Selection of stain fungi on rubberwood (Hevea brasiliensis) and its growth response against chitosan

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Introduction
Rubberwood (Hevea brasiliensis Muell.Arg.) is one of the hardwood species that is native from tropical rain forest, Brazil. Rubberwood firstly planted in Indonesia and Malaysia for rubber sap production. The wood has bright color, medium brown streak, the average of density in 0.6 g/cm³ modulus of rupture (MOR) in 103 MPa, modulus of elasticity (MOE) in 9900 MPa, tensile strength in 1.4-1.9 MPa, and hardness in 6000 N. Therefore, rubberwood widely used as furniture, particle wood, wood floors, plywood and sawn timber (Riyaphan et al. 2015). On the other hand, rubberwood has low natural durability that makes it susceptible to attack by stain fungi as mold or blue stain fungi. Stain fungi is filamentous fungi that cause discoloration on wood surface like blue, greyish, green and black. The colorization influenced by melanin crystallization around hyphae or from secretion of extracellular material. Stain fungi do not affect of strength loss, but decrease the aesthetic and cost of wood. Previous studies reported that three stain fungi were identified from rubberwood, which were Aspergillus niger, Aspergillus flavus, and Penicillium citrinum (Olderttroen et al. 2016). In fact, stain fungi were found to attack the part of piano produced by one company in Indonesia.

Many case wood that attacked by stain fungi became the main reason of some industry to preserving wood with chemicals compound. However, most of chemicals are harmful to humans and environment. Wilson et al. (2014) reported that 80% residue of fungicide azoxystrobin and 71% boscalid found in fly larva which is not as main target. Furthermore, other results report that methylene dithiocyanate as an active compound of fungicide can also cause respiratory disorder and death. Based on these negative impacts, many studies conducted to find preservatives that are safe for humans and environment. One of the natural materials that can be used as wood preservatives is chitosan.

Chitosan is unbranched polymer that consists of glucosamine (GlcN) and N-acetyl glucosamine (GlcNac). The polymer has the potency to be used as wood preservative considering its properties, such as biodegradable, non-toxic, abundant, antimicrobial, and antifungal activity (Zivanovic et al. 2015). Some studies are reporting the activity of chitosan inhibited stain fungi. Waewthongrak et al. (2015) confirmed that chitosan can inhibit the growth of Penicillium digitatum. The addition of chitosan concentration 1 and 5 mg mL⁻¹ in potato dextrose agar (PDA) medium showed 100% inhibition rate, while the diameter colony of control was 20.5 mm after five days incubation. On the other hand, chitosan only gives 40% spore viability of Aspergillus parasiticus, whereas control with 100% spore viability (Hernández-téllez et al. 2017).
More recently, Da Silva et al. (2018) reported that chitosan inhibited the growth of Aspergillus flavus. The sample showed only 35.34 ± 4.04% infection at chitosan concentration 2 g L⁻¹, while control has the highest infection 100% after seven days incubation. Several studies reported that chitosan concentration 0.01% caused damage in plasma membrane and cytoplasm material, whereas the higher concentration of chitosan (0.15%) caused higher destruction of membrane structure and complete loss of cytoplasm integrity of Sphaeropsis sapinea (Singh et al. 2008). This was also reported by Divya et al. (2018) that chitosan showed damage to membrane permeability and abnormality of cell Rhizoctonia solani, Fusarium oxysporum, Colletotrichum acutatum, and Phytophthora infestans. Since there is no research about isolate and identify stain fungi from rubberwood and in vitro assay of chitosan against stain fungi in Indonesia, the aim of present study was to identify stain fungi on seasoned rubberwood and determine the antifungal activity of chitosan.

MATERIALS AND METHODS

Isolation and selection of fungi
Rubberwood sample that infected by stain fungi were obtained from the one company in Indonesia. Wood samples were cut in 2×2×1 cm and soaked in aquadest for 24 hours and then treated with surface sterilization based on Hughes et al. (2018) with modification. The wood samples were soaked in sterile aquadest for two minutes and shaken occasionally. This process was carried out in five times and then passed on fire slightly (Sudirman 2018, pers. com). The woods were incubated in PDA medium for seven days at room temperature (28±2°C). The tip of hyphae was taken and cultivated in PDA for purification.

The purified isolates were then selected to observe percentage growth of wood test samples. Wood test samples were prepared with size 5×1×0.5 cm and sterilized in autoclave for 121°C for 15 minutes. Sterile wood test samples and each isolate stain fungus were put in Petri dish (ø = 9 cm) and incubated for six weeks incubation at room temperature (28±2°C) based on European Standard 152 (1984) with modification. The wood test samples were removed after six weeks and dried up at 60°C for three days. The percentage growth of isolate to the surface area of the wood test samples was represented by score. Score 0 indicated that there is no growth of isolate on wood test samples and scores 5 indicated the heavy stain fungi growth, which was 0: no growth, 1: 20% growth of stain fungi on wood test samples, 2: 40%, 3: 60%, 4: 80% and 5: 100% (Jantamas et al. 2013).

