Lignin Degradation: A Review

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ABSTRACT

Lignin is regarded as the most plentiful aromatic polymer contains both non-phenolic and phenolic structures. It makes the integral part of secondary wall and plays a significant role in water conduction in vascular plants. Many fungi, bacteria and insects have ability to decrease this lignin by producing enzymes. Certain enzymes from specialized bacteria and fungi have been identified by researchers that can metabolize lignin and enable utilization of lignin-derived carbon sources. In this review, we attempt to provide an overview of the complexity of lignin’s polymeric structure, its distribution in forest soils, and its chemical nature. Herein, we focus on lignin biodegradation by various microorganism, fungi and bacteria present in plant biomass and soils that are capable of producing ligninolytic enzymes such as lignin peroxidase, manganese peroxidase, versatile peroxidase, and dye – decolorizing peroxidase. The relevant and recent reports have been included in this review.

Keywords: Fungi, bacteria, ligninolytic enzymes, lignin peroxidase, manganese peroxidase, versatile peroxidase, dye-decolorizing peroxidase, degradation

INTRODUCTION

Lignin is the most common aromatic organic compound found in the lignocellulose component of the plant cell wall. Its characteristic ability to absorb UV (ultraviolet) radiation makes it susceptible to degradation on being exposed to sunlight. The source of lignin in soil can be of plant origin or lignocellulosic waste from the food processing industry. It represents a significant part of plant litter input (approximately 20%) into the soil. Lignin is an amorphous three-dimensional polymer composed of phenylpropanoid subunits. It acts as a binding material and is involved in cross-linking of cellulose that provides extra strength, rigidity, and stiffness to the cell wall. Lignin protects plant cells from environmental stress conditions. The complex structure of lignin makes it recalcitrant to most degradation methods and continues to pose a critical challenge.

Lignin is the most abundant source of carbon in the soil after cellulose. Lignin degradation can thus play a major role in improving earth’s biofuel resources and also serve as an alternative to harsh technologies used in the paper and pulp industry. Degradation studies are mainly biotic, aerobic, and co-metabolic. Studies have shown that certain bacteria and fungi are able to break down various biopolymers in soil. Lignocellulosic biomass degradation has been widely studied in wood-rotting Basidiomycetes fungi due to their potential to degrade lignin.

In this review, we discuss the structural diversity of the lignin molecule, its distribution in soil, and also biodegradation of lignin by various groups of fungi and bacteria by reviewing the pertinent literature.
Lignocellulose

On the world lignocelluloses are the main part of biomass, because it is a renewable resource and the prominent structural component of plant cell wall as well. Lignocellulosic wastes are released in large amounts by many industries. Plant cell wall is generally composed of cellulose (35% to 50%), hemicellulose (20% to 35%), and lignin (15% to 20%). Cellulose is the dominant part of lignocellulose and consist of a linear chain of D-glucose linked by β (1-4)-glycosidic bonds to each other. The cellulose strains are connected to each other deliver cellulose fibril. A number of intra- and intermolecular hydrogen bonds are linked cellulose fibers together. Hemicellulose is the second plentiful constituent of lignocellulose, is comprised of diverse pentoses (arabinose, xylose) and hexoses (mannose, galactose, glucose). So, that large amount of hemicelluloses must be eliminated to improve the cellulose digestibility for the enzymatic hydrolysis.

Lignin primarily is consisted of \( p \)-coumaryl phenol (H), guaiacyl (G) and sinapyl alcohol (S). Polymerization of these constituents mainly synthesize lignin and their proportion is different between crops, woody plants and also in the primary and secondary cell wall.

Lignin Structure and Its Biosynthesis

In the plant cell, lignin is biosynthesized by the combination of three basic hydroxycinnamoyl alcohol. In the monomers plant cell, or monolignols: ligninisbiosynthesized by the combination of three basic hydroxycinnamoyl alcohol monomers or monolignols:

1. p-Coumaryl alcohol;
2. Coniferyl alcohol;
3. Sinapyl alcohol.

These monolignols are often referred to as phenylpropanoids, which differ in the substitutions at the 3-C and 5-C positions in the aromatic ring.

Lignin synthesis starts with the random self-replicating radical coupling of phenoxy radical to form an oligomeric product. After polymerization, these polymers are referred as \( p \)-hydroxyphenyl (H), guaiacyl (G), and syringyl (S) (from \( p \)-coumaryl alcohol, coniferyl alcohol, and sinapyl alcohol, respectively). Monolignols are linked either by C- C bond or C-O–C bond, and more than two third of monolignols are joined by either linkages.

Lignin in Soil

Because of the high inflow of organic aromatic matter into the soil, lignin is considered a major component of soil organic matter. The High stability and low degradability of lignin soil contribute to increasing humus formation. The copper oxidation (CuO) method is commonly used for the characterization and quantification of lignin in soil. Oxidation by CuO yields phenolic compounds such as vanillyl, syringyl, and cinnamyl type compounds. These compounds reflect the origin and extent of lignin decomposition in the soil. The sum of the above three monomeric phenolic components gives an estimate of total lignin in the soil, whereas the carboxylic acid to aldehyde ratio conveys the extent of lignin decomposition. Studies have shown that biotic, aerobic, and co-metabolic degradation are the main processes involved in lignin degradation.

