THE CORELATIONS BETWEEN WHITE ROT (RIGIDOPORUS LIGNOSUS L.) INCIDENCE AND SOIL CHARACTERS OF RUBBER ECOSYSTEM IN PENUMANGAN BARU, LAMPUNG

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ABSTRACT

The corelations between White Rot (Rigidoporus lignosus L.) incidence and soil characters of rubber ecosystem in Penumangan Baru, Lampung. A study was conducted to evaluate the corelation between soil characters and white rot incidence. The study was done in Penumangan Lampung from June 2007 to July 2008. Seven sites were observed on the incidence of white rot of indicator plants (cassava). Physical, chemical, and biological soil characters were correlated and regressed with white rot incidence. The results showed that there were correlation between white rot incidence and soil characters such as pH KCl, number of fungal genus (fungal diversity), Ca⁺, total cation, CEC, and saturation base. Ion Ca⁺, total cation, saturation base correlated positively with white rot incidence, on the contrary, fungal diversity, cation exchange capacity, and pH KCl correlated negatively with white rot incidence. Thus, efforts to increase fungal diversity, cation exchange capacity, and pH (in acidic soil) could be used as basic strategies to develope intergrated control of white rot disease.

Key words: Rigidoporus lignosus, rubber, soil character

INTRODUCTION

White rot disease (*Rigidoporus lignosus* L.) is very important disease in rubber plantation. The results of 1998 IRRDB disease survey that the disease is severe in Cameroon, Côte d'Ivoire, Indonesia, Malaysia, Sri Langka, and Thailand (Anonym, 2009). The disease causes economic lost not only for the lost of production but also the expensive control of the disease. The disease causes lost on production about 3-15% with the financial value about 2.1 billion rupiahs every year particularly in small holder plantation. The damage in small holder plantation occured often in the areas where stumps of rubber and primary forest trees exist, and in sandy and loose soil areas (Widjanarko, 1969; Basuki, 1981; Basuki, 1982; Situmorang & Budiman, 1990).

The control of white root disease is rarely conducted such as in Penumangan (Figure 1). This fact caused the high incidence of the disease in the area. The reasons are lack of knowledge on the disease and its control and lack of farmer's income.

Thus it is necessary to introduce control technology suitable to the social-economic condition of the farmers. In order to get the effective, cheap, and

friendly control technique, many factors involved in white root progress must be understood.

In the concept of disease triangle, the progress of white rot disease is depended upon three factors: rubber (host), *Rigidoporus lignosus* (pathogen) (Figure 2), and environment. *R. lignosus* is a polyphage pathogen, it infects many host such as rubber, cassava, coffea, cocoa, and many other trees.

There is no resistant clone of rubber available (Holliday, 1980). Where ever rubbers are grown, they are threathen by white rot disease. The component of environment to increase white rot disease is soil. Soil characters are important to disease progress.

White rot fungi attact many plant stages, but the disease, usually appears in young plantation. The symtomp will have appeared since two years old plants. And then after 5 or 6 years old the progress of the disease is slow but sure, and in the old plantation many rubber trees suffered from white rot disease (Semangun, 2000).

The plantation from forest or rubber sometimes will suffer more severe from white rot. Many infected root residues left in the soil become the food base of new infection. In central java, white rot diseases are

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Figire 1. The rubber tree were falling down attacked by R. lignosus (arrow)





Figure 2. Rhizomorphs (A), and Basidiocarps (B) of R. lignosus

found in the plantation with the history of teak forest (Soewitooetomo, 1970). The duration of the white rot fungi in the soil is depended upon number of the infected residues in the soil and many other factors involved in the root residues decomposition. According to John (1960) on the root with diameter 0.6; 2.5; and 7.5 cm, the fungi survived in 6, 20, and 40 months respectively.

Other factors in the soil role in white root progress are pH and porosity of soil. White root fungi is favoured by porous soil and neutral pH (6-7) (Semangun, 2000). The disease occured in liparite soil, in East Sumatra and Southern hill of East Java. In Southern Sumatra, the disease occured in many small holder plantation, particularly in sandy soil area (Situmorang & Budiman, 1990).

