the structure of spores [8].

2.3 Dual Culture Assay

To carry out an antagonism test, small pieces of pure culture of the fungus *Fusarium* oxysporum Schltdl. (\pm 5 mm) and each endophytic fungal isolate (\pm 5 mm) was inoculated in one petri dish containing PDA media at a distance of 3 cm. As a negative control, small pieces of *F. oxysporum*. (\pm 5 mm) was planted in the center of the Petri dish containing PDA media. Isolates were incubated for seven days [9].

2.4 Antifungal non-volatile compounds test

Pure cultures of endophytic fungi were inoculated in 10 mL of PDB media. The endophytic fungal culture was incubated for ten days at room temperature (25-28°C). The fungal culture was filtered using filter paper, and the filtrate was filtered again using a 0.22 μ m sterile syringe filter. The cell-free filtrate was mixed with sterile PDA media in petri dishes. Pieces of fungal culture of *Fusarium oxysporum* Schltdl. (± 5 mm) were inoculated at three points of the petri dish. As a negative control, the fungus *F. oxysporum*. It was inoculated on a petri dish containing PDA medium only. The isolates were incubated for five days [10,11].

2.5 Antifungal volatile compounds test

Pure culture of *Fusarium oxysporum* Schltdl. (\pm 5 mm) and endophytic fungi (\pm 5 mm) were inoculated in the center of a petri dish containing PDA media separately. Both Petri dishes were cupped facing each other (*F. oxysporum*. fungus on top, endophytic fungus on bottom). As a negative control, the petri dish containing the fungus *Fusarium oxysporum* Schltdl. It was cupped with a petri dish containing PDA medium only. The fungus was incubated for six days [12]. The percentage of inhibition in the antifungal non-volatile compounds test and antifungal volatile compounds test were measured using the formula:

Inhibition (%) =
$$\left(\frac{C-T}{T}\right) \times 100$$
 (1)

Where:

C = Diameter of the fungal colony in the control plate

T = Diameter of the fungal colony in the treated plate

2.6 Fungal identification

Molecular identification of fungal strains was completed by DNA amplification and sequencing of internal transcribed spacer (ITS) regions. Fungal mycelia (20 mg) were frozen overnight. Fungal DNA isolation was performed according to the manufacturer's protocol using the Wizard® Genomic DNA Purification Kit Protocol (Promega). Then, the isolated DNA was amplified by polymerase chain reaction (PCR). PCR was performed using GoTaq® Master Mix (Promega). The primers used, namely ITS 1 (TCCGTAGGTGAACCTGCGG) and ITS4 (TCCTCCGCTTATTGATGC) (references?), were mixed with the Master Mix Kit and DNA template with a total volume of 50 μ L. The mixture was then added into a thermal cycler using programmed PCR (BioRad, USA). The amplified fungal DNA (PCR products) was then sent to a commercial service for sequencing, and base sequences were compared using the BLAST Algorithm with publicly accessible databases, including GenBank.

3 Results and Discussion

The results of the isolation of endophytic fungi from corn plants (*Zea mays* L.) obtained 28 endophytic fungi, with 16 isolates from leaf organ parts, one isolate from stem organ parts, and 11 isolates from root organ parts. Fig. 1 shows the isolation results of endophytic fungi from maize plants (*Zea mays* L.), highlighting the diversity and distribution of fungal isolates obtained. More endophytic fungi were found in the leaf organ. The isolation results of endophytic fungi were old

leaves, so the results of endophytic fungi isolation were more widely obtained in leaf organs. More endophytic fungal isolates were found in leaf organs, especially in older leaves than young leaves [14]. Fewer endophytic fungi were found in the stem organ than in the leaf organ because the epidermis layer of corn stems has a dense structure that functions as a protector called silica cells, making it difficult for endophytic fungi to penetrate plant tissue. The epidermis has a dense structure and can thicken because it contains silica to strengthen plant tissue structure and function as a protector [15]. Eight-eight isolates are known to inhibit the growth of the pathogenic fungus *Fusarium oxysporum* Schltdl. Through the mechanism of nutrient competition, the diameter of the development of endophytic fungi was more significant than the diameter of the fungus *Fusarium oxysporum* Schltdl so that the growth of endophytic fungi can fill the media and suppress the growth of *Fusarium oxysporum* Schltdl fungus.



Fig. 1. Isolation results of endophytic fungi in maize plants (Zea mays L.).