Lignin Distribution in Different Soil Horizons

Distribution of lignin in different soil limits has been discussed by many groups. The lignin content in the soil decreases from the upper soil horizon to the lower soil horizon. However, in a few cases, an increase in lignin content of soil organic matter (SOM) with depth has been observed. Lignin distribution in soil could also vary with location. A relatively higher rate of lignin degradation is found in lower horizon of soil as compared to the upper horizon because the acid-to-aldehyde ratios of the vanillyl and syringyl units are greater in lower horizon. An increase in gradient is found from organic to mineral horizon. This is in accordance with the decreasing vanillyl, syringyl, and cinnamyl phenolic-lignin contents with the depth and limited supply of fresh organic materials in deep soil horizons.

The particle size of soil components also influences the lignin content and acid-to-aldehyde ratio. The lignin content of SOM decreases from the coarse to the finest particle-size fractions while the acid-to-
aldehyde ratio increases with decreasing particle size and is the highest in the clay fraction.

**Lignin Degradation in Soil**

The Lignocellulosic complex in the plant cell wall contains approximately 40 to 60% cellulose, 20 to 40% hemicellulose, and 10 to 25% lignin, which provides rigidity to the cell wall structure. Certain enzymes from specialized bacteria and fungi have been identified by researchers that can catalyze a number of oxidative and hydroxylation reactions, depolymerize the phenolic and non-phenolic lignin polymer, and also mineralize the insoluble lignin. The orientation, adsorption, and diffusion of the ligninolytic enzymes in the soil solid phase affect the lignin degradation in soil. In laboratory studies, the impact of soil particle size on soil respiration was observed by Datta et al., which can, in turn, affect lignin degradability in soil.

The biodegradation of lignocellulosic biomass has been widely studied in wood rotting Bacilomyecetes microorganisms. These bacilomyecetes are categorized as white-rot and brown-rot fungi. White-rot fungi are the most effective bio-degraders of lignocellulosic biomass (e.g., Phanerochaete chrysosporium) and can degrade lignin faster than other microorganisms. Su et al. reported that the degradation rate of lignin by Phanerochaete chrysosporium in tobacco stalk was 53.75% in 15 days.

White-rot fungi produce a number of extracellular enzymes that directly attack lignin, cellulose, and hemicellulose of the plant cell wall to decompose it. These enzymes include laccases and peroxidases, such as lignin peroxidase (LiP), manganese peroxidase (MnP), and versatile peroxidase (VP). Laccases and peroxidase enzymes can also cause lignin degradation through low molecular weight free radicals such as OH, depolymerize the phenolic and non-phenolic lignin polymer, and mineralize the insoluble lignin.

On the other hand, the brown-rot fungi are less efficient in degrading lignin compared to white-rot fungi. Lignin degradation by brown-rot fungi mainly involves non-enzymatic oxidation reactions producing hydroxyl radicals via Fenton chemistry. Brown-rot fungi partially oxidize lignin via aromatic ring demethylation. During this process, the phenolic hydroxyl content of the reaction mixture increases due to partial oxidation and partially due to the addition of new carboxyl and carbonyl groups.

Research in terms of lignin degradation by bacteria has been limited. Although fungi are reported to be more potent in lignin degradation, many soil bacteria such as Actinomycetes are also capable of mineralizing and solubilizing polymeric lignin and lignin-related compounds. Spiker showed that bacteria (e.g., *Streptomycetes viridosporus*) could oxidize phenolics but not the non-phenolic compounds. Various types of cleavages in lignin molecules, e.g., aromatic ring, demethylation, and oxidation, are catalyzed by bacterial enzymes.

Lignin-degrading fungi and bacteria can produce primarily four major extracellular heme peroxidases, including lignin peroxidase (LiP, EC 1.11.1.14), manganese-dependent peroxidase (MnP, EC 1.11.1.13), versatile peroxidase (VP, EC 1.11.1.16), and laccase (Lac, EC 1.10.3.2). These enzymes can directly attack lignin, cellulose, and hemicellulose of the plant cell wall to decompose it. Litter decomposing fungi of families such as strophariaceae, tricholomataceae, and bolbitiaceae have been found to have an evident expression of MnP. Recently, a new group of peroxidases has been identified in fungi and bacteria that is capable of degrading lignin, known as dye-decolorizing peroxidases (DyPs, EC 1.11.1.19).

**Steps in Lignin Degradation**

Lignin biodegradation involves both depolymerization and aromatic ring cleavage. Extracellular Enzyme brought about oxidation of lignin in the following steps:

1. Oxidation of β-O-4 linkages to arylglycerol compounds;
2. Aromatic rings cleavage, mostly follows the β-ketoadipate pathway;
3. Cleaved aromatic rings coupled with β-O-4 oxidation leads to the formation of cyclic carbonate structures.
The lignin–protein and lignin theories hypothesize that during the formation of humic acid from lignin, lignin first breaks down into smaller constituents, and later, these small constituents recombine to form more complex organic molecules.

### Biodegradation of lignin

#### Humification

Humification is a process of conversion of dead organic matter (leaves, twigs, etc.) into humus by the action of decomposers such as bacteria and fungi. Humification affects soil property and nature. Due to its complex nature, chemical composition, and structure, humus is still not well understood. Different theories have been proposed on humification but doubt still exists, and none of them are universally accepted. The heterogeneous nature of soil makes it difficult to generalize different experimental results. A study was done in 2009 by Alaniello et al. to investigate the different steps of humification and the changes that occur during humification. At the beginning of the experiment, a high degree of humification is seen in lignin treated soil. The overall result shows that all the natural substances tend to be almost completely mineralized or converted to substances similar to those of the treated soil, excluding lignin, which after a year incubation is still analytically recognizable in the soil.

