EVALUATION OF Lysinibacillus SP, ISOLATED FROM Coptotermes curvignathus GUT, FOR THE DELIGNIFICATION OF OIL PALM RESIDUES

AVALIAÇÃO DE Lysinibacillus SP, ISOLADO A PARTIR DO INTESTINO DE Coptotermes curvignathus, PARA A DESLIGNIFICAÇÃO DE RESÍDUOS DE ÓLEO DE PALMA (DENDÊ)

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ABSTRACT. The application of ligninolytic bacteria and enzymes is a green pre-treatment alternative in the production of paper and biofuel from oil palm residues. In this study we investigated the ability of Lysinibacillus pakistanensis isolated from termite gut in degrading the lignin component of oil palm residues. The residues were biotreated with the bacterial strain in an aerated submerged fermentation system for 7 days at 30 °C, pH 7 and compared with untreated control. Enzyme activities were determined using specific substrates. Peak lignin peroxidase (377.6 U/L), manganese peroxidase (218.19 U/L), and laccase (405.4 U/L) activity were recorded after 4, 4, and 5 days of incubation respectively, using oil palm leaf as substrates. Lignin loss of 4.5%, 5.7% and 6.6% in oil palm leaf, oil palm trunk and empty fruit bunch respectively was achieved after treatment with the microorganism. SEM images revealed structural changes in the cell wall of the residues. Pre-treatment with this bacterial strain has promising prospects of improving the efficiency of the pulping process in an environmentally safe manner.

KEYWORDS: Biodelignification. Oil palm. Pulping. Lignin. Enzymes.

INTRODUCTION

The pulp and paper industry is one of the high demand sectors, resulting from the accelerating demands for paper-based products. Despite the increasing deployment of paperless communication with the advent of information technology, world paper consumption is continually increasing rather than decreasing as expected (SZABO et al., 2009). The removal of lignin is one of the key processes in paper production as it imparts stiffness on mechanical pulp fibres and yellow coloration on newsprints (PALIWAL et al., 2012). Moreover, the efficient hydrolysis of cellulose either for pulp or biofuel production is contingent upon degradation of lignin. Conventional methods of dissolving lignin in wood involve the use of chemicals and bleaching agents (LIEW et al., 2011; AYERONFE et al., 2018). These processes do not only require high inputs of energy but also produce highly toxic effluents which are not environmental friendly (SUMATHI; HUNG, 2006). The continuous increase in demand for paper products and growing environmental concerns necessitate the need to look towards technological improvement in the conventional pulping methods.

To this effect, research on the use of microbial enzymes as a pre-treatment method has gained interest. Pre-treatments of wood pulp with ligninolytic enzymes might provide milder and cleaner processes of delignification with no substantial loss to cellulose. This will not only save cost and energy, but will also improve the fibre bonding and the quality of paper produced (LIEW et al., 2011; FERRAZ, 2008).

The ligninolytic capabilities of fungi have been explored over the years (CRISTINA et al., 2013, OBRUCA et al., 2012; TANAKA et al. 2012). However, long weeks of incubation and ligninase enzymes expression in fungi being a secondary metabolism have limited its commercial deployment (BUGG et al., 2008). Fungi lack of stability under practical conditions also underscores the need to explore bacterial ligninolytic potential. Exploring the diversity in lignin-degrading environments; such as soils in forestry locations and also in wood digesting insects, may also reveal novel enzymatic activities for lignin degradation (TAYLOR et al., 2012).
Few reports exist on degradation of lignin by some termites species gut flora (GEIB et al. 2008; Ke et al., 2011). GEIB et al. (2008) demonstrated the depolymerisation, demethylation and ring hydroxylation of lignin by gut microflora of Zootermopsis angusticollis and Anoplophora glabripennis. Sunil et al. (2016) investigated the lignin degrading ability of Trabulsiella sp, which was reported to have degraded up to 60% of guaiacylglycerol-β-guaiacyl ether (GGE). Lignin degrading capabilities of gut flora of of Rhyynchophorus Ferrugineus have also been reported (KASSIM et al., 2016). There are also reports of aromatic degradation capabilities of bacteria isolated from termite guts. An aromatic-degrading strain of Burkholderia and Citrobacter has been isolated from the gut C. formosanus (HARAZONO et al., 2003). A strain of Rhodococcus erythropolis capable of degrading polychlorinated biphenyl was isolated from Reticulitermes speratus (CHUNG et al., 1994). These reports put forward the practicability of termite symbiotic systems as screening sources for biocatalysts of industrial interest. However, the integration of enzymes secreted by termites gut microflora in pulping process has not been explored commercially, as much effort have been focused on lignin degradation by fungi (SINGHAL et al., 2015; SINGH et al., 2013).