Fig. 2. Antagonistic activity of the endophytic fungus from maize plants (*Zea mays* L.) against *Fusarium oxysporum* Schltdl. (a.) Control (b.) isolate A3.2.1 and (c.) isolate A2.1.1

An isolate with code A3.2.1 isolated from the root organ of corn plants (*Zea mays* L.) has an antibiotic mechanism. Although the results showed that the pathogenic fungus was more significant in diameter than the endophytic fungus, it could still suppress the growth of pathogenic fungi because pathogenic fungi grow slowly and do not fill the space. A clear zone is formed from the results of the inhibition. The mechanism of nutrient competition occurs due to the growth of endophytic fungal mycelium that fills the media and pathogenic fungi so that the development of pathogenic fungi is suppressed in the media space. In contrast, the antibiosis mechanism occurs due to the antibiotic substances produced by endophytic fungi, characterized by the inhibition zone formed [16]. Fig. 2 shows the antagonistic activity of endophytic fungi from maize plants (*Zea mays* L.) against *Fusarium oxysporum* Schltdl.

Seven isolates were isolates that can inhibit pathogen growth by producing non-volatile compounds. In comparison, six other isolates cannot impede the development of F. *oxysporum*, in this case, proving that endophytic fungi have the effectiveness to inhibit the

growth of pathogenic fungi by producing a specific metabolite. The efficacy of the inhibitory activity of endophytic fungi against other pathogens is thought to be that the isolate produces metabolite compounds that have activity as antifungal [17]. Table 1 shows the percentage inhibition of *Fusarium oxysporum* by endophytic fungi, evaluated using antifungal, nonvolatile, and volatile compounds tests. Fig 3 shows the results of the antifungal non-volatile compounds test, illustrating the inhibition of *Fusarium oxysporum* growth by endophytic fungal isolates. The isolate with the highest percentage of inhibition is isolate B1.1.1, which amounted to 39.83% inhibition.



Fig. 3. Antifungal non-volatile compounds test. (a.) Control (b.) isolate B1.1.1 and (c.) isolate D2.2.1.

 Table 1. Percentage inhibition of Fusarium oxysporum by endophytic fungi using antifungal non-volatile and volatile compounds tests

No	Isolate	Inhibitior	ı (%)
INO	No Isolate	Antifungal non-volatile compounds test	Antifungal volatile compounds test
1	A1. 2. 1	$9,72 \pm 1,37$	25.29 ± 16.92
2	A2. 1. 1	n.d.	43.40 ± 7.95
3	A2. 2. 1	n.d.	30.72 ± 12.02
4	A3. 2. 1	$30,74 \pm 6,05$	18.04 ± 16.08
5	A3. 2. 2	n.d.	33.46 ± 2.96
6	B1. 1. 1	$39{,}83\pm2{,}58$	39.93 ± 15.62
7	D2. 1. 1	n.d.	19.41 ± 8.60
8	D2. 1. 2	[8,79 ± 6,22]	41.31 ± 8.13
9	D2. 2. 1	n.d.	44.38 ± 9.34
10	D3. 1. 1	n.d.	19.41 ± 8.60
11	D3. 1. 2	n.d.	21.50 ± 8.41
12	D3. 2. 1	n.d.	18.30 ± 10.17
13	D3. 2. 2	n.d.	17.45 ± 5.82

Figure 4. shows the inhibitory effect of endophytic fungi on the growth of *Fusarium* oxysporum, measured by the reduced diameter of the fungal colony. Antagonism activity test using the volatile compound production method aims to see whether endophytic fungi can produce a volatile compound that can inhibit the growth of pathogenic fungi. The results of the volatile compound production test showed that endophytic fungi could produce a volatile compound (metabolite) that could inhibit the growth of pathogenic fungi. Endophytic fungi inhibit the growth of pathogenic microbes through competition for space and nutrients and by producing volatile and non-volatile compounds (antibiosis) [18]. The diameter of pathogen growth was measured in the volatile compound production test. The percentage of growth inhibition was then calculated based on the antagonistic activity between endophytic

Commented [A3]: Decimal formating

fungi and the pathogenic fungus *Fusarium oxysporum* Schltdl. Figure 5 shows the electrophoresis results depicting the amplified PCR products of the ITS gene, demonstrating bands at approximately 600 base pairs in length. Identification through ITS gene amplification confirmed that isolate D2.2.1 is *Trichoderma harzianum*, with a similarity score of 97.30% based on BLAST results.



Fig. 4. Antifungal volatile compounds test. (a.) Control (b.) isolate D2.2.1. The red circle indicates the diameter of the fungal growth of *Fusarium oxysporum* Schltdl.



Fig. 5. Electrophoresis Results of PCR Products for ITS Gene (600 bp).