#### Lignin-degrading fungi

Lignin degraders in nature are mainly white-rot fungi. There are several thousand species of white-rot fungi, most of them Basidiomycotina, in addition to a few Ascomycotina which can cause white-rot (Eriks-son et al., 1990). White-rot fungi belonging to the sub-division Basidiomycotina attack either hardwood or softwood, while Ascomycotina probably degrade only hardwood (Kirk and Farrell, 1987). Lignin degradation by white-rot fungi is faster than that of any other organisms and they are responsible for most of the lignin decomposition in nature. However, the growth substrate is not only lignin, but also hemicelluloses and cellulose (Buswell and Odier, 1987; Kirk and Farrell, 1987; Blanchette, 1995). The growth of fungi decreases in nitrogen- or carbon-depleted conditions and ligninolytic activity appears as a form of secondary metabolism (Brown, 1985; Kirk and Farrell, 1987).

White-rot fungi can cause selective or nonselective delignification of wood. In selective delignification, lignin is removed without any marked loss of cellulose, and in nonselective delignification all the major cell wall components are degraded (Eriks-son et al., 1990; Blanchette, 1995). Although some white-rot fungi al-ways attack wood either in a selective or in nonselective way, there are fungi which are capable of both forms of degradation, e.g. Heterobasidion annosum (Eriks-son et al., 1990; Blanchette, 1995). Among the best studied white-rot fungi are Phanerochaete chrysosporium and Phlebia radiata, which degrade lignin selectively, and Trametes versicolor which degrades lignin nonselectively (Hatakka, 1994).

White-rot fungi degrade lignin by means of oxidative enzymes (Hatakka, 1994). Because of the nature and size of the lignin molecule, the enzymes responsible for the initial attack must be extracellular and nonspecific (Kirk and Farrell, 1987; Hatakka, 1994).

The best studied extracellular enzymes of white-rot fungi are lignin per-oxidases (LiPs), manganese peroxidases (MnP) and laccase. The role of LiP and MnP in lignin degradation has been verified, while that of other enzymes is still uncertain (Hatakka, 1994). Di- erent white-rot fungi produce di erent combinations of enzymes: there are fungi producing LiP and MnP, fungi producing MnP and laccase, fungi producing LiP and laccase and fungi which produce neither LiP nor MnP, but laccase and aryl alcohol oxidase (AAO) or some other enzyme (Hatakka 1994). P. chrysosporium, P. radiata and T. versicolor belong to the LiP±MnP group. Usually the members of this group also produce laccase, but P. chrysosporium is an exception, although very recently laccase production by this fungus has been reported under specific conditions (Srinivasan et al., 1995). Several fungi belonging to the LiP±MnP and the MnP-laccase groups are eective lignin degraders, while the lignin degradation capability of fungi belonging to the LiP-laccase group is much lower (Hatakka, 1994). This may be due to the essential role of MnP in lignin de-gradation. However, lignin degradation is a complex process and the enzymes have probably synergistic effects on each other. The mineralization of $^{14}$C-lignin in optimal conditions by
the most effective white-rot fungi can reach 50% or even 70%. In an experiment with several strains of P. chrysosporium 45±50% of \( \text{b}^{14}\text{C-DHP} \) was mineralized and 69% of the \( \text{14} \text{C-lignin} \) from fir by one of the strains in 22 days (Hatakka et al., 1984).

Brown-rot fungi extensively degrade cellulose and hemicelluloses in wood, but lignin degradation is limited. Lignin is chemically modified by demethylation of its phenolic and nonphenolic units (Kirk and Farrell, 1987; Eriksson et al., 1990), and limited aromatic hydroxylation and ring cleavage of lignin also occurs (Kirk and Farrell, 1987). Brown-rot fungi are able to mineralize the methoxyl groups of lignin, but the mineralization of other parts is much lower (Buswell and Odier, 1987; Kirk and Farrell, 1987). Brown-rot fungi mainly colonize softwoods and only 6% of all wood-rotting Basidiomycotina are brownrotters (Rayner and Boddy, 1988). Brown-rotted wood is brown in colour, consisting of a high proportion of modified lignin residue and little carbohydrates, and it persists in the forest for a long time (Blanchette, 1995).

Soft-rot fungi are Ascomycotina or Deuteromycotina which degrade both hardwood and softwood, although hardwoods are degraded to a greater extent than soft-woods (Kuhad et al., 1997). All wood components are degraded, but the rate of degradation is minimal compared to that of white-rot or brown-rot fungi (Eriksson et al., 1990). Soft-rot fungi degrade wood in environments that are too severe for white- or brown-rot fungi, generally in wet environments (Blanchette, 1995). They also decompose plant litter in soils. Haider and Trojanowski (1980) and Rodriguez et al. (1996) studied the lignin mineralization capacity of soft-rot fungi. In 35 days, Chaetomium piluliferum mineralized 17% of \( \text{14} \text{C-labelled corn stalk lignin} \), but the amount mineralized by other soft-rot fungi was much less (Haider and Trojanowski, 1980). Mineralization of side chain- and methoxy-labelled lignins by all fungi was slightly higher (Haider and Trojanowski, 1980). The soil fungi Peni-cillium chrysogenum, Fusarium solani and Fusarium oxysporum mineralized 20±27% of \( \text{14} \text{C-MWL} \) from wheat straw in 28 days. P. chrysogenum can also attack kraft and organosolv lignins (Rodriguez et al., 1996). Little is known about the enzyme system of soft-rot fungi or their lignin degradation capacity as litter de-composing organisms (Haider and Trojanowski, 1980; Kirk and Farrell, 1987). Lignin-related compounds, like vanillic acid and phenols, are rapidly degraded by soft-rot fungi (Haider and Trojanowski, 1980). This could mean that soft-rot fungi are an important group of lignin degraders in mixed populations, although their lignin degradation capacity alone is limited.