Oil palm is an economical perennial crop widely cultivated in West Africa and Southeast Asia. In Malaysia, Oil palm plantation increased from 2.3 million in 1994 to about 3.87 million ha in 2010 (MPOB 2011). As a consequence, the palm oil production has recorded a rocketing upswing over the years, from about 7.8 million tons in 1995 to about 15 million tonnes in 2005 and to 18.8 million tonnes in 2012 (MPOB 2012).

The explosive growth of the oil palm plantation has generated massive amounts of waste which are, at present, not efficiently utilized, thereby creating obstacles in replanting operations and colossal environmental concerns. The exploration of these oil palm residues in the production of value added products will not only help alleviate the disposal and environmental problems associated with it, but as well add value to the creation of rural agricultural-based economy. This study therefore, aimed to explore the green delignification of oil palm residues using bacteria isolated from termite’s gut.

**MATERIAL AND METHODS**

**Microorganism**

Pure cultures of bacterial isolates from Coiptotermes curvignathus gut was obtained from the Department of Crop Science, University Putra Malaysia, Sarawak. The isolate was re-streaked on LB agar plate for 24hrs and used for further analysis. Stock cultures were kept in 25% glycerol stock.

**Screening for lignin degrading potential of the isolate**

The lignin degrading capacity of the isolate was assayed for based on its ability to grow on Mineral salt media (MSM) supplemented with Kraft lignin as sole carbon source, as well as its ability to decolorise commercialized dyes. MSM-Lignin agar was prepared according to Bandounas et al. (2011). 200 mL M9 Salt solution (64 g NaHPO₄, 7H₂O, 15 g KHPO₄, 7H₂O, 2.5 g NaCl, 5.0 g NH₄Cl, 1 L ddH₂O), 100 µL 1M CaCl₂, 1mL 1M MgSO₄, 5 g of lignin powder, 15 g of agar powder, 800 ml of ddH₂O and autoclaved at 120 °C for 20 minutes. MSM-lignin agar plates were observed for colonies formation after incubation for 7 days at 37 °C. Un-inoculated MSM-Lignin plates served as controls.

The isolates which showed good growth on MSM-Lignin agar were further screened using indicator dyes (methylene blue and azure B) as described by (Bandounas et al., 2011), colonies on MSM-Lignin agar were streaked on LB agar plates supplemented with 2g/l and 2g/l methylene blue and azure B respectively. The plates were incubated at 37°C for 72hrs and observed for zone of clearance.

**Identification of lignin degrading microorganisms**

The Microbial genomic DNA of the isolate was extracted from the pure using DNeasy Qiagen Kit according to manufacturer’s instruction. The 16S rRNA gene amplification and sequencing were carried out by First Base Laboratory, Malaysia. The primers 27F (5'-AGAGTTTGTATCMTGGCTCAG-3') and 1495R (5'-GGTACCCTTGATCAGACTTT-3') were used for the amplification of the 16S rRNA gene. The PCR reactions were carried out at 95°C for 5 min, 30 cycles of 94°C for 40s, 55°C for 40s, 72°C for 80s and a final extension of 7 min at 72°C. The PCR products were analysed on 0.8% (w/v) agarose gel and sequenced. All sequences were blasted using BLAST (http://blast.ncbi.nlm.nih.gov/blast/Blast.cgi) against the NCBI 16S ribosomal RNA sequences (Bacteria only) Database, excluding uncultured Bacteria bacterium (taxid: 77133). Sequence data were aligned with software package MEGA and the
Phylogenetic tree was constructed using the neighbour-joining method.