The endophytic fungus genus *Trichoderma harzianum* can inhibit the growth of the pathogenic fungus *Fusarium oxysporum* Schltdl. Because it antagonistically inhibits other pathogenic fungi by producing a volatile metabolite. *Trichoderma harzianum* can make a volatile compound that can inhibit or kill other pathogenic fungi, so *Trichoderma harzianum* fungi include endophytic fungi with high antagonism [19]. *Trichoderma harzianum* produces volatile compounds such as 6-pentyl pyrrole, viri dins, haziarnic acid, gliotoxin, kininginins, and cytosperone [20].

4 Conclusion

Endophytic fungi successfully isolated from maize plants. These fungi can inhibit the growth of the pathogenic fungus *Fusarium oxysporum* Schltdl through multiple mechanisms. Among all the fungi studied, *Trichoderma harzianum* exhibits the most potent inhibition and has the potential to be used as a biological control agent against the fungal pathogen *Fusarium oxysporum* Schltdl.

Acknowledgment

The authors thank LPPM Universitas Ahmad Dahlan for supporting this research.

Commented [A4]: low Image quality, the resolution needs to be increased

References

- L. Asniah, D. Lestari, Mariadi, L. Darlian, Potensi cendawan endofit nonpatogen asal akar tanaman cabai (*Capsicum annuum* L.) sebagai biofungisida patogen *Fusarium* oxysporum. AGRIPLUS. 24, 177-183 (2014)
- Soenartinigsih, M. Aqil, N. N. Andayani, Strategi pengendalian cendawan *Fusarium* sp. dan kontaminasi mikotoksin pada jagung. Iptek Tanam. Pangan. 11, 85-93 (2016)
- 3. I. M. Sudantha, A. L. Abadi, Uji efektivitas beberapa jenis jamur endofit *Trichoderma* spp. isolat lokal NTB terhadap jamur *Fusarium oxysporum* f. sp. vanillae penyebab penyakit busuk batang pada bibit vanili. Crop Agro. **4**, 64-73 (2011)
- 4. Suriani, A. Muis, Fusarium pada tanaman jagung dan pengendaliannya dengan memanfaatkan mikroba endofit. Iptek Tanam. Pangan. **11**, 133-141 (2016)
- 5. Harni, A. Munif, Pemanfaatan agens hayati endofit untuk mengendalikan penyakit kuning pada tanaman lada. Buletin RISTRI . **3**, 201-206 (2012)
- Yurnaliza, I. N. P. A. Aryantha, R. R. Esyanti, A. Susanto, Antagonistic Activity of Fungal Endophytes from Oil Palm Tissues Against *Ganoderma boninense* Pat. Plant Pathol. J. 13, 257-267 (2014). https://doi.org/10.3923/ppi.2014.257.267
- F. R. Nuraini, R. Setyaningsih, A. Susilowati, Screening and characterization of endophytic fungi as antagonistic agents toward *Fusarium oxysporum* on eggplant (*Solanum melongena*). Biodiversitas. 18, 1377-1384 (2017) https://doi.org/10.13057/biodiv/d180413
- 8. J. G. Cappuccino, C. T. Welsh, Microbiology: A Laboratory Manual. Pearson Education: English, Twelfth Edition (2020)
- M. C. Landum, M. D. R. Felix, J. Alho, R. Garcia, M. J. Cabrita, F. Rei, C. M. R. Varanda, Antagonistic activity of fungi of *Olea europaea* L. against *Colletotrichum acutatum*. Elseiver. **183**, 100-108 (2016) https://doi.org/10.1016/j.micres.2015.12.001
- A. J. Rabha, A. Naglot, G. D. Sharma, H. K. Gogoi, V. Veer, In vitro evaluation of antagonism of endophytic *Colletotrichum gloeosporioides* against potent fungal pathogens of *Camellia sinensis*. Indian J. Microbiol. 54, 302-309 (2014) <u>https://doi.org/10.1007/s12088-014-0458-8</u>
- 11. T. N. T. Hamzah, S. Y. Lee, A. Hidayat, R. Terhem, I. R. Hanum, R. Mohamed, Diversity and characterization of endophytic fungi isolated from the tropical mangrove species, *Rhizophora mucronata*, and identification of potential antagonists against the soil-borne fungus, *Fusarium solani*. Front. Microbiol. 9, 1-17 (2018) https://doi.org/10.3389/fmicb.2018.01707
- M. Katoch, S. Pull, Endophytic fungi are associated with *Monarda citriodora*, an aromatic and medicinal plant, and their biocontrol potential. Pharm. Biol. 55, 1528-1535 (2017)

https://doi.org/10.1080/13880209.2017.1309054

 Zafitra, Y. Elfina, M. Ali, Uji antagonis jamur Trichoderma, Verticillium dan Torulomyces terhadap *Ganoderma boninense* Pat. Secara in vitro. JOM FAPERTA. 4, 1-6 (2017)