**Lignin-degrading bacteria**

There are many genera of actinomycetes and eubacteria which can degrade extracted lignin and DHP (Buswell and Odier, 1987). Many bacterial strains, especially actinomycetes, can solubilize and modify the lignin structure extensively, but their ability to mineralize lignin is limited (Buswell and Odier, 1987; Ball et al., 1989; Eriksson et al., 1990; Godden et al., 1992). Actinomycetes degrade lignin as their primary metabolic activity and at high nitrogen levels compared to white-rot fungi, most of which degrade lignin via their secondary metabolism. Some species of Streptomyces and Nocardia have been shown to degrade some lignin in soil (Haider and Trojanowski, 1980). Streptomyces badius mineralizes 3.2% of \( \text{14} \text{C-MWL} \) in 30 days, and mineralization was further enhanced to 11% when cel-lulose and yeast extract were added to the medium (Eriksson et al., 1990). There is some evidence that the grass lignins are attacked by actinomycetes more efficiently than the wood lignins (Buswell and Odier, 1987).

The lignin-degrading eubacteria can be divided into erosion, cavitation and tunnelling bacteria (Eriksson et al., 1990; Blanchette, 1995). Wood is degraded by bacteria under certain extreme environmental conditions, e.g. wood saturated with water, almost anaerobic conditions or wood with a high extractive content. However, the rate of degradation is very slow (Eriksson et al., 1990; Blanchette, 1995). Erosion bacteria grow towards the middle lamella of the wood cells and cause erosion of the fibre wall, while tunnelling bacteria grow within the cell wall. The species of the tunnelling bac-teria have not been identified (Eriksson et al., 1990). Bacteria of several genera such as Pseudomonas, Al-caligenes, Arthrobacter, Nocardia and Streptomyces can degrade single-ring aromatic compounds. The role of bacteria may be significant in consuming the small molecular weight intermediate compounds produced by fungi (Vicuna, 1988; Ruttimann et al., 1991).
Although aerobic microorganisms are primarily lignin degraders in most environments, it has been shown that anaerobic rumen microorganisms are capable of degrading plant fibre cell walls (Kuhad et al., 1997). Colberg (1988) has reviewed the anaerobic microbial degradation of lignin compounds and concluded that the intermediate metabolic products called oligolignols, released during aerobic degradation, may be partially degraded to CO$_2$ and CH$_4$ by anaerobic microorganisms. Also, polymeric lignin is mineralized in anoxic sediments at a slow rate. In many studies, however, no anaerobic degradation of lignin has been observed (e.g. Hackett et al., 1977; Micales and Skog, 1997). Benner and Hodson (1985) reported that an elevated temperature of 55°C enhances the anaerobic degradation of lignin, nevertheless mineralization of DHP stays low. Chandler et al. (1980) have formulated an equation for the anaerobic degradation of different organic materials. Degradation is directly proportional to the lignin content of the material:

\[ B = 0.830 - 0.028X \]

where B is biodegradable fraction and X is the lignin content, % of the volatile solids. The data were collected from the anaerobic fermentation process using a wide range of lignocellulosic materials. Degradation is directly proportional to the lignin content of the material:

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Bacteria are typically unicellular with a size ranging from 0.5 to 3.0 μm. Because of their small size, bacteria have a very high surface/volume ratio, which allows rapid transfer of soluble substrates into the cell. As a result, bacteria are usually far more dominant than larger microorganisms such as fungi. Some bacteria, e.g. Bacillus spp., are capable of producing thick-walled endospores which are very resistant to heat, radiation and chemical disinfection (Haug, 1993).

A wide range of bacteria have been isolated from different compost environments, including species of Pseudomonas, Klebsiella and Bacillus (Nakasaki et al., 1985; Strom, 1985a,b; Falcon et al., 1987). Typical bacteria of the thermophilic phase are species of Bacillus, e.g. B. subtilis, B. licheniformis and B. circulans. Strom (1985b) reports that as much as 87% of the randomly selected colonies during the thermophilic phase of composting belong to the genus Bacillus. Many thermophilic species of Thermus have been isolated from compost at temperatures as high as 65°C and even 82°C (Bea et al., 1996).

Actinomycetes are bacteria which form multicellular filaments, thus they resemble fungi. They appear during the thermophilic phase as well as the cooling and maturation phase of composting, and can occasionally become so numerous that they are visible on the surface of the compost. Thermophilic actinomycetes have been isolated from a wide range of natural substrates, e.g. from desert sand and compost (Cross, 1968). The genera of the thermophilic actinomycetes isolated from compost include Nocardia, Streptomyces, Thermoactinomyces and Micromonospora (Waksman et al., 1939b; Strom, 1985a). Actinomycetes are able to degrade some cellulose, and solubilize lignin, and they tolerate higher temperatures and pH than fungi. Thus, actinomycetes are important agents of lignocellulose degradation during peak heating, although their ability to degrade cellulosic lignin is not as high as that of fungi (Crawford, 1983; Godden et al., 1992). Under adverse conditions actinomycetes survive as spores (Cross, 1968).