**Biodegradation experiment**

The degradation experiment was conducted in an aerated submerged fermentation system (OBRUCA et al., 2012). To evaluate the lignin degrading capacity of the isolate, the isolate was cultured for a period of 7 days in LB medium containing 2g (oven dried weight) of the different oil palm residues (oil palm leaf (OPL), empty fruit bunch (EFB) and oil palm trunk (OPT)) as sole substrates. The residues were washed and autoclaved to get rid of contaminants prior to inoculation with the bacterial strain. Cultures were incubated at 30°C at 120 rpm for 7 days in an incubator shaker. The control and cultured samples were centrifuged at 5000 rpm for 20 min to remove biomass and suspended solids. The supernatants containing the crude enzymes extract were used to determine the activity of ligninolytic enzymes.

**Ligninolytic enzymes assay**

Lignin peroxidase, LiP activity was measured by monitoring the oxidation of veratryl alcohol to veratryl aldehyde in the presence of H₂O₂, this was determined by the increase in absorbance at 310nm. (ε310 = 9300 M⁻¹ cm⁻¹) in 100 mM citrate buffer (pH 3.0) at 30 °C. Manganese peroxidase, MnP activity was assayed spectrophotometrically with phenol red (ε610 = 22000 M⁻¹ cm⁻¹) as substrate in 50 mM sodium lactate at (pH 5.0) 30°C (OBRUCA et al., 2012). Laccase activity was measured using 2, 2-azinobis (3-ethylbenzthiazoline-6- sulphonic acid) (ABTS) in 0.1M acetate buffer (pH4.5) at 30 °C. Oxidation of ABTS was determined by the increase in A420 (ε420 = 36,000 M⁻¹ cm⁻¹) (CHEN et al., 2012). The enzyme activity was expressed in international unit s (U), defined as the amount of enzyme required to convert 1 μmole of substrate to product per minute.

**Determination of lignin loss**

The lignin content of the control and biotreated biomass was determined prior to and after degradation experiment by treatment with cold sulphuric acid according to T222om-02 test. Percentage lignin loss was calculated as follows (Tappi Test T222om-02):

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\text{% Lignin loss} = \frac{\text{Lignin content of control - Lignin content after treatment}}{\text{Lignin content of control}} \times 100
\]

**Scanning electron microscopy**

At the end of the degradation experiment, the cultured samples were centrifuged to extract the delignified residues. The residues were washed and dried. The dried residues were sputter-coated with gold-palladium alloy to increase conductivity. Morphological changes in the samples, resulting from lignin degradation, were then observed under a scanning electron microscope (Carl Zeiss EVO LS 10).

**Statistical analysis**

Data obtained were subjected to analysis of variance (ANOVA, p<0.05) using appropriate statistical software (Minitab). This test was used to determine the differences between the mean of the samples. The significance of the results was determined using p-value (p<0.05). The results indicate the mean of three replicates, whereas the error bars in graphs denote standard error.

**RESULTS AND DISCUSSION**

**Screening and identification of lignin degrading bacteria**

The bacterial strain was selected based on the ability to grow on mineral salt medium supplemented with kraft lignin as sole carbon source, as well as the ability to decolorise industrial dyes evident by the formation of zone of clearance around the plates. The selected bacterial isolate was identified based on 16S rRNA sequencing and was found to exhibit maximum homology with *Lysinibacillus pakistanensis*. The identification of *Lysinibacillus sp* from termite gut bacteria has not been reported neither has the ligninolytic potential of this genus been reported. However, *Lysinibacillus spp* were found to be present in dye contaminated soil (ANJANEYA et al., 2011; Liang et al. 2009). Saratale et al. (2003) isolated *Lysinibacillus sp*. RGS from dyes contaminated soil, capable of decolorising sulfonated dyes and textile effluents. Chaudhari et al. (2013) also reported the decolourization of reactive Orange M2R dye and chromate reduction by *Lysinibacillus sp*. KMK-A. These reports lend credence to the insinuation of a possible link between aromatic degradation and lignin metabolism, which is tenable given that much of the aromatic material found in the soil are lignin derivatives.

**Enzymes Assay**

The production of ligninolytic enzymes during degradation of oil palm residues by *Lysinibacillus sp* is being reported for the first time. The result indicates that *Lysinibacillus sp* can produce the three different ligninolytic enzymes
(Lignin peroxidase, Manganese and laccase (Figure 1).

![Graphs showing enzyme activity over time for lignin peroxidase, laccase, and manganese peroxidase.]

**Figure 1.** Lignin peroxidase (a), Laccase (b), and Manganese peroxidase (c) activity profile during the degradation of oil palm residues by *Lysinibacillus* sp. Results are mean of three independent samples with standard error.