Morphological identification of stain fungi
Morphological identification was carried out on isolates that had the highest score on wood test samples. The isolates were cultivated in the center of malt extract agar (MEA) medium at room temperature (28±2°C) for macroscopic analysis based on Silva et al. (2011). Microscopic analysis was conducted by cultivated the isolates in three points on MEA for seven days at room temperature (28±2°C). Macroscopic analysis includes colony diameter, colony color of the top, and bottom, while microscopic analysis includes conidia diameter, conidia shape, hyphae, present of metulae, and size of vesicle (Samson et al. 2014).

Bioactivity test of chitosan against stain fungi
Chitosan was obtained from Aquatic Product Technology Laboratory, Faculty of Fisheries and Marine Sciences, Bogor Agricultural University, Bogor, Indonesia. Chitosan powder from shrimp shells was dissolved in acetic acid 2% (w/v) to make concentration 0, 0.5, 1, and 2%. The solution was then blended to get a good mixture.

The effect of chitosan on mycelial growth was evaluated in solid medium. PDA medium and chitosan solution were sterilized for 15 minutes at 121°C, separately. Sterile PDA medium (15 mL) was poured into Petri dish and mixed with 1 mL chitosan solution with each concentration. PDA medium mixed with acetic acid 2% was prepared as a control (Dewi and Nur 2017). Inhibitory effect was conducted by inoculating a 0.5 cm diameter isolate in PDA and incubated for seven days at room temperature (28±2°C). The experiments were repeated in three replicates. Determination of the percentage of growth inhibition is based on previous research with formulations (Pratheesh et al. 2017):
\[ \text{PIDG} = \frac{D_1 - D_2}{D_1} \times 100 \]

Where:
- PIDG: Percentage of Inhibition of Diameter Growth (%)
- D1: Colony diameter in control plates (cm)
- D2: Colony diameter in test plates (cm)

The experiments were analyzed by one-way analysis of variance (ANOVA) with four chitosan treatments, which were: [P1]: 2%, [P2]: 1%, [P3]: 0.5% and [P4]: control with acetic acid 2%. The difference of mean was determined by Duncan based on 95% confidence interval using SPSS software version 22.

**RESULTS AND DISCUSSION**

**Stain fungi isolates and its discolorization effects**

There were eight isolates obtained from seasoned rubberwood after seven days incubation and two isolates showed the highest percentage growth on the surface area of the wood test samples for six weeks, which were 90.6% for isolate I4 and 96.2% for isolate II4. Based on the percentage of growth, both isolates were given scores 5 (Figure 1).

The stain fungi that have been obtained probably belong to the blue stain or sap stain fungi group and not to the mold group. The discoloration of wood caused by the melanin of fungi that persist in wood even after the fungus death. The blue stain fungi have the ability to penetrate deeper into wood compared to mold that only grows on the wood surface. It is caused by the hyphae that penetrate from the pits in wood or directly through the cells by forming boreholes (Daniel 2016). Therefore, the blue stain fungi can not be removed by brushing treatment, in contrast to mold fungi. Based on this condition, many parts of wood will be wasted to remove blue stain and caused dimensional changes on wood.

The ability of stain fungi to grow on wood was caused by the presence of simple sugars and starches as a source of nutrition. Schmidt et al. (2016) in his research informed that A. niger was able to colonize wood samples in first-week incubation depends on the condition of substrates rich in glucose and starch of 162.3 and 33.5 nmol mg-1 respectively. Another study also reported that the carbohydrate content most commonly found in rubberwood is starch (Ketkakomol et al. 2014). Stain fungi do not attack cellulose, hemicellulose, and lignin so that high weight loss does not occur in wood. Stain fungi will reduce the economic value due to decrease in the aesthetic of wood. Another research had stated that the presence of stain fungi on wood was increased the water absorption. The higher water sorption, the higher moisture content. Based on this condition, the higher moisture content will increase the possibility of the growth of decay fungi (Feng et al. 2014).

![Figure 1](image-url)
Figure 2. Characteristics of Isolate I4. A. Colony at above view, B. Colony at reverse view, C. Microscopic structure. Scale bar: A and B = 1 cm, C = 10 µm

Figure 3. Characteristics of Isolate II4. A. Colony at above view, B. Colony at reverse view, C. Microscopic structure. Scale bar: A and B = 1 cm, C = 10 µm

Figure 4. Neighbor-joining of isolates IPBCC 19.1482 (I4) and IPBCC 19.1483 (II4) with *Fusarium solani* as outgroup

**Morphology of stain fungi**

Isolate I4 has macroscopic character consisting of a colony that was black at the top and pale white at reverse color, and the colony diameter in 7.35 cm after seven days incubation in MEA at room temperature (28±2°C). In microscopic structures, isolate I4 showed the hyaline of vegetative hyphae and conidiophore, uniseriate (absence of metulae), septate hyphae, vesicle size 28.82 (28.15-29.20) µm with globose shape, conidia size 2.85 (2.09-3.09) µm with globose shape and brownish color (Figure 2).