Factors affecting fungal growth

Temperature is one of the most important factors affecting fungal growth. Other important factors are sources of carbon and nitrogen, and the pH. A moderately high level of nitrogen is needed for fungal growth although some fungi, mainly wood-rotting fungi, grow at low nitrogen levels. Indeed, a low nutrient nitrogen level is often a prerequisite for lignin degradation (Eriksson et al., 1990; Dix and Webster, 1995). However, low nutrient nitrogen is a rate-limiting factor for the degradation of cellulose (Dix and Webster, 1995). Most fungi prefer an acidic environment but tolerate a wide range of pH, with the exception of the Basidiomycotina which do not grow well above pH 7.5. Coprinus species are the only Basidiomycotina which prefer an alkaline environment (Dix and Webster, 1995).

The majority of fungi are mesophiles which grow between 5°C and 37°C, with an optimum temperature of 25±30°C (Dix and Webster, 1995).
However, in the compost environment the elevated temperature means that the small group of thermophilic fungi is an important biodegradation agent. In nature, thermophilic fungi grow in garden compost heaps, bird nests, coal tips, power plant cooling pipes and e, vents, in the storage of many agricultural products (like hay, grain etc.) and in piles of wood chips and peat (Sharma, 1989; Dix and Webster, 1995). In mushroom compost, thermophilic fungi are responsible for the degradation of lignocellulose, which is a prerequisite for the growth of the edible fungus (Sharma, 1989).

Cooney and Emerson (1964) define thermophilic fungi as fungi with a maximum growth temperature of 50°C or higher and a minimum growth temperature of 20°C or higher. Thermotolerant species have a maximum growth temperature of about 50°C and a minimum well below 20°C (Cooney and Emerson, 1964). Crisan (1973), however, defines thermophilic fungi as fungi with a temperature optimum of 40°C or higher. Thermophilic and ther motolerant fungi which are known to have cellulolytic or ligninolytic activity, or which have been found growing in lignocellulose substrate or compost.

Cooney and Emerson (1964), Crisan (1973), Rosenberg (1975), Brock (1978) and Mouchacca (1997) have enumerated several species and varieties of thermophilic fungi. The growth temperature optimum for most thermophilic fungi is between 40°C and 50°C and the maximum is 55°C (See Table 2). According to Cooney and Emerson (1964), 60°C is the temperature limit of fungi and, according to Brock (1978), 60±61.5°C is the temperature limit for all eukaryotes. Temperature tolerance varies within the genera and even between the isolates of fungal species. The substrate may also have some influence on the temperature tolerance (Ofosu-Asiedu and Smith, 1973). There are indeed some reports of fungal growth above 61.5°C. El-Naghy et al. (1991) reported saccharification activity of *Sporotrichum thermophile* (syn. *Myceliophthora thermophile*) at 65°C, and Ofosu-Asiedu and Smith (1973) found *Talaromyces emersonii* still to be very active after four weeks at 60°C indicating that the fungus might be active at even higher temperatures. However, the growth of the fungus near the temperature maximum is very slow compared to that at the optimum temperature (Rosenberg, 1975).

The ligninolytic capacity of all thermophilic fungi is not known. However, most of them are known to be able to degrade wood or other lignocellulose, cellulose or hemicelluloses (e.g. Fergus, 1969; Ofosu-Asiedu and Smith, 1973; Sharma, 1989; Kuhad et al., 1997). The ability of fungi to hydrolyse hemicelluloses is probably more common than cellulose hydrolyzation (Dix and Webster, 1995).

The most effective lignin degraders are Basidiomycotina, but according to Cooney and Emerson (1964) and Mouchacca (1997) all Basidiomycotina are mesophilic. However, a few Basidiomycotina grow well at elevated temperatures. *Phanerochaete chrysosporium* (anamorph *Sporotrichum pulverulentum*) is a white-rot fungus with an optimum temperature of 36±40°C and maximum temperature 46±49°C (Mouchacca, 1997). *Ganoderma colossum* is another white-rot fungus which is still capable of growing at 45°C and has an optimum temperature of 40°C (Adaskaveg et al., 1990; 1995). In the genus *Coprinus* there are some species that have an optimum temperature of above 40°C (Crisan, 1973). Some of the wood-rotting *Coprinus* species are brown-rot fungi which modify rather than degrade lignin (Rayner and Boddy, 1988).

A thermophilic Ascomycotina, *Thermoascus aurantiacus*, has a high ligninolytic capacity (Machuca et al., 1995), and it has been isolated from compost (von Klopotek, 1962). Heat-tolerant soft-rot fungi, such as *Thielavia terrestris*, *Paecilomyces sp.* and *Talaromyces thermophilus*, are weakly ligninolytic (Dix and Webster, 1995).

Nusbaumer et al. (1996), Thambirajah and Kuthubutheen (1989), Thambirajah et al. (1995), von Klopotek (1962) and Waksman et al. (1939a,b) have studied the occurrence of fungi during composting. Waksman et al. (1939a,b) studied composting on a laboratory scale at temperatures of 28°C, 50°C, 65°C and 75°C. At 28°C the population was heterogeneous with bacteria being dominant throughout the whole period, and fungi appearing later. Fungi, together with bacteria and actino-mycetes, formed the microbial population in the compost at 50°C. At the beginning there were active thermophilic fungi,
which were followed by bacteria and actinomycetes, some of them growing on the fungal mycelium. The composts used in the other studies mentioned above were full scale.

