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# Antagonistic evaluation of *Trichodermaviride* and Aspergillus flavus against wood-decay fungus **Pleurotusostreatus**

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Abstract. Pleurotusostreatus cultivation is often limited by the presence of antagonistic microorganisms that can also grow together on a variety of substrates that are used as growth media of *P. ostreatus*. As a preliminary study, this research aims to evaluate the antagonism of Trichodermaviride and Aspergillus flavus in interacting with the P.ostreatus fungus by invitro. An antagonism test using a dual culture method of these fungi with 1 month incubation time on Poteto Dextrose Agar (PDA) media was performed. Evaluation of the antagonistic properties of these fungi is conducted through macroscopic and microscopic observations. Macroscopically, the growth of T. viride and A. flavus mycelium showed that both colonies were more dominant than P. ostreatus colonies. Microscopically, hyphae of T. viride and A. *flavus* indicate the presence of very strong micoparasitic properties that characterized by attachment and convolution of both hyphae to P. ostreatus hyphae. It was concluded thatthe growth of T. viride and A. flavus fungi was more dominant and had potential as an inhibiting agent of P. ostreatus fungal growth, respectively, in dual culture on PDA media.

#### 1. Introduction

Pleurotusostreatus, has long been recognized as one of the most widely used wood fungi (mushroom) for food because of its high levels of protein, minerals and vitamins [1]. Naturally, this fungus is not only grows on dead wood but also easily cultivated on various agricultural industry and residual material substrates, and other agriculture-cellulosic materials [2] because they produce oxidative enzymes [laccase and manganese peroxidase (MnP)] and hydrolytic (cellulase, xylanase and tannase) [3]. The use of different substrates is able to effect on biological and economic yield of *P. ostreatus* (oyster mushroom) cultivation [4].

However, it is common to find problems in the cultivation of the *P. ostreatus* mushroom, one of which is the existence of contamination on the growth substrate media which regularly found, so that variations of heat treatment are required to eliminate the growth of other fungi such as Trichoderma sp., Altemaria sp., Aspergillus sp., Fusariumsp., Monilia sp., Mucor sp., Rhizopus sp. etc.[5]. The growth of these other fungi as a competitor often leads to impaired growth of *P. ostreatus* so that it can reduce the production of its cultivation.

In the same habitat, it is well known that mushrooms have mutually supportive relationships and inhibit the growth of one another in obtaining nutritional sources for its survival. Schmidt[6] notes that

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the inhibiting growth relationship or better known as the antagonism (competitive reciprocal effect) is based on production of toxic metabolites, on mycoparasitism, and on nutrient competition.

In the effort of handling *P. ostreatus* growth disorders, the initial information required on how the characteristics of interdependent or antagonistic relationship between *P. ostreatus* with competitors in a substrate. Therefore, antagonistic studies of fungi were performed using two species of antagonistic fungi, *Trichodermaviride* and *Aspergillusflavus* to find out the interaction pattern of each of these fungi in inhibiting the growth of *P. ostreatus* fungus by in-vitro.

#### 2. Materials and methods

#### 2.1. Fungal isolates and culture media

The fungal isolates used were *Trichodermaviride*, *Aspergillusflavus* and *Pleurotusostreatus* stored in the collecting of the Forest Protection Laboratory of Faculty of Forestry, Mulawarman University, Samarinda.

Potato Dextrose Agar (PDA) material is made aseptically in laboratory following the procedure ofHiMedia[7]. PDA is used as a culture media for fungal growth of *P. ostreatus*, and antagonistic test as well.

## 2.2. Antagonistic test

An antagonistic test using a dual culture method of Bruce and Highley[8] on PDA media was performed on *T. viride* and *A. Flavus* against *P. ostreatus*, respectively, with an incubation period of 1 month.

Materials and equipment used in the antagonistic test were conducted aseptically in Laboratory of Wood Biology, Faculty of Forestry, Mulawarman University, Samarinda.

#### 2.3. Observations

Interaction of two fungal colonies in vitrodual culture on PDA media wasexaminedon macro-and microscopic observations.

2.3.1. *Macroscopic observation*.In macroscopic observation,NIKON SMZ645 Stereoscopic Microscope at 10-30x magnification was used to evaluate mycelium growth and the contact limit between the two fungal colonies growing on the PDA media plates.

2.3.2. *Microscopic observation*. By using NIKON Eclipse E400 Light Microscope at 40-400x magnification, microscopic observation was made on the growth and interaction of the two fungal hyphae growing on the PDA media plates.

#### 3. Results and discussion

#### 3.1. Growth inhibition of Pleurotusostreatus by Trichodermaviride

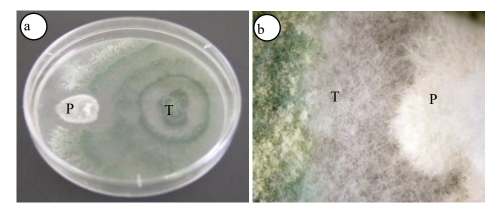
Macroscopically, in the dual culture method, the growth of both fungi in one Petri dish at 1-2 weeks incubation showed *Tricodermaviride* colony with mycelium and greenish spores more dominant than *Pleurotusostreatus* colony with whitish mycelium (Figure 1a).

Since the beginning contact, *T. viride* colony dominates almost thoroughly the surface area of PDA media. On the contrary, the growth of *P. ostreatus* mycelium with very limited growth area, cover the surface of PDA media, even tend to grow in the opposite direction from the contact zone. In the contact zone, there is no clear boundary separating between the two fungal colonies (Figure 1b).