The observation values obtained are different from Silva et al. (2011) that observed *Aspergillus foetidus*. Previous study reported that size of colony diameter and conidia were 6.2-6.6 cm and 4-5 µm respectively after
seven days incubation in MEA medium at 25°C. The differences were thought to be due to the influence of temperature, giving a physiological effect on the species of fungi. Another reference informed that there is a different growth among black Aspergillus when cultivated in different temperatures during ten days. The optimal growth was showed at temperature 33-36°C, while in lower temperature (15°C) the growth was still found with smaller colony diameters (Meijer et al. 2011).

Isolate II4 has the macroscopic characters consisting of a colony that was yellow green at the top and pale green at reverse color, and the colony diameter in 8.30 cm after seven days incubation in MEA at room temperature (28±2°C). In microscopic structures, isolate II4 showed the hyaline of vegetative hyphae and conidiophore, uniseriate (absence of metulae), septate hyphae, vesicle size in 19.96 (16.67-22.19) µm with subclavate shape, conidia size in 2.85 (2.18-3.84) µm with globose shape and yellowish-green color (Figure 3).

These observations have similarities with the results from Frisvad et al. (2019) that observed Aspergillus aflatoxiformans with macroscopic characters were white mycelium and yellow-green color of conidia. Based on microscopic analysis, A. aflatoxiformans showed yellow-green conidia, uniseriate, hyaline conidiophore, vesicle size 23-38 µm, and conidia size of 3.5-5 and 3-4.5 µm in MEA medium at 25°C during seven days incubation.

**Phylogenetic of stain fungi**

The results of molecular identification are presented on phylogenetic tree (Figure 4). Isolate I4 has 100% similarities with Aspergillus foetidos with accession number of MN542769, whereas isolate II4 is 100% closer to Aspergillus aflatoxiformans with accession number of MN542794. The isolates were then stored in IPB Culture Collection (IPBCC) with the code IPBCC 19.1482 for isolate I4 and IPBCC 19.1483 for II4.

Based on these results, the stain fungi isolates obtained from rubberwood are different results with Oldertrøen et al. (2016) who obtained Aspergillus niger, Aspergillus flavus, and Penicillium citrinum. However, these results still showed the similarities in genus level. Some studies reported that Aspergillus can also be found in spruce wood (Picea canadensis), other products that are made from wood, and decayed wood (Emoghene et al. 2014; Abdel-Azeem et al. 2019; Al-tememe et al. 2019). This suggests that Aspergillus spp. has the ability to grow on a variety of substrates.

Aspergillus foetidos is a fungus that is included in the Aspergillus section Nigri and can be found on various substrates apart from rubberwood. Another study reported that A. foetidos had been isolated and identified from raisins, beans, coffee, and chocolate (Silva et al. 2011). Based on the references, it was found that A. foetidos had been isolated and stored in one of the culture collections in Indonesia, but the isolates obtained were not from rubberwood. Hence, A. foetidos would be a new record in Indonesia regarding the distribution and description of its colony. Aspergillus aflatoxiformans that was obtained relatively new and has not been recorded at several culture collections in Indonesia. The isolate was grouped in Aspergillus section Flavi that first discovered by Frisvad et al. in its publication in 2019. Aspergillus aflatoxiformans were first identified from agricultural soil, stored rice grains, and sesame kernels from market in Nigeria, while our result this species was obtained from rubberwood.

According to this consideration, A. aflatoxiformans would be a new record for Indonesia.

**Bioactivity of chitosan on stain fungi**

The increase of chitosan concentration was showed negative response to the growth of A. foetidos. In all experiments, addition of chitosan in PDA medium showed the inhibition in each concentration after seven days incubation. This research had shown that the largest growth diameter was in the control of 9 cm, while the smallest diameter was found in chitosan 2% of 7.18 cm (Figure 5).

The highest growth inhibition of 2% chitosan concentration was 20.28%. Analysis of variance showed that addition of chitosan significantly affected the growth of A. foetidos. Based on the differences among mean, concentration 2% distinctly different among other treatments (Figure 6).