The raw material of compost contains about $10^6$ microbial counts of mesophilic fungi/g of raw material and thermophilic fungi $10^5 \pm 10^6$/g (von Klopotek, 1962; Thambirajah and Kuthubutheen, 1989; Thambirajah et al., 1995). The predominant mesophilic fungus in the raw material has been *Geotrichum sp.* (von Klopotek, 1962; Nusbaumer et al., 1996) and the thermotolerant *fungus Aspergillus fumigatus* (von Klopotek, 1962). Counts of fungi decrease as the temperature rises, and at 64°C all the thermophilic fungi disappear. However, a mesophilic fungus, *Cladosporium cladosporioides*, grew well at 64± 65°C, but no fungi were detected at 67°C (von Klopotek, 1962). In the studies of Thambirajah et al. (1995) and Waksman et al. (1939a,b), no fungi were detected when the temperature was over 60°C. In the study of Thambirajah and Kuthubutheen (1989) fungi survived at high temperature probably due to the short duration of peak heating. When the temperature decreases below 60°C mesophilic and thermophilic fungi reappear in compost (von Klopotek, 1962; Waksman et al. 1939a,b). The dominating fungus after peak heating is *Aspergillus sp.* (Nusbaumer et al., 1996) or *Thermomyces lanuginosus* (von Klopotek, 1962), which was also found to dominate at 50°C. *T. lanuginosus* can decompose cellulose, hemi-celluloses and even lignin, although to a much smaller extent than the other components (Waksman et al., 1939a,b). In the studies of Thambirajah and Kuthubutheen (1989) and Thambirajah et al. (1995) the number of mesophilic and thermophilic fungi $(10^4 \pm 10^6)$g in mature compost were the same, but in the study of von Klopotek (1962) thermophilic fungi dominated especially in the drier parts of mature compost.

Coprinus sp. (von Klopotek, 1962; Nussbaumer et al., 1996), Panaeolus sp., Corticium coronilla and possibly Mycena sp. (von Klopotek, 1962) are Basidiomycotina occurring in compost. They were all isolated from compost during the cooling and maturation phase or from mature compost (von Klopotek, 1962; Nussbaumer et al., 1996).

### Microorganisms during composting

#### Compost environment

During composting microorganisms transform organic matter into CO$_2$, biomass, thermoenergy (heat) and humus-like end-product, (Figs. 1 and 2). The organic substrates, bulking agents and amendments used in composting are mostly derived from plant material. The main components of the organic matter are carbo-hydrates (e.g. cellulose), proteins, lipids and lignin. The capacity of microorganisms to assimilate organic matter depends on their ability to produce the enzymes needed for degradation of the substrate. The more complex the substrate, the more extensive and comprehensive is the enzyme system required. Through the synergistic action of microorganisms complex organic compounds are degraded to smaller molecules which can then be utilized by the microbial cells (Golueke, 1991, 1992).

Microorganisms require a carbon source, macronutrients such as nitrogen, phosphorous and potassium, and certain trace elements for their growth. Carbon serves primarily as an energy source for the microorganisms, while a small fraction of the carbon is incorporated into their cells. Some of the energy formed is used for microbial metabolism, and the rest is released as heat. Nitrogen is a critical element for microorganisms because it is a component of the proteins, nucleic acids, amino acids, enzymes and coenzymes necessary for cell growth and functioning. If nitrogen is a limiting factor during composting the degradation process will be slow. In contrast, if there is excess nitrogen, it is often lost from the system as ammonia gas or other nitrogen compounds. The optimum C/N ratio has been reported to be 25±40, but the value varies depending on the substrate (Golueke, 1991).

Microorganisms are able to use organic molecules which dissolve in water. If the moisture content falls below a critical level, microbial activity will decrease and the microbes become dormant. On the other hand, too high a moisture content can cause a lack of aeration and the leaching of nutrients. In the subsequent anaerobic conditions the decomposition rate decreases and odour problems arise (Golueke, 1991).
Under optimal conditions, composting proceeds through three phases: (1) the mesophilic phase, (2) the thermophilic phase, which can last from a few days to several months, and (3) the cooling and maturation phase which lasts for several months. The length of the composting phases depends on the nature of the organic matter being composted and the efficiency of the process, which is determined by the degree of aeration and agitation. At the start of composting the mass is at ambient temperature and usually slightly acidic. Soluble and easily degradable carbon sources, such as mono-saccharides, starch and lipids, are utilized by microorganisms in the early stage of composting. The pH decreases because organic acids are formed from these compounds during degradation. In the next stage microorganisms start to degrade proteins, resulting in the liberation of ammonium and an increase in the pH. After the easily degradable carbon sources have been consumed, more resistant compounds such as cellulose, hemicellulose and lignin are degraded and partly trans-formed into humus (Crawford, 1983; Paatero et al., 1984).

Humus is the end product of the humification process, in which compounds of natural origin are partially transformed into relatively inert humic substances. Humic substances can be considered as a major reser-voir of organic carbon in soils and aquatic environments (Aiken et al., 1985). The definitive structure of humic substances is not known, but they are usually divided into groups on the basis of chemical fractionation.

Organic matter present in compost is usually chemically complex and difficult to fractionate. Extraction procedures may remove only a fraction of the organic matter, and, consequently, nondestructive methods such as $^{13}$C-NMR (Almendros et al., 1992) and FTIR (Fourier transform IR) (Tseng et al., 1996) spectroscopy have been used for the analysis of humus. Aiken et al. (1985) divided humic substances into the following groups: humin (not soluble in water at any pH), humic acids (not soluble in water under acidic conditions) and fulvic acids (soluble in water under all pH conditions).