However maximal production of lignin peroxidase varies with the substrates used. This variation could be attributed to the differences in the chemical compositions of the substrates. Ligninolytic enzymes production was higher using OPL as substrate than observed using EFB and OPT. The high lignin content of OPL (29.58%), compared to EFB (20.35%) and OPT could be a stimulant of ligninolytic enzymes production. This is consistent with a previous study on the production of ligninolytic enzymes by *Pleurotus eryngii* on agro-industrial wastes (AKPINAR; UREK 2014). It was reported that higher lignin contents in apricot wastes significantly stimulated the lignin
metabolism when compared to pomegranate wastes. Laccase and lignin peroxidase were the predominating enzymes observed during pre-treatment with the bacterial strain. Laccase activity was undetected after a day of incubation.

However, activity steadily increases in the following 3 days of incubation, reaching maximal (405.4 U/L, 375.56 U/L, 376.66 U/L) in medium supplemented with oil palm leaf, empty fruit bunch and oil palm trunk respectively, after 5 days of incubation (Figure 1). Laccase activity thereafter declined. Decline in activity could result from the build-up of lignin degradation derivatives which may exhibit inhibitory effects on enzyme production. This is consistent with the results obtained during the degradation of kraft lignin by *Pandoraee* sp. B-6 isolated from bamboo chips. It was reported that laccase activity was maintained at a low level in the initial 2 days of incubation, peak activity (1120.6 U/L) was recorded after 5 days of incubation (Shi et al., 2013). The peak laccase activity after 5 days of incubation also correlates with peak growth/beginning of cell death in the isolates (data not shown). It could be that laccase enzymes mainly functions when the exponential growth of phase is almost over. A similar conclusion was reached in a previous study (Shi et al., 2013). Maximal lignin peroxidase activity was recorded between 4-6 days of incubation. A similar finding was reported during kraft lignin metabolism by bacterial strains isolated from oil palm plantation soils (Hashimah et al., 2013). Lignin peroxidase was highest (377.6 U/L) using with oil palm leaf as substrates, as compared to 372.92 U/L and 324.16 U/L recorded using oil palm trunk and empty fruit bunch respectively. However, in contrast to decline in LiP activity observed during degradation of oil palm leaf, activity increases steadily throughout the degradation experiments using oil palm trunk and empty fruit bunch as substrates. MnP activity increases significantly in the initial 3-4 days of incubation, which was immediately followed by a marked decline. Interestingly, activity steadily increases in the following days of incubation. Decline in activity and a sudden rise in activity could be explained by the breakdown of inhibitory compounds to yield products that can be metabolised by the isolate. Maximal MnP activity observed in the initial 3 and 4 days also shows that manganese peroxidase production is crucial in lignin metabolism by *Lysinibacillus* sp.

**Determination of Lignin Loss**

Lignin losses in oil palm leaf, empty fruit bunch and oil palm trunk bio-treated with *Lysinibacillus* sp were 4.5%, 6.6% and 5.7% respectively. The loss in lignin content after degradation experiment could be attributed to the action of the ligninolytic enzymes which have also been implicated in previously reported lignin degradation experiments (Kamsani et al., 2016; Ntougias et al., 2015). There is a significant difference in the percentages of lignin loss recorded for each residue after treatment with *Lysinibacillus* sp. This could be due to the structural differences of the lignin component of each residue, as well as the degree of susceptibility of each residue to microbial degradation (Skyba et al., 2013). Colonization of wood by microorganism facilitates the softening of lignin thereby reducing the input of energy and chemicals required in subsequent pulping process. The higher the lignin loss, the more easily fibres will bleach with the utilization of few chemicals. This will ultimately improve the pulp production economics (Ferráz et al., 2008; Li ew et al., 2011).

Percentage lignin loss during lignin degradation by white rot fungi on spruce wood shavings ranged from 2.5% - 7.2% (Fackler et al., 2006). However, it took 2 weeks of incubation to achieve such extent of degradation in the fungi studied. Interestingly, only 7 days of incubation was sufficient to record comparable results in the bacterial strains involved in this study. Lignin metabolism being a primary metabolism rather than secondary as reported in fungi is propitious for potential biotechnological applications. Lignin loss of 4% was also reported during of biopulping of wood chips with *Phlebia brevispora* after 30 days incubation (Fonseca et al., 2014). Singh et al. (2010) reported a 9.35% reduction in lignin content of oil palm trunk after 4 weeks of treatment with white-rot fungus *Trametes versicolor*. 14.5% wheat straw-lignin was also degraded by *Pleurotus eryngii* (Skyba et al., 2013). Nevertheless, production parameters were optimized to enhance lignin metabolism. Higher degradation could also be achieved by the selected bacterial strains in this study with longer incubation periods if production parameters were optimized.

