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The saprotrophic wood-degrading abilities of *Rigidoporus microporus*

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Highlights

- Rigidoporus microporus isolates displayed varying saprotrophic capabilities on wood blocks of Rubber tree (*Hevea brasiliensis*).
- Percentage mass loss of (*Hevea brasiliensis*) wood blocks caused by the pathogenic *Rigi*doporus microporus was significantly higher than that observed with the endophytic isolate.
- The endophytic isolate has very poor saprotrophic ability on Hevea brasiliensis wood blocks.

Abstract

Saprotrophic wood-decaying abilities of *Rigidoporus microporus* (Polyporales, Basidiomycota) syn. *Rigidoporus lignosus* and the structural alterations induced in wood blocks of *Hevea bra-siliensis* Muell. Arg were studied. Mass loss of wood blocks was analyzed after 3 and 6 months respectively and the patterns of decay by pathogenic and endophytic isolates of this fungus were investigated using light microscopy. Effects of temperature on growth of the isolates on malt extract agar were also investigated. The *R. microporus* isolated from a non-*H. brasiliensis* host caused the highest percentage mass loss (27.2% after 6 months), followed by isolates ED310 (21.1%) and M13 (15.7%), both collected from diseased *H. brasiliensis* plantations. The isolate initially identified as an endophyte showed very low saprotrophic wood decay capability (4.3% after 6 months). The optimal temperature for growth of the isolates was 30 °C; except for the endophytic isolate which showed highest growth at 25 °C. Wood samples degraded by the *R. microporus* isolates showed simultaneous attack of wood cell walls, typical of white rot fungi. Results of the study indicate variability in the wood degrading abilities of the isolates and the potential differences in their physiology are discussed. Our findings further support the need for a taxonomical revision of the *Rigidoporus* genus.

Keywords simultaneous decay; wood degradation; white rot; delignification
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1 Introduction

Rigidoporus microporus (Polyporales, Basidiomycota) syn. *Rigidoporus lignosus* is the most economically important pathogen of tropical rubber tree (*Hevea brasiliensis* Muell. Arg) plantations (Liyanage 1997) where it causes white rot disease (WRD). White rot disease was shown as a major problem for 43% of farmers in a smallholdings survey in Malaysia (Sail and Ahmad 2009). In Nigeria, WRD is responsible for 96 % of incidences of root diseases and results in killing of up to five trees per hectare per year in plantations (Omorusi, 2012). The taxonomy of the *R. microporus* complex is problematic. The source of the type specimen of *R. microporus* (Sw.) Overeem 1788 was from West Indies, but the source of the type specimen of the synonym *R. lignosus* (Klotzsch) Imazeki is not given in the literature and probably does not exist anymore (Ryvarden, 1976). In an earlier multigene phylogenetic study, we were able to separate *R. microporus* isolates causing WRD in Africa and Asia into two different groups (Oghenekaro et al. 2014). Non-pathogenic isolates from Peru and a herbarium specimen of a sample from Cuba identified as *R. microporus* were also separated into different groups (Oghenekaro et al. 2014). White rot disease attack of rubber wood plantations is well known throughout Southeast Asia, central, east and West Africa.

White rot fungi are known as active wood degraders secreting a wide range of hydrolytic and oxidative enzymes involved in plant polymer biomineralization (Daniel 2014). Cellulose degradation is carried out by a range of cellobiohydrolases and endoglucanases (Hori et al. 2013). Lytic polysaccharide monooxygenases are also implicated in cellulose breakdown (Bey et al. 2013). White rot fungi degrade lignin using a plethora of ligninolytic peroxidases and laccases (Daniel 2014). Ligninolytic peroxidases include lignin, manganese and versatile peroxidases that may be produced in different amounts according to the white rot species involved (Floudas et al. 2012; Fernandez-Fueyo et al. 2012). Laccases may also be produced in high amounts by white rot basidiomycetes and are implicated in various biological processes in addition to lignin biodegradation in both bacteria and fungi (Furukawa et al. 2014). A typical example is *Pycnoporus cinnabarius* (Jacq.) Fr. which has been shown to secrete large amounts of laccase into culture media (Levasseur et al. 2014).