In general, immature compost contains high levels of fulvic acids and low levels of humic acids. As the decomposition proceeds, the fulvic acid fraction either decreases or remains unchanged while humic acids are produced. The degree of humification and compost maturity can be evaluated by means of the humification index. The humification index is the ratio between humic acids and fulvic acids ($C_{HA}/C_{FA}$) expressed as a percentage of the total organic carbon (Rialdi et al., 1986; Inbar et al., 1990; Chen et al., 1996). Chen et al. (1996) studied the formation and properties of humic substances during composting. They found that the humification index increases during the process. The $^{13}$C-NMR spectra of humic acids have been used to study the presence of aliphatic and aromatic structures. Both $^{13}$C-NMR analysis and FTIR spectra indicate that the humic acid fraction extracted from a mature com-post contains more aromatic structures and carboxyl groups and less carbohydrate components than that from an immature compost (Chen et al., 1996).

It has been claimed that humus is mainly formed from lignin, polysaccharides and nitrogenous compounds (Varadachari and Ghosh, 1984; Fustec et al., 1989; Inbar et al., 1989). The chemical pathway from organic matter to humus is very complex and involves a number of degradative and condensation reactions. Several schemes for the formation of humus have been proposed (e.g. Varadachari and Ghosh, 1984; Brown, 1985; Colberg, 1988). According to Varadachari and Ghosh (1984), lignin is first degraded by extracellular enzymes to smaller units, which are then absorbed into microbial cells where they are partially converted to phenols and quinones. The substances are discharged together with oxidizing enzymes into the environment, where they polymerized by a free-radical mechanism.

Composting is a dynamic process carried out by a rapid succession of mixed microbial populations. The main groups of microorganism involved are bacteria, including actinomycetes, and fungi (Golueke, 1991). Although the total number of microorganisms does not significantly change during composting, the microbial diversity can vary during the different phases of composting (Atkinson et al., 1996a). The precise nature of succession and the number of microorganisms at each composting phase is dependent on the substrate and on the preceding microorganisms in the succession (Crawford, 1983).
At the beginning of composting mesophilic bacteria predominate, but after the temperature increases to over 40°C, thermophilic bacteria take over and thermophilic fungi also appear in the compost. When the temperature exceeds 60°C, microbial activity decreases dramatically, but after the compost has cooled mesophilic bacteria and actinomycetes again dominate (McKinley and Vestal, 1985; Strom, 1985a).

Composting is an aerobic process in general, but anaerobic microenvironments may develop. Atkinson et al. (1996b) estimated that almost 1% of all the bacteria found in municipal solid waste compost were anaerobic. All the anaerobic bacteria found were highly cellulolytic and thus may play a significant role in the degradation of macromolecules. The majority of the mesophilic anaerobic bacteria were facultative, while under thermophilic conditions more obligate anaerobic bacteria were found (Atkinson et al., 1996b).

Fungal Incubation

Sacrau poplar was harvested from the experimental farm at Beijing Forestry University, Beijing, China. Before treatment, the wood sample was ground to pass through a 0.9-mm-sized screen. Two fungal strains, *T. pubescens* C7571 and *T. versicolor* C6915, were collected from Guangdong and Hebei provinces in China, respectively. Biological treatment was carried out in a 250-mL Erlenmeyer flask with 5 g of air-dried poplar wood and 12.5 mL of distilled water. The samples were sterilized in the autoclave for 20 min at 121 °C and inoculated with 5 mL of inoculum. The cultures were incubated statically at 28 °C for 4, 8, 12, and 16 weeks. The non-inoculated sample served as the control. All experiments were performed in triplicate.

Lignin Reduction by Fungi

Aerobic and anaerobic organisms produce carbon dioxide and glucose release, relatively. The most important group of microorganisms in cellulose reduction are *Basidiomycetes* (white- and brown-rot fungi), *Ascomycetes, Deuteromycetes* (soft-rot fungi), and anaerobic (rumen) fungi. Lignin biodegradation because of its complex structure and macromolecular features is complicated. A few microorganisms are identified to reduce lignin partly, which decrease just the polysaccharide component. Nevertheless, lignin virtually includes sugars, it is probably that these procedures rely on energy obtained from the sugars. It takes maximum time to achieve roughly 10% lignin degradation under 1,000 Daltons in molecular weight. For the hydrolysis of lignin oxygen is an essential principle.

Many different environmental factors influence lignin degradation. The environmental parameters motivate lignin degradation by increasing growth and metabolism of the fungi. Temperature, acidity, carbon and nitrogen sources are the major prominent parameters that affect fungal growth. In the fungal growth, combined nitrogen at the low and high contents perform an efficient function. Lignin degradation is optimum at low nitrogen level. The hydrolysis of cellulose by fungi is more harder than hemicelluloses hydrolyzation. White-rot fungi are one of the extremely skilled fungi in lignin reduction in nature.