It is intriguing that maximum enzyme production was observed using oil palm leaf as substrates. However, higher lignin degradation was recorded in empty fruit bunch. This could result due to the low lignin content of empty fruit bunch thereby making it easily degraded than others.

The structure of lignin, which is determined by the composition of monolignols, varies between and within species (Skyba et al., 2013). The susceptibility of wood to degradation is largely
dependent on the composition, as well as distribution of lignin in the different cell layers (SKYBA et al., 2013). Moreover, the level of enzymes activity sometimes does not correlate with the degree of lignin loss, which is consistent with previous reports (FACKLER et al., 2006; KNEZEVIC et al., 2013). In some cases, the reduction in the lignin content by the fungal species correlated with the enzymes expression pattern during lignin degradation by white rot fungi on spruce wood shavings (FACKLER et al., 2006). However, it was concluded that the cumulative total peroxidase activity of the fungal cultures after 14 days with the decrease of the lignin content within the same time showed a very low correlation (FACKLER et al., 2006). Percentage lignin loss in Ceriporiopsis subvermispora correlated with increased peroxidase activity after 8 days of cultivation. However, delignification increases despite decreased peroxidase activity in the following days of cultivation. Treatment with Phlebia brevispora resulted in higher reduction in lignin content (6.2%), as compared to that with Dichomitus squalens BS1000.73 (4.4%) even though peroxidase activity in cultures of the latter was 19 fold higher than that observed in the cultures of the former. Knezevic et al. (2013) also reported that significant lignin breakdown seemed to occur in later stages of wheat straw fermentation, even though the peak of enzyme activity was reached much earlier. Apparently, the rate of lignin breakdown is not necessarily correlated with the level of enzyme activity.

Scanning electron microscopy

To gain insight into the structural changes in the cell wall of the residues, the treated and untreated samples were observed using a scanning electron microscope (Carl Zeiss EVO LS 10) (Figure 2).

The surface morphology of the untreated samples (Fig. 2) appeared more compact and unaltered as opposed the rough and fractured surfaces of the treated samples. This modification could be attributed to the loss of lignin in the treated samples. Similar observations have been reported in previous studies (SINGHAL et al., 2015; SINGH et al., 2013). Interestingly, such structural changes are crucial in pulping processes as less chemicals and energy will be required in subsequent chemical and mechanical pulping process respectively. Formation of pits on the surface of bagasse fibres was reported in the biopulping of bagasse by Cryptococcus albidus (SINGHAL et al., 2015). Alteration in the cell wall of sugarcane bagasse as a result of acidic treatments has also been reported (MORAES ROCHA et al., 2011)
Evaluation of *lysinibacillus sp.*

**Figure 2.** SEM micrographs of (a) control OPL; (b) biotreated OPL; (c) control EFB; (d) biotreated EFB; (e) control OPT; and (f) biotreated OPT
CONCLUSIONS

The application of ligninolytic bacteria and enzymes is a green pre-treatment alternative in the production of paper and biofuel from oil palm residues.

In this study we investigated the ability of *Lysinibacillus sp* isolated from termite gut in degrading the lignin component of oil palm residues. All three ligninolytic enzymes were expressed by the organism. However, the level of production of the ligninolytic enzymes, LiP, MnP and Lac varies with the different carbon sources. This could be attributed to the nature and components of the different residues. Although, highest enzyme activity was recorded using oil palm leaf as substrate, the highest lignin loss was observed in empty fruit bunch, suggesting that the level of enzyme activities does not necessarily correlate with the extent of lignin degradation. Interestingly, these observations propose the utilization of these residues not only in paper production, but also as substrates in the cost-effective production of ligninolytic enzymes for industrial applications. However, optimization of production parameters such as pH, temperature, presence of inducers, is necessary in order to enhance the degradation efficiency of the bacterial strain.

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