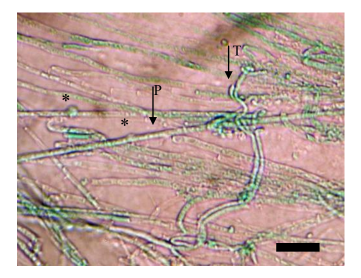
The dominance of *T. viride* colonies is not only from the fungal growth fully covering the surface of the PDA media but also the fungus grows on top of the *P. ostreatus* mycelium (Figure 1). This condition shows *T. viride* plays a role micoparasite[9] by utilizing the mycelium of *P. ostreatus* as a

source of nutrients other than obtained from the media. It is certain that with the dominant colonization, *T. viride* is antagonistic fungus to *P. ostreatus*.

The interaction stage of both colonies can also be shown by microscopic observations (Figure 2) in which appears to be attachment and convolution of *T. viride* hyphae on the *P. ostreatus* hyphae. The attachment is an early stage of the convolution process, which, according to Klein Dand Eveleigh[10], is a frequent response between the *Trichoderma* spp. with other fungi in case of contact.



**Figure 1.**The dominant growth of the *Tricodermaviride* colony(T) against *Pleurotusostreatus* (P) on PDA dual culture media: (a) The overall surface of the media in Petri dishes; (b) At the contact boundary of the colonies' encounter [28 x magnifications].



**Figure 2.**Attachment and convolution of *Tricodermaviride* hyphae (T) on *Pleurotusostreatus* hyphae (P) on dual culture of PDA media. Sign \* denotes clamp connection of *P. ostreatus* hyphae, which is known as main characteristic of basidiomycetefungi [Bar =  $20 \mu m$ ].

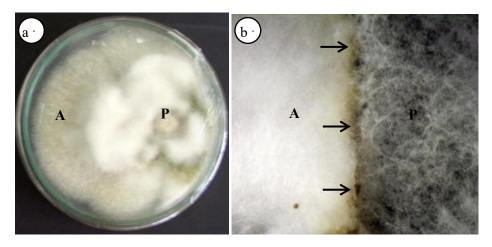
Indication of the micoparasite process of T. *viride* as an antagonist fungus, according to Ranasingh *et al*[9], it is usually followed by releasing process of various enzymes such as chitinase, glucanase and pectinase.

Basically, the antagonistic properties are due to the ability of *T. viride* to produce extracellular enzymes capable of metabolizeon polycyclic aromatic hydrocarbons (PAHs) such as naphthalene, phenanthrene, chrysene, pyrene, and benzo[a]pyrene [11], which is indicate his dominating ability and

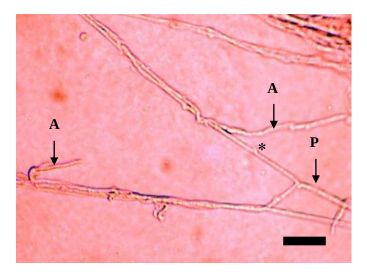
even the destructive structure of the *P. ostreatus* mycelium. Such event was also observed in dominating of other Tricoderma species, i.e. *T. citrinoviride*, *T. harzianum and T. Atroviride* associated with the *Camptomyia* pestagainst the mycelial growth of ashiitake mushroom, *Lentinulaedodes* [12].

## 3.2. Growth Inhibition of Pleurotusostreatus by Aspergillusflavus

The *Aspergillusflavus* mycelial colony also showed growth dominance fulfilling the PDA media against *Pleurotusostreatus* (Figure 3a). The condition also shows that the *P. ostreatus* mycelium could not develop as it could grow in mono-culture media.



**Figure 3.***Aspergillusflavus* colony (A) Dominates the surface of the media and covers the mycelium of *Pleurotusostreatus* (P) on PDA dual culture media method: (a) The overall surface of the media in Petri dishes; (b) At the contact boundary of the colonies (arrows) [28 x magnifications].



**Figure 4.**Attachment and convolution of *Aspergillusflavus* hyphae (A) to the *Pleurotusostreatus* hyphae (P) on dual culture of PDA media Sign \* denotes clamp connection of *P. ostreatus* hyphae, which is known as main characteristic of basidiomycete fungi [Bar =  $20 \mu m$ ].

Covering of PDA media by *A. flavus* mycelium was also characterized by presence of a limiting zone that forms like a long line span on the second encounter of mycelium (Figure 3b). The properties

of *A. flavus*micoparasite on dual-culture media were demonstrated by exploiting *P. ostreatus*mycelium as a source of nutrients beside those that obtained from PDA. This could be indicative that for the fact that *A. flavus* has in common with *T. viride*, which acts as an antagonist fungus against *P. ostreatus*.

Microscopically, the antagonistic features are seen that *A. flavus* hyphae also attached and convoluted *P. ostreatus* hyphae (Figure 4). The antagonistic ability of *A. flavus* hyphae is related to the extracellular xylanase enzymes that released [13], which may play a role as a metabolic toxin [5] and also has the greatest cellulolytic ability [14] against *P. ostreatus* hyphaeso that its growth in PDA media becomes inhibited. Meanwhile, aflatoxinB<sub>1</sub> (AFB<sub>1</sub>)as a secondary metabolite that produced by several members of Aspergillus fungi could be degraded by ligninolytic enzymes of *P. ostreatus* strains [15,16].

#### 4. Conclusions

Based on results of this study, we conclude that *Trichodermaviride* and *Apergillusflavus* mycelial colonies dominate the *Pleurotusostreatus* mycelia colony in utilizing nutrient in dual culture of PDA media. Both of the antagonistic fungi are very powerful microparasite against *P. ostreatus* hyphae, therefore the prevention inhibition of theirgrowth in media of *P. ostreatus* cultivation must be undertaken early.

#### Acknowledgements

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