White-rot fungi

White-rot fungi is the only organism that can decrease lignin faster than other organisms. Moreover, in nature the responsibility of white-rot fungi in lignin saccharification is high. *Basidiomycetes* are the important identified white-rot fungi. Under aerobic environment *P. chrysosporium* can reduce one gram of different separated lignins in two days. It influence in creation of about 70% CO₂ and 30% low-molecular-weight water-soluble compounds. The fungus uses lignin, hemicelluloses, and cellulose as substrate. The lignin reduction happens at the end of primary growth by cooperation of other metabolism like nitrogen. The oxidation reactions are involved in the fungal attack which decreases methoxy, phenolic, and aliphatic content of lignin. These reactions also cleave aromatic rings, and forms new carbonyl groups. Selective type of degradations involves degradation of lignin and hemicellulose compared to cellulose. For example, *C. subvermispora, Dichomitus squalens, P. chrysosporium* follow selective decay mechanism. Similarly, simultaneous, or non-selective type of degradation is the type of digestion in which all the components of lignocellulose are decayed irrespectively (e.g. *Trametes versicolor* and *Fomes*...
fomentarius). White-rot fungi are identified to grow on woody and herbaceous plants. The most examined white-rot fungi for lignin reduction in a selective way are *P. chrysosporium* and *Phlebia* while in a non-selective way *Trametes versicolor* decrease lignin.

At the high concentration, white-rot fungi can decrease pollution. In addition, since the lignin reduction system is non-particular and free white-rot fungi have an ability to decrease various contaminations. For the cultivation of fungi cost-effective substrates and various liquid medias are being used. Fungi have special oxygenic radicle which has ability to oxidize to biomolecules of other organisms that cause the death of the particular microorganism. By altering medium pH other microbe cannot easily growth into the medium, because of preventing of fungus. Moreover, many different genes create by fungi which can convert lignin into water-soluble compounds.

**Soft-rot fungi**

Most of the soft-rot fungi have identified from *Deuteromycotina* or *Ascomycotina*. These fungi are very skilled to decrease lignin in woody plants more than herbaceous crops. Soft rot fungi are degrading wood components very slowly as compared to white-rot and brown-rot fungi. The best place for the growth of soft-rot fungi are compost, soil, piles of woodchips, straw. Soft-rot fungi can reduce cellulose and hemicellulose of woody plants in a slowly way, whereas lignin reduction is somewhat slight. The adaptation of soft-rot fungi in various temperature, different pH and limited oxygen is higher than other fungi.

Soft rot fungi are no doubt the most efficient fungi to degrade lignin in mixed microbial populations.

**Brown-rot fungi**

Brown-rot fungi degrades cellulose and hemicellulose more faster than lignin. Moreover, Compare with other fungus and bacteria the way of digestibility of plant cell wall by brown-rot fungi is entirely different, because the reduction mechanism is non-enzymatic and lacks of *exoglucanases*. Phenolic and non-phenolic de-methylation result in chemical alteration in lignin which outcome of aromatic hydroxylation and ring splitting . In the wood presence lignin de-methylation is operated by brown-rot. Brown-rot fungi more effectively grown on herbaceous crops than woody plants. Among the brown-rot fungi generally *Serpula lacrymans* and *Gloeophyllum trabeum* can destruct the structure of woody plants without difficulty. The residues of brown-rot fungi is brown in colour which composed of changed lignin and also remains in the nature lacking any more hydrolysis.

**Molds**

Most of deuteromycetes and certain ascomycetes which are actually called Microfungi or molds, i.e. are usually thought to degrade mainly carbohydrates in soil, forest litter, and compost, can also degrade lignin in these environments. These molds are able to mineralize grass lignin up to 27%. Among the molds the *Penicillium chrysogenum*, *Fusarium oxysporum*, and *Fusarium solani* have been identified for their lignnolytic activities in forests. These molds or microfungi mineralized 27.4%, 23.5%, and 22.6% of a 14 C-labeled lignin prepared from milled wheat straw. However, lignin prepared from pine was much less degraded, and mineralization rate of less than 3% was obtained. The degradation of lignin has been also being studied in the red mold of bread (*Chrysonilia sitophila*). These fungi caused 20% weight loss of pine wood in 3 months, with the losses of carbohydrate and lignin being 18% and 25%, respectively. Analysis of the decayed lignin suggested that oxidative Ca ± Cb and b-O-aryl cleavages occurred during lignin degradation. Recently another mold *Neurospora discrete* was found to degrade lignin in sugarcane bagasse and produced nearly 1.5 times the amount of lignin degradation products in submerged culture. Based on this data, *N. discrete* is recorded to have high lignin degrading capability than previously reported lignin degrading fungi.

1. **Lignin-degrading enzymes from fungi**

The structure of lignin mainly composed of phenolic and non-phenolic components. Lignin forms an integral part of secondary walls in plants and it plays an important role in enhancing the efficiency of water conduction in vascular plants. In the lignin, digestibly many various fungi and bacteria are...
proficient for genes production. These enzymes comprising of lignin peroxidases (e.g. lignin peroxidase (E.C. 1.11.1.7), and manganese peroxidase (E.C. 1.11.1.7)) and laccases as well. These are hemecontaining glycoprotein which requires hydrogen peroxide as oxidant. Lignin peroxidase degrades non-phenolic lignin units. Manganese peroxidase acts on phenolic and non-phenolic lignin units through lipid peroxidation reactions. It oxidizes Mn2+ to Mn3+ which oxidizes phenol rings to phenoxy radicals leading to decomposition of compounds.

P. chrysosporium, Ceriporiala cerata, Cyathus stercolerus, C. subvermispora, Pyenoporus cinnarbinus, and Pleurotus ostreae produce enzymes which are involve in lignin degradation.

Laccases (E.C. 1.10.3.2.) are copper containing enzymes which are involved in lignin degradation