Short Communication: Microscopic decay pattern of yellow meranti (Shorea gibbosa) wood caused by white-rot fungus Phlebia brevispora

ERWIN
Faculty of Forestry, Universitas Mulawarman, Kampus Gunung Kelua, Jl. Penajam, Samarinda 75119, East Kalimantan, Indonesia. Tel.: +62-541-735089, 749068, Fax.: +62-541-735379, email: merwin0903@gmail.com

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Abstract. Erwin, 2016. Microscopic decay pattern of yellow meranti (Shorea gibbosa) wood caused by white-rot fungus Phlebia brevispora. Biodiversitas 17: 417–421. The anatomical changes of wood decaying caused by white-rot fungus Phlebia brevispora could provide the basis for evaluating and analysis of decay on yellow meranti (Shorea gibbosa) heartwood. By using soil-block test procedure of JIS K-1571 and microscopic analysis, a progressive decay in vitro of S. gibbosa wood caused by P. brevispora was well characterized. The percentage of wood weight loss was ranged from 0.91% to 12.34% in 2-12 weeks' incubation. On the first 6 weeks of incubation of S. gibbosa infected with P. brevispora, the early stages decay, in which pit erosion and slight erosion of cell walls facilitated by hyphal spreading among cells. The intermediate decay features of numerous and conspicuous holes as well as erosion troughs in cell walls were found after 8 weeks' incubation. Furthermore, complete degradation of wood cell components, defined as the advanced stage of decay, was found in some areas of wood after 12 weeks' incubation. The pattern of wood decay was similar to those of the decayed xylem of S. gibbosa stem canker in field conditions.

Keywords: Cell degradation, microscopic, Phlebia brevispora, Shorea gibbosa, wood decay

INTRODUCTION

White rot basidiomycetes that cause the decay are especially important in wood decomposition because they are the only fungi capable of degrading all cell wall components (cellulose, lignin, hemicelluloses) of wood (Blanchette 1991; Schmidt 2006; Schwarze 2007). Micromorphological aspects of two main types of white rot, selective delignification and simultaneous rot, have been distinguished (Blanchette 1984; Otjen et al. 1987; Anagnost 1998; Schwarze 2007). In selective delignification, lignin in the secondary wall and middle lamella is almost entirely removed, whereas large quantities of cellulose in the S2 layer of the cell wall are left intact and are separated from one another. Simultaneous rot is characterized by removal of both cellulose and lignin, leaving cells either riddled with bore holes and erosion troughs, or with extensively thinned secondary walls.

A white-rot fungus has been isolated from decayed xylem of Shorea gibbosa stem canker, namely Phlebia brevispora (Erwin et al. 2010) and was suspected to cause serious wood decay on this tree species (Erwin 2012).

Shorea gibbosa is known as a member of yellow meranti group (Ogata et al. 2008) which has been long managed for timber production and used for many wood products, therefore, the microbial decay processes go along with a loss of wood quality will affect the lumber value and the wood products in use.

The present study is intended to the previous reports of Erwin (2010) and Erwin et al. (2012) with presented the anatomical features of S. gibbosa heartwood infected with P. brevispora under laboratory conditions (in vitro). Although microscopic observation techniques are not applicable in the field use, however, the decay pattern of the infected wood can clearly be characterized and very useful for providing valuable information and understanding the stages of wood degradation by the fungal attack.

The aim of this study was to (i) evaluate the ability of P. brevispora to degrade S. gibbosa heartwood, and (ii) confirm the decay pattern of the fungus in artificial laboratory conditions by microscopic observations.

MATERIALS AND METHODS

Fungal strain

The decay fungus isolated from decayed xylem of S. gibbosa stem canker, and designated as YM3, was genetically identified by their internal transcribed spacers (ITS) sequence as Phlebia brevispora (Erwin et al. 2010). The fungal strains were maintained at 4°C on PDA slants.

Decay test procedure

For this experiment, inoculation procedures followed the JIS K 1571 soil-block test procedure (JIS K 1571 2004). A medium of 250 g quartz sand and 80-85 ml of nutrient solution (4.0% glucose, 0.3% peptone, and 1.5% malt extract) used for culture media. Twelve sound wood-blocks (20 mm x 20 mm x 10 mm) in radial, tangential, and longitudinal directions, respectively, were obtained from uninfected heartwood of the stem disks of S. gibbosa. The
blocks were oven dried and weighed, then sterilized with gaseous ethylene oxide at 50 °C for 5 h. The blocks were introduced in four glass jars (each glass jar containing three blocks), and inoculated with the liquid fungal culture of the isolated fungus, then aseptically incubated at 26 ± 2 °C and 70-80% RH for each 2, 4, 6, 8, 10 and 12 weeks. The blocks of each incubation period were brushed clean to remove superfluous mycelia. Nine blocks were oven-dried at 70 °C until a constant dry weight was reached. Percent weight loss due to decay was then calculated; three blocks were reserved for microscopic observations.