Apart from being a serious pathogen, *R. microporus* is a typical white rot basidiomycete with saprotrophic abilities degrading the major components of wood including lignin. Epidemiological studies of *H. brasiliensis* natural forest and plantations showed high fungal density in soil (Nandris et al. 1988) suggesting a capacity for high biodegradative ability of plant residues by *R. microporus*. Majority of previous studies on *R. microporus* has been directed at population biology and molecular phylogeny (Kaewchai et al. 2010; Oghenekaro et al. 2014), pathogenicity (Farid et al. 2009; Kaewchai et al. 2009; Madushani et al. 2013), host-parasite interactions (Nicole et al, 1985; Nicole et al. 1986a; 1986b), peroxidases (Geiger et al. 1989) and laccases, with the latter being isolated, purified and studied in detail (Nicole et al. 1992; Bonomo et al. 1998; 2001; Cambria et al, 2000; 2011; 2012). In vitro studies have focused on biological control of the pathogen (Kaewchai and Soytong, 2010; Ogbebor et al. 2015).

There are presently no studies on the effects of temperature on in vitro growth of isolates of the fungus causing WRD and almost nothing is known about the role of the fungus when the tree is eventually killed. The present study primarily focused on assessment of saprotrophic wood decay ability of a tropical rubber tree pathogen (*Rigidoporus microporus*) compared to isolates described as endophytes or saprotrophs on host and non-host trees. To achieve this, we conducted initial studies to assess wood decay capacity of a sub-set of representative *R. microporus* isolates including non-pathogenic South American isolate previously identified as an endophyte (Martin et al. 2015).

2 Materials and methods

2.1 Fungal isolates

The *Rigidoporus microporus* isolates used were: ED310 (Nigeria), M13 (Malaysia), MUCL45064 (Cuba) and MS564b (Peru). The isolates, ED310 and M13 were isolated from diseased *H. bra-siliensis* plantations. MUCL45064 was isolated from an un-identified angiosperm in 2003 and identified as *R. microporus* according to records from BCC/MUCL [Belgian Co-ordinated collections of Micro-organisms (http://bccm.belspo.be/about-us/bccm-mucl)]. MS564b was isolated from sapwood of *H. brasiliensis* from the wild and identified as *R. microporus* (Martin et al. 2015). Isolates ED310, M13 and MS564b are deposited in culture collections of Forest Pathology Group at Department of Forest Sciences, University of Helsinki. The isolates used in this study were derived from isolates that were identified in our earlier work using multigene phylogeny and found to belong to 4 different clades. The isolates were previously grouped together as *R. microporus*. The African and Asian isolates were more closely related and were isolated from WRD diseased *H. brasiliensis* trees (Oghenekaro et al. 2014). The fungi were routinely maintained on 2% w/v malt extract agar plates (MEA).

2.2 Wood samples and decay tests

Wood decay studies were carried out using wood blocks obtained from Rubber tree (*H. brasiliensis*). Wood blocks were generated from the tree clone NIG 801 (Umar et al. 2010) of H. brasiliensis at the Rubber Research Institute of Nigeria. Three wood blocks measuring 3 x 1 x 0.5 cm were dried at 65 °C to constant mass. Three wood blocks per treatment were then weighed and placed in 100 ml Erlenmeyer flasks containing vermiculite (fraction size -1 mm) and nutrient solution (gl⁻¹: $NH_4NO_3 - 0.6$, $K_2HPO_4 - 0.4$, $KH_2PO_4 - 0.5$, $MgSO_4.7H_2O - 0.4$ and glucose - 1.0) in the ratio 1:6 (1 g vermiculite to 6 ml nutrient solution). Flasks containing wood blocks were stoppered with cotton wool and aluminum foil and autoclaved for 20 mins. Three agar square plugs (5 mm²) from actively growing isolates of R. microporus were inoculated into each flask with wood blocks. Sterile agar square plugs of 2% MEA inoculated into separate flasks containing wood blocks served as controls. There were three replications. Inoculated flasks were placed in an improvised incubation chamber overlaid with wet paper towels to provide humidity and prevent water loss. The chamber was covered with a lid and incubated at 25 °C. Humidity in the chamber was maintained at 60-80% by wetting the paper towels periodically at intervals of 2 weeks. Two time points were chosen, with sets of blocks harvested after three months and six months. At the end of each incubation period, the wood blocks were removed from the flask and adhering mycelia scraped off using a scalpel. Wood blocks were then dried in an oven at 105 °C for 24 hrs. Percentage dry matter loss with respect to the original dry mass of the wood blocks was then calculated.

2.3 Growth rate and temperature studies

Growth rates of the isolates at different temperature (15, 20, 25, and 30 °C) were studied in vitro using MEA. Five mm² agar plug of an actively growing culture was placed on the center of 9 cm Petri dish containing MEA. Three replicates were prepared for each isolate and temperature. Measurement of the diameter of colony growth was done daily for each replicate. Mean values were calculated.

2.4 Light microscopy observations of wood decay

To observe patterns of fungal colonization and decay by the *R. microporus* isolates, sections were cut from the wood blocks using a razor blade and examined using light microscopy. Transverse (T.S), radial and tangential longitudinal (L.S) sections were stained with either 1% w/v Safranin O or 1% w/v aniline blue (to stain and visualize hyphae) respectively. Sections were mounted in fresh glycerol on glass slides and observed with a Leica DMLB light microscope with images recorded using a Leica DC 300 camera.

2.5 Statistical analysis

The Levene test was used for testing homogeneity of variances (p > 0.05) and normality of the error variances. Normality of variances and uniformity of residuals was confirmed by the skewness and kurtosis values for the data distribution. To better satisfy conditions for normality and uniformity of variance, logarithmic values of the data distribution was used for the skewness and kurtosis test. Percentage mass loss was compared separately at 3 and 6 months, using one-way analyses of variance to determine significance at 5% level ($p \le 0.05$). Interaction of temperature and isolate on hypha length on media was tested using two-way analysis of variance at 5% level ($p \le 0.05$). Differences between the means (within groups) of the percentage mass losses caused by the different fungal isolates were tested using Tukey HSD (Honestly Significant Difference) test. Statistical analysis was performed with SPSS 21.0 Statistical package (IBM Corporation, New York, U.S.A).

3 Results

3.1 Mass loss

The *R. microporus* isolates studied showed large variations in their saprotrophic wood decay ability on rubber wood blocks. Wood blocks were densely colonized with characteristic mycelial strands of *R. microporus* by three of the isolates while isolate MS564b showed poor and faint colonization. Isolate MUCL45064 showed the greatest capacity for saprotrophic attack degrading 19.3% and 27% of the wood after 3 and 6 months respectively. This was followed by ED310 (13.7% and 21.2%) and M13 (11.3% and 15.7%) isolates (Fig. 1). The isolate MS564b showed only weak attack and 4.3% mass loss after 6 months. According to Tukey's HSD test, there was no significant difference between isolates ED310 and M13 after 3 months. (p>0.05). There was no significant difference in percentage mass loss between the non-pathogenic endophytic isolate (MS564b) and the control both at 3 and 6 months (p>0.05) (Fig. 1).

3.2 Effects of temperature on growth rate

On MEA, isolates MUCL45064 and MS564b grew at all the temperatures tested while ED310 and M13 did not grow at 15 °C after 6 days (Fig. 2). Highest hyphae growth length after 6 days (78.3 mm) was observed for M13 at 30 °C (Fig. 2). A two way ANOVA at the 5% level ($p \le 0.05$) to test interaction of temperature and isolate on hyphae length on media showed that there was significant effect (F = 103.26, p = 0.001). Consequently, we did not consider the effects of temperature and isolate type individually. However, there was generally an increase in growth as temperature increased except with the endophytic isolate, MS564b. For the endophytic isolate, mean growth at 25 °C (52.1 mm) was higher than at 30 °C (41 mm) after 6 days (Fig 2).