In North Sumatra, plantation with Yellow-red podzolic soil less suffer from white rot disease compared to aluvial soil. The reason is that yellow-red podzolic soil is more acid. In acid soil, the activity of *laccase*

decrease (Bonomo et al., 2001). Laccase is an enzym belongs to polyphenoloxydase involved in lignin degradation by R. lignosus (Geiger et al., 1986). Other reason is that in acid soil Trichoderma koningii, antagonist of white rot fungi, is favoured (Semangun, 2000).

Indeed, many other factors involved in the dynamic of white rot, other than mention above. Thus, it is important to investigate the relationship between other soil characters and white rot incidence.

The study is conducted to evaluate the corelation between physical, chemical, and biological soil characters and white rot incidence. It is also intended to get the best combination of soil characters to suppress white rot. So, the understanding of white rot dynamics in relation to soil characters will be unraveled, and in the long run ecofriendly control of white rot disease in small holder plantation would be established.



Figure 3. Dilution Plate method to identify and count number of soil fungi



Figure 4. Bacterial colonies are growing seperately

MATERIALS AND METHODS

The observation was conducted in small holder rubber plantation in Panumangan Baru Village, Central Tulang Bawang District, Tulang Bawang regency, Lampung Province. White rot incidences were documented across seven sites in the village and the physical, chemical, and biological characters of soil in the sites were assesed.

Within each sites, soil samples were taken from nine points of the certain area diagonally and mixed together. From the mixed soil, 2 kg of soil is placed into two plastic bags, each bag contains 1 kg of soil sample. One plastic bag was sent to soil laboratory in Bogor (Bogor Soil Research Indonesia) to assess its soil characters (texture, pH, organic matter, cation exchange capacity, etc). The other plastic bag was sent to the plant disease laboratory in the University of Lampung to assess the diversity of fungi and abudance of other soil microbes (nematode, common fungi, and bacteria).

The diversity of fungi and the abundance of the soil microbs and soil characters then be correlated with the incidence of white root rot.

Nematode Observing. Soil samples for observation of nematodes community were taken from the same point of mycorrhizal observation. From the sample points, soil sample were taken using trowel with the depth 0-20 cm as much as 0.5 kg per point and kept in polyphenyl plastic bag for the next process. The soil samples must be avoided from direct sunshine and must be kept in cool place before laboratory process.

Nematode extraction on 300 cc of soil samples were conducted using varied level of sieving tools and then centifuse in sugar dilution (500 g in 1 l solvent) according to Gafur & Swibawa (2004). Laboratory process and the enumeration of abundance of soil nematodes is the same as explain in survey of nematode in method of plant parasitic nematode management on the cofee base agricultural land.

Observing of Fungal diversity. Soil Samples for the observation of fungi other than white root rot fungi ($Rigidoporus\ lignosus$) were taken from the same point with nematode observation. From each point, 500 g soil was taken using garden trowel at 0-20 cm depth. Soil Samples were placed in plastic bags and kept in cool box before laboratory process.

The observation of fungal abundance and their diversity is conducted with *plate-counting methods* (Figure 3). Soil sample was diluted to get suspension series and the existing fungi were grown in rose-bengal chloramphenicol media on petridish.

The observation was conducted by grouping, counting, and identifying fungal colony with the same morphology. Identification of fungi were conducted at genus level based on fungal morphology (Von Arx, 1981, 1987; Rossman *et al.*, 1987; Scots *et al.*, 1994; Lane, 1997; Ellis & Ellis, 1997; Barnett & Hunter, 1998; CABI, 2001).

Observing of Bacteria. Bacteria in soil occur mostly in aggregates. To count the number of bacteria in soil, the soil must be diluted in a serial dilution and then plated. Identification of bacteria was done after a pure culture was obtained. From each point, 500 g soil was taken composited, and 100 g soil was diluted in 100 ml sterile water to make 10⁻². From the 10⁻², aseptically transfered 1 ml of dilution fluid to 9 ml blank to obtain 10⁻³. Using a fresh, sterile pipette for each succeeding step, trasfer 1 ml from the 10⁻³ dilution to another 9 ml blank to obtain 10⁻⁴ dilution. The similar procedure was repeated to obtain 10⁻⁶ dilution. From each dilution tube, aseptically placed 1 ml into each of two sterile petri plates and poured 15 ml sterile molten nutrient agar into each petri plate. Gently swirled each plate on the bench so that the inoculum thoroughly mixed with the agar. Allowed the plates to stand and solidified before inverting the plates and incubated them at room temperature for 2