Microscopic observation

The dried blocks were sectioned with razor blade then fixed in 2.5% glutaraldehyde in 0.1 M phosphate buffer (pH 7.2) at 4 °C overnight, washed four times in 0.1 M phosphate buffer at pH 7.2 for 15 minutes each, and rinsed three times in distilled water for 5 minutes each. The blocks were placed in an ethanol dehydration series of 50, 80, 95, 100% each for 20 minutes, and then three times in 100% ethanol. The dehydrated blocks were freeze-dried, and mounted on SEM stubs, then coated with gold-palladium using Jeol JFC-1200 Fine Coater. The coated samples were observed under a JEOL Scanning Microscope (JSM-5310) and the EDAX application program used to obtain SEM images of the altered properties of wood.

RESULTS AND DISCUSSION

The weight loss of S. gibbosa wood after P. brevispora decay over 2-12 weeks is shown in Table1. Based on classification of natural durability of Indonesian woods (Seng 1990), the infected wood of S. gibbosa with 12.34% weight loss, were categorized non-resistant (class IV) against P. brevispora attack. The result indicated the fungus capable of attacking the heartwood of S. gibbosa under laboratory conditions, thus, it should be taken as a consideration for wood protection, and otherwise, the fungus can produce an extensive degradation into wood under favorable temperature and humidity. Meanwhile, microscopic observations of this decay showed various stages of decay, as shown in Figures1-4.

After 2-4 weeks' incubation, the wood blocks had lost 0.91-2.24% in weight. Abundant clamped hyphae colonizing the lumina of vessels were observed in transverse, radial and tangential views (Figures 1.A-D). However, in axial parenchyma cells-rays and fibers adjacent to heavily infected vessels-hyphae were not observed. In this case, hyphae propagated mainly in vessels where they could either grow parallel to the cell axis and diagonally across the lumina, supported at their points of attachment with the cell walls, or in the central part of the lumina, where they are held in place by hyphal branches extending from the main hyphae attached to the cell walls. Hyphae passing through the perforation plates were also detected (Figure 1.E). Despite hyphae being attached deep within the vessel walls, they did not severely damage cell walls (Figure 1.F).

Table 1. Weight loss in S. gibbosa wood infected with P. brevispora for periods of 2, 4, 6, 8, 10 and 12 weeks

<table>
<thead>
<tr>
<th>Incubation period (weeks)</th>
<th>Weight loss percentage Mean ± SE</th>
</tr>
</thead>
<tbody>
<tr>
<td>2</td>
<td>0.91 ± 0.10</td>
</tr>
<tr>
<td>4</td>
<td>2.24 ± 0.60</td>
</tr>
<tr>
<td>6</td>
<td>5.02 ± 1.03</td>
</tr>
<tr>
<td>8</td>
<td>8.23 ± 1.22</td>
</tr>
<tr>
<td>10</td>
<td>11.80 ± 5.15</td>
</tr>
<tr>
<td>12</td>
<td>12.34 ± 2.76</td>
</tr>
</tbody>
</table>

After 6 weeks' incubation, wood blocks had lost 5.02% of their weight. Fungal hyphae had extended from heavily infected vessels into rays, axial parenchyma cells and fibers mainly through pits, causing slight erosion of the cell wall (Figure 2). In vessels, rounded pit erosion was seen (Figure 2.A). Hyphal penetration into rays, parenchyma cells and fibers could also be seen (Figures 2.B-D).

Initial colonization of vessels by fungal hyphae is the typical decay pattern of simultaneous rot in hardwood caused by white-rot fungi (Zabel and Morrell 1992). Such typical decay appeared in these S. gibbosa wood samples, where fungal hyphae became quickly established, first in vessels (in 2-4 weeks' incubation), then spreading from these vessels into adjacent rays, parenchyma cells and fibers until 6 weeks decay process was reached.

At the early colonization phase of decay, damage is limited, and any visible evidence is not easily observed on the lumen surfaces as termed. This is the incipient or hidden stage of decay (Zabel and Morrell 1992; Schwarze 2007). Nowadays, this decay stages can be detected within several days by FT-NIR (Fourier transform near-infrared) spectroscopy (Fackler et al. 2006, 2007a,b) and multiplex PCRs methods (Nicolotti et al 2009).

After 8 weeks' incubation, wood blocks had sustained an average weight loss of 8.23%. Rounded pit erosions of vessels were enlarged enzymatically and coalesced to form numerous and conspicuous holes (Figure 3.A). Numbers of fungal hyphae in rays and parenchyma cells increased, and hole formation and cell wall destruction became clear (Figure 3.B-D). Figure 3.E shows the lysis zones that developed around elongated holes and which were frequently observed in parenchyma cell walls. Portions of the secondary walls were removed as well as the compound middle lamella, resulting in erosion troughs within cell walls. Hyphae had also begun to colonize fibers intensively but did not damage cell walls (Figure 3F).