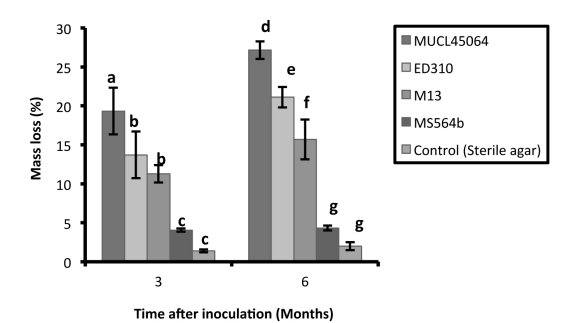


Fig. 1. Mass loss of *Hevea brasiliensis* wood blocks after 3 and 6 months caused by isolates of *Rigidoporus microporus*. Values with different letters differed significantly within groups according to Tukey HSD test (P < 0.05). Error bars indicate standard errors of the mean.

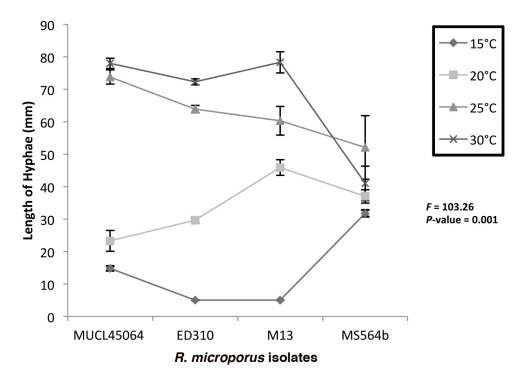


Fig. 2. Growth of *Rigidoporus microporus* isolates at different temperatures on malt extract agar after 6 days. Two way ANOVA shows interactive effect of isolates and temperature is significant (P < 0.05) and only this interaction was significant. The F and P values for the time x isolate combinations are indicated. Error bars indicate standard deviations of the mean.

3.3 Light microscopy observations of wood decay

Light microscopy was carried out to confirm the mass loss results and to visualize the attack on the wood material. All four isolates produced simultaneous attack and characteristic cell wall thinning typical of white rot fungi. The ED310, MUCL45064 and M13 isolates showed typical hyphal colonization and spreading into the rays and vessels as well as to other cellular elements. First signs of delignification occurred in the ray cells where the red staining of lignified cells changed to pinkish indicating a gradual selective lignin loss from the ray cells towards the fibers. Corresponding cell wall thinning occurred as delignification progressed (Fig. 3A). As decay continued, ultimately, the middle lamella holding the fibers together was delignified resulting in fiber separation (Fig. 3B). Examples of fungal delignification spreading from the rays to the secondary wall (S2) of fibers are shown in Figure 3A, B, C. Hyphae of most of the isolates after staining with aniline blue were observed in the vessels and ray cells (Fig. 3D, E, F). While hyphae colonization of vessels was evident, the vessel walls showed strong resistance to decay by the isolates.

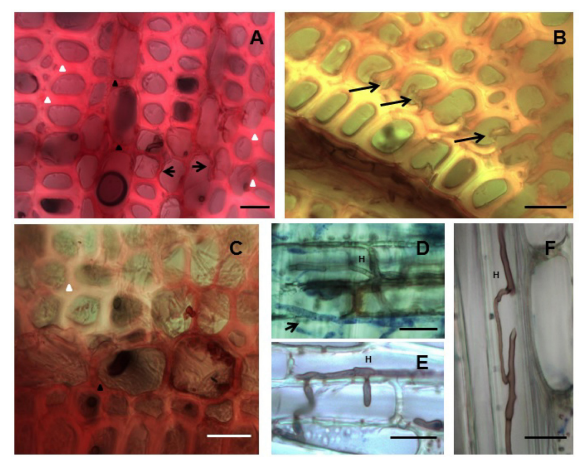


Fig. 3. Simultaneous white rot decay of *Hevea brasiliensis* wood by *Rigidoporus microporus*. (A) MUCL45064, Decay progressing from ray towards (arrows) the fibers and cell wall thinning through simultaneous attack. (B) ED310, Early stage of fibre separation due to attack of middle lamellae. (C) MS564b, Delignification spreading from across fibers as seen by the loss of staining (upper fibre regions). (D) ED310, Hypha (H) stained with aniline-blue (arrow) in the rays. (E) ED310, Hypha (H) penetrating vessel. (F) MS564b, hyphae (H) spreading through a fibre. Fig 3A-C –Transverse sections stained with safranin. Fig. 3D-F-Longitudinal sections stained with aniline blue. Bars: 20 μm.

4 Discussion

In this study, the results indicated that the *R. microporus* isolates previously isolated from trees displayed varying abilities to cause mass loss of rubber wood suggesting differences in their saprotrophic abilities. Interestingly, the values for percentage mass loss recorded were comparable to other studies using strains of the sapstain fungus Lasiodiploidia theobromae (Pat.) Griffon & Maubl. cultivated on *H. brasiliensis* and inoculated under similar conditions (Encinas and Daniel 1996; 1997). The isolate MUCL45064 collected from a non-H. brasiliensis host showed higher wood decay ability than the isolates which are proven pathogens on rubber trees. This isolate also had the highest growth on MEA at 25 °C, the same temperature used for the wood decay experiment. It is difficult to speculate on whether the higher saprotrophic ability of this isolate also suggests a potential to be pathogenic on *H. brasiliensis* plantations especially if it undergoes a host-shift. Possibility for a host-shift of R. microporus from other trees to H. brasiliensis or vice-versa has been proven through molecular studies (Oghenekaro et al. 2014). The fungus has recently been reported for the first time as pathogenic on Artocarpus nobilis Thw. (Moraceae family) in Sri Lanka (Madushani et al. 2013). An interesting feature concerned the endophytic isolate which showed very poor saprotrophic ability; perhaps this is associated with its behavior as a sapwood endophyte of H. brasiliensis in the natural forest (Martin et al. 2015).

Of the four isolates studied, the MUCL45064, ED310 and M13 isolates showed characteristic mycelial strand development in culture. *R. microporus* produces rhizomorphs in the soil and also mycelial strands under culture conditions (Richard and Button 1996). The absence of mycelial strands in the non-pathogenic isolate further points to an endophytic life-style. This isolate was previously isolated from the sapwood of healthy *H. brasilieneis* trees showing no symptoms of WRD (Martin et al. 2015). The effect of different temperature on the growth rates of the fungi also varied among the isolates. The varying saprotrophic abilities of the different isolates suggests that the isolates collectively considered as *R. microporus* (Mycobank, GenBank) and used for this study are most likely different species as suggested by Oghenekaro et al. (2014) through molecular phylogenetic studies. This supports the consensus that the *R. microporus/lignosus* species complex should be revised.

Light microscopy observations of *H. brasiliensis* degraded wood confirmed typical white rot decay. The differential pink and red Safranin staining of cell walls showed evidence of delignification as reported in previous studies (Srebotnik and Messner 1994; Schwarze and Engels 1998). The isolates colonized the wood blocks through the rays, vessels and then fibers. Direct hyphal penetration through wood cell walls was observed for all isolates except for the endophytic isolate whose hyphae only colonized and spread in the rays. There was widespread distribution of hyphae throughout vessels, fibres and tracheids in the wood blocks especially with isolate ED310 (Schwarz et al., 2004; Schwarz 2007). Simultaneous cell wall thinning and preferential degradation of cell walls was observed for the MUCL45064, ED310 and M13 isolates. Degradation of cell wall materials particularly around the middle lamella regions was also evident. The saprotrophic decay patterns observed are similar to those reported for other basidiomycete white rot fungi like *Heterobasidion annosum* (Daniel et al. 1998); *Lenzites stereoides* (Fr.) Ryvarden and *Ganoderma lucidum* (Curtis) P. Karst (Nagadesi et al. 2013); *Phanerochaete chrysosporium* Burdsall (Koyani and Rajput 2014); *Schizophyllum commune* Fries and *Flavodon flavus* (Klotzsch) Ryvarden (Padhiar and Albert 2012).

The results of the present study reveal differences in the wood decay capability of host and non-host pathogenic and endophytic isolates as well as in their physiology. The study further sheds light on the potential contribution of the isolates to carbon cycling through their saprotrophic decay activities on dead trees already killed by this fungus. This imperatively should have important implications on the management of trees in rubber plantations infected by the pathogens.

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