Commented [A5]: need more international journals as references

- S. H. Ramadhani, Samingan, Iswadi, Isolasi dan identifikasi jamur endofit pada daun jamblang (*Syzygium cumini* L.). Jurnal Ilmiah Mahasiswa FKIP Unsyiah. 2, 77-90 (2017)
- B. I. Malak, Identifikasi anatomi tumbuhan sirih hutan (*Piper aduncum* L.). Biolearing J. 8, 50-55 (2017)
- Izzatinnisa, U. Utami, A. Mujahidin, Uji antagonisme beberapa fungi endofit pada tanaman kentang terhadap *Fusarium oxysporum* secara in vitro. J. Riset Biologi Aplikasinya. 2, 18-25 (2019) <u>https://doi.org/10.26740/jrba.v2n1.p18-25</u>
- N. P. L. Sunariasih, I. K. Suada, N. W. Suniti, Identifikasi jamur endofit dari biji padi dan uji daya hambatnya terhadap *Pyricularia oryzae* Cav. secara in vitro. E-Jurnal Agroekoteknologi Tropika. 3, 51-60 (2014)
- F. Vinale, K. Sivasithamparam, E. L. Ghisalberti, S. L. Woo, M. Nigro, R. Marra, M. Lorito, Trichoderma secondary metabolites are active on plants and fungal pathogens Open Mycol. J. 8, (Suppl-1, M5) 127-139 (2014) https://doi.org/10.2174/1874437001408010127
- B. Dendang, Uji antagonisme *Trichoderma* spp. terhadap *Ganoderma* sp. yang menyerang tanaman sengon secara in-vitro. J. Penelitian Kehutanan Wallacea. 4, 147-156 (2015)

https://doi.org/10.18330/jwallacea.2015.vol4iss2pp147-156

 R. Harni, W. Amaria, A. H. Mahsunah, I. Lakani, Pengaruh metabolit sekunder *Trichoderma* spp. dan fungisida nabati untuk mengendalikan penyakit VSD pada tanaman kakao. J. Tanaman Ind. Penyegar. 6, 109-118 (2019) <u>https://doi.org/10.21082/jtidp.v6n3.2019.p109-118</u> EMAIL Letter of Acceptance (LOA) 2 JULI 2024 Editor 2024-07-02 01:52 PM Subject: [ICoBEAF] Editor Decision

Oktira Roka Aji:

Congratulations! Your submission entitled "Antagonistic Activity of Endophytic Fungi from Maize Plants (Zea mays L.) Against Fusarium oxysporum Schltdl." has been accepted for presentation at the INTERNATIONAL CONFERENCE OF BIOLOGICAL, ENVIRONMENT, AGRICULTURE, AND FOOD, which will be held on July 16-17, 2024.

Please complete the following administrative steps for your presentation article. One author must register as a full participant to present this article. Please follow these instructions:

1. Pay the presenter and publication fee (early bird rate applies if you register within 7 days of acceptance) to BTPN bank, account number 90310414509 (account holder: Fitri Rilivo Kristine).

a. Presenter early bird + publication: \$160 | IDR Rp. 2.450.000

b. Presenter regular + publication: \$170 | IDR Rp. 2.750.000

2. Fill in the registration form and send your PowerPoint presentation along with evidence of payment to https://s.uad.id/icobeafpresenter.

3. Follow the comments of the editors and reviewers, and upload your revised paper to the ICoBEAF Submission system before July 14, 2024.

4. The committee will send further information about the presentation after you register as a presenter.

Please note the following ICoBEAF procedures:

1. Non-presented papers will not be considered for publication in the BIO Web of Conferences.

2. Each author is allowed a maximum of two papers as an author/co-author in the ICoBEAF conference.

We would like to thank you for your contribution to ICoBEAF 2024. Our conference will provide you with a forum to share your specialized research with international colleagues. If you need further information, please contact us by email at icobeaf@uad.ac.id or visit our website https://adis.uad.ac.id/ad-icobeaf/for the latest updates on the conference.

Best regards,

Assist. Prof. Much Fuad Saifuddin General Chair of ICoBEAF 2024

sender by admin icobeaf icobeaf@uad.ac.id

INTERNATIONAL CONFERENCE ON BIOLOGICAL, ENVIRONMENT, AGRICULTURE, AND FOOD http://seminar.uad.ac.id/index.php/icobeaf

Close