Early in the degradation process, depressions could be seen on the inner surfaces of the secondary walls, the S3 layer, under and in the neighborhood of the hyphae, as shown in Figures 1 and 2. In later stages of degradation (in 8 weeks' incubation), the hyphae caused wide and deep erosion troughs. In this decay stage, the lysis zones that developed around bore holes and axially elongated troughs showed clearly the effects of fungal enzymes on cell walls, which were gradually eroded. Anagnost (1998) and Schwarze (2007) expressed that numerous bore-holes appear between two neighboring cells, showed an intermediate stage of decay had occurred.
Figure 1. Decay in S. gibbosa wood blocks caused by fungus P. brevispora after 2-4 weeks' incubation. A. Hyphal colonization in the lumen of two neighboring vessels at after 2 weeks' incubation (arrows). Bar 50 μm; B. Transverse view of hyphal colonization (arrow) in vessels after 2 weeks' incubation. Bar 20 μm; C. Tangential view of hyphal colonization (arrow) in vessels after 2 weeks' incubation. Bar 40 μm; D. Radial view of hyphal colonization (arrow) in vessels after 2 weeks' incubation. Bar 40 μm; E. Fungal hyphae (arrows) passing through perforation plates of vessels after 4 weeks' incubation. Bar 10 μm; F. Hyphae attached deep within the vessel walls after 4 weeks' incubation. Bar 5 μm.

Meanwhile, the area of decay in cell walls was found at an extended distance from the hyphae, in accordance to Takano et al. (2006), suggesting that extracellular enzymes of white-rot fungus can diffuse some distance from the fungal cell wall. The lysis zones indicated predelignification before the cell walls were completely removed. The extracellular enzymes of P. brevispora as reported by Arora and Rampal (2002) and Ponting et al. (2005) were known as laccase, then, Sharma and Arora (2011) identified xylanase and carboxymethyl cellulase.
Fig. 3. Decay in *S. gibbosa* wood blocks after 8 weeks' incubation. A. Rounded pit erosion (arrow) and coalesced holes (head arrow) in vessels. *Bar* 20 µm; B. Hyphae begin to heavily colonize parenchyma cells (arrows). *Bar* 10 µm; C. Rounded pit erosion (arrow) and coalesced holes (head arrow) in parenchyma cells. *Bar* 10 µm; D. Enlarged holes in rays (arrows). *Bar* 10 µm; E. Erosion troughs (arrows) and lyses zone (head arrow) in parenchyma cells. *Bar* 10 µm; F. Hyphae begin to heavily colonize fibers. *Bar* 10 µm

Fig. 4. Decay in *S. gibbosa* wood blocks after 10-12 weeks' incubation. A. Partial thinning of fiber cell wall (arrow). *Bar* 2 µm; B. Coalesced holes appear enlarged (arrow). *Bar* 10 µm; C. Erosion channels in parenchyma cells adjacent to infected vessels (arrow). *Bar* 10 µm; D. Complete removal of wood cells (arrow). *Bar* 10 µm

could also be released. They were responsible for degradation of lignin and cellulosic materials of wood cell walls. Due to its ability to produce such extracellular enzymes, *P. brevispora* was classified as one of hydrolytic fungi (Mitui 2012).

After 10 and 12 weeks' incubation, wood block sustained an average weight loss of 11.80% and 12.34%, respectively. Partial thinning of fiber walls was frequently observed adjacent to the completely removed cells (Figure 4a). In some vessels, the rounded and coalesced holes...
appeared to be enlarged, resulting in severe cell wall damage (Figure 4b). Parenchyma cell walls adjacent to infected vessels appear partially removed, forming a channel-like appearance (Figure 4c), and in some decay areas, due to advanced delignification, parenchyma cells have been completely removed. This decay process exhibited complete degradation of the compound middle lamella and cell corners that recognized as advanced stages of decay (Schwarze 2007). Meanwhile, complete degradation of cell wall components resulted in large voids that appeared in transverse sections of the decayed areas, as shown in Figure 4d. It seems to be a general sign of this decay stage that large holes appeared in transverse section, where all cell types had already been disintegrated, for instances Populus sp decayed by Trametes trogii (Levin and Castro 1998) and decaying of Populus deltoides by Pycnoporus sanguineus (Luna et al 2004).

In conclusion, S. gibbosa wood is susceptible to colonization and decay caused by P. brevipora under favorable temperature and humidity with a progressive decay pattern that has been well characterized here. The first 6 weeks of incubation was classified as the early stages decay, in which pit erosion and slight erosion of cell walls facilitated hyphal penetration among cells. Numerous and conspicuous holes as well as erosion troughs in cell walls, which were found at the end of 8 weeks incubation, showed that an intermediate stage of decay had occurred. Furthermore, complete degradation of wood cell components, termed the advanced stage of decay, was found in some areas of wood blocks after 12 weeks incubation.

The decay pattern in vitro that presented in this study was similar to those of the decayed xylem of S. gibbosa stem canker as reported in previous work of Erwin (2012). Therefore, a further inoculation experiment is necessary to confirm the pathogenicity of P. brevipora to S. gibbosa standing trees and also to clarify whether this fungus is one of causal agents of wood decay on the trees.

REFERENCES