![Figure 5](Image)

**Figure 5**. Growth of Aspergillus foetidos on PDA medium with various chitosan concentration. A. Control acetic acid 2%, B. Chitosan 0.5%, C. Chitosan 1%, D. Chitosan 2%. Scale bar = 1 cm
The growth of inhibition obtained was still lower than other previous study. The lower inhibition was suspected that Aspergillus foetidus resistance to chitosan. Ing et al. (2012) informed the resistance of A. niger to chitosan caused by high chitin content in the cell wall. The content of chitin and β-1,3-glucans in cell walls influences the tolerance of fungi to chitosan. The presence of chitin plays a role in helping maintain cell walls, whereas glucans regulate the flexibility and consistency of cell walls (Aranda-Martinez et al. 2016). The inhibition growth was suspected by the interaction between chitosan and the plasma membrane which increased the oxidative stress (Fawzya et al. 2019). Another research informed that chitosan damage membranes and penetrates into fungal cells, thereby damaging intracellular composition, for instance, inhibit the absorption of nutrients, interfere with mRNA and protein synthesis (El-Guilli et al. 2015).

On the other hand, A. aflatoxiformans showed a different response to chitosan. The addition of chitosan on PDA media showed a positive response to growth of A. aflatoxiformans. The higher of chitosan concentration, the higher growth of A. aflatoxiformans. The addition of chitosan concentration 2% showed 8.28 cm of colony diameter, while control only 7.72 cm. Analysis of variance confirmed that there is no inhibition effect of chitosan to A. aflatoxiformans. Based on the differences among mean, concentration 2% distinctly different from control (Figure 7).

Another response was showed of A. aflatoxiformans when chitosan added to medium. There was a clear zone in medium around the colony. The clear zone was only showed by A. aflatoxiformans in the media treated with chitosan, while there was no clear zone in control medium (Figure 8).

The positive response on the growth of this isolate is suspected that A. aflatoxiformans is able to degrade chitosan in the medium by chitosanase enzyme. Nampally et al. (2015) reported that a similar response was showed by Aspergillus niger when cultivated in Luria-Bertani (LB) agar medium with addition chitosan 0.9%. Another research confirmed that clear zone was also reported by A. flavus in chitosanase detection agar (CDA) medium after five days incubation (Zu et al. 2012). The degradation of chitosan was suspected by the activity of chitosanase enzymes. Chitosanase of Aspergillus fumigatus hydrolyzed 100% chitosan with deacetylated degree of 70%. This enzyme showed the ability to split β-1,4 glycosidic bonds between glucosamine (GlcN-GlcN) and N-acetyl

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**Figure 6.** Percentage of growth inhibition of *Aspergillus foetidus*. The same letters indicated not significantly different of mean according to Duncan’s test ($p < 0.05$)

**Figure 7.** Diameter growth of *Aspergillus aflatoxiformans*. The same letters indicated not significantly different of mean according to Duncan’s test ($p < 0.05$)

**Figure 8.** Clear zone of *Aspergillus aflatoxiformans*. A. Control acetic acid 2%, B. Chitosan 0.5%, C. Chitosan 1%, D. Chitosan 2%. Arrow = Clear zone; Scale bar = 1 cm
glucosamine (GlcNAc-GlcN) (Hirano et al. 2012). In addition, Zhang et al. (2015) confirmed that chitosanase subclass II was produced by A. clavatus. Chitosanase was classified into three subclasses based on the specificity of cleavage, which was subclass I that split both GlcN-GlcN and GlcNAc-GlcN linkages, subclass II can split only GlcN-GlcN linkages and subclass III can cleave both GlcN-GlcN and GlcN-GlcNac linkages. According to this result, chitosan showed the low inhibition effect on the A. foetidus growth and there is no inhibition against A. aflatoxiformans. Therefore, in order to inhibit growth of A. foetidus and A. aflatoxiformans could be tested using plant or microbial extraction and other microbial secondary metabolites that have been published in several references.

In summary, A total of eight isolates of fungi were isolated from seasoned rubberwood and two of them showed the highest score of growth on wood test samples. Both isolates were identified as Aspergillus foetidus and A. aflatoxiformans and belong to the group of blue stain fungi. According to publication and distribution in culture collections, A. foetidus was not isolated and identified yet from rubberwood, therefore this isolate can also be said to be a new record in Indonesia. Based on morphological, molecular characteristics, and the search of publications and culture collections of A. aflatoxiformans, this isolate would be a new record in Indonesia, thereby this information was increasing the references about the habitat and distribution of A. aflatoxiformans species. The chitosan showed the various response to inhibit the growth of blue stain fungi. Chitosan concentration 2% was shown 20.28% growth inhibition of A. foetidus, while there was no growth inhibition showed of A. aflatoxiformans. The addition of chitosan concentration 2% in medium increased the growth of A. aflatoxiformans up to 8.28 cm, while control only 7.72 cm. Another response was shown by A. aflatoxiformans in chitosan treatment by forming a clear zone around the colony and it still unknown that the clear zone was formed by degradation of chitosan as a nutrition source or response to unfavorable environmental conditions.

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