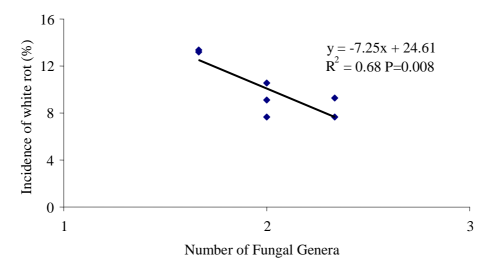


Figure 5. The relation between number of genera with white rot incidence

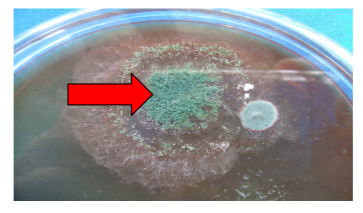


Figure 6. Colony of Trichoderma sp. (arrow) isolated from Penumangan Baru soil

days. The number of colonies were determined from countables plates (plates from dilution that yield at least 30 colonies and less than 300 colonies, and the colony separate each other (Figure 4). Grouping of the isolated soil bacteria specimens were done based on their function grouping (antagonist, pathogen, saprophyte). Antagonist group will be noticed by the presence of clear zone around the bacterial colonies.

The pathogenicity of the bacterial isolates were determined based on HR reaction on tobacco. Further identifications up to genus level were done based on biochemical test (Fahy & Persley, 1983).

Data Analysis. Analysis of correlation and regression were done using t-test at $\alpha=0.01$ or 0.05 level. The white rot incidences were correlated and regressed with each of the following variables: soil texture, soil organic matter, macro and micro-nutrients, CEC, base saturation, and pH.

RESULTS AND DISCCUSSIONS

The results of the study showed that there were correlation between white rot incidence and soil characters such as fungal diversity, Ca, total cations,

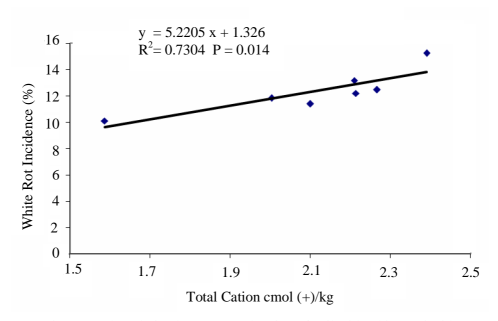


Figure 7. The relation between total cation of soil with white rot incidence

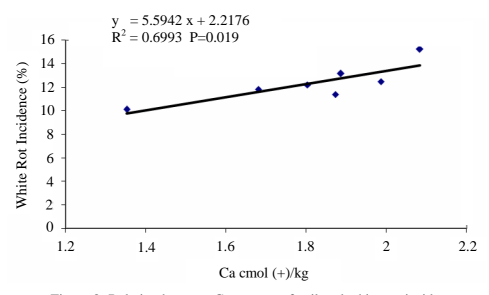


Figure 8. Relation between Ca content of soil and white rot incidence

CEC, saturation base, and pH of the soil. The correlation occured at P<0.05. There was no correlation between the diversity and abundance of nematode and bacteria investigated with white rot incidence.

There was a negative correlation between white rot incidence and fungal diversity of the soil. Soil in Penumangan belongs to acid and sandy soil. In this soil, in fact, the higher the fungal diversity of the soil the lower the white rot incidence (Figure 5). It was indicated significantly the presence of many fungal genera among them were antagonistic fungi such as *Trichoderma* sp. (Figure 6). The role of *Trichoderma* spp. in acid soil were reported by Pegg (1977). Jayasuriya & Thenakoon (2007) found *Trichoderma* isolates from rubber growing area were antagonistic to *R. lignosus*.

On the contrary, there was a positive correlation between incidence of white rot and the Ca, total cation, and saturation base of the soil in Penumangan Baru. When total cation was higher, the white rot incidence significantly to be higher (Figure 7).

The relation between total cation and white rot incidence mainly was determined by the relation between Ca and white rot incidence (Figure 8).

The increase of Ca content also increased white rot incidence. It was due to the decrease of fungal population, including antagonist of *R. lignosus* (Figure 9).

In a line with Ca and total cation, saturation base was also positively correlated with white rot incidence (Figure 10). It indicated that the higher the saturation base of the soil, the higher the incidence of white rot. It

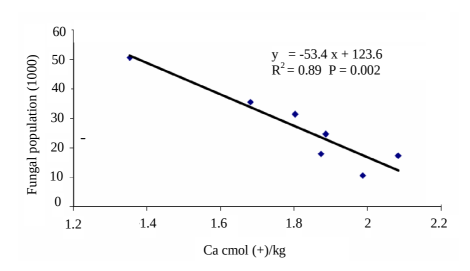


Figure 9. Relation between Ca content of soil and fungal population

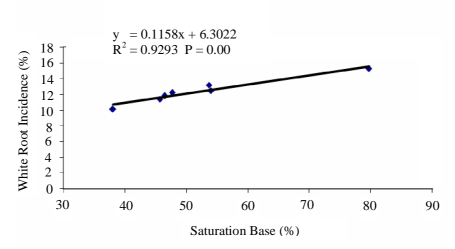


Figure 10. The relation between saturation base of soil and white rot incidence

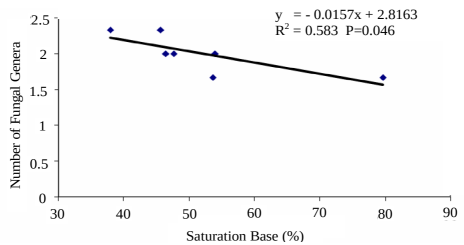


Figure 11. The relation between saturation base and fungal diversity

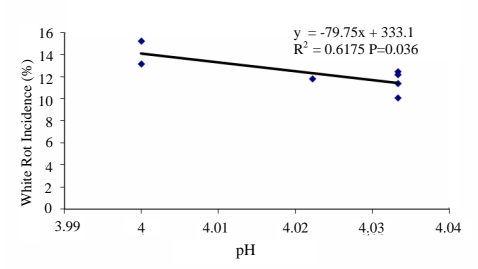


Figure 12. The relation between pH of soil and white rot incidence

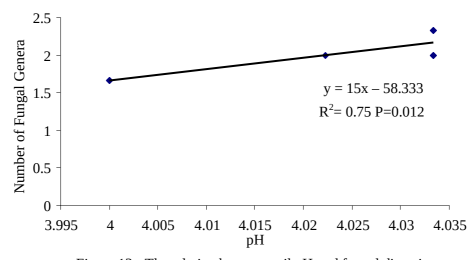


Figure 13. The relation between soil pH and fungal diversity

was due to the availability of condition nearer to neutral pH in the soil. The *R. lignosus* needed neutral condition for activity.

Other reason is that increase in base saturation decrease fungal diversity (Figure 11), including the diversity of antagonistic fungi and finally increasing white rot fungi.

It was sureprice that pH of soil correlated negatively with incidence of white rot (Figure 12). In the common sense the disease is favoured by neutral pH (Situmorang & Budiman, 1990; Semangun, 2000). It is suspected that in more acidic soil, as in Penumangan Baru (pH 4.0-4.03), the activity of *R. lignosus* is slow down for the activity *laccase* decreases for destabilization (Bonomo *et al.*, 2001). So the decrease of white rot incidence as a result of the increase of soil pH is a new phenomena. May be, It was due to the presence of *R. lignosus* antagonist.

It is alledged that the antagonist from other fungi as supported by the fact that the increase of soil pH would be followed by the increase of fungal diversity (Figure 13).

Cation exchange capacity (CEC) correlated negatively with white rot incidence. The higher CEC value the lower the white rot incidence. It was due to the availability cations for the indicator plants (cassava) and finally increase the resistance of indicator plants against *R. lignosus*.

CONCLUSION

There were correlation between white rot incidence and soil characters such as pH KCl, fungal diversity, total content of calcium, total cation, CEC, and saturation base. Calcium, total cation, saturation base correlated positively with white rot incidence, on the contrary, fungal diversity, cation exchange capacity, and pH KCl correlated negatively with white rot incidence. Any efforts to increase fungal diversity, cation exchange capacity, and pH (in acidic soil, pH 4.0-4.03) could be used as basic strategies to develope intergrated control of white rot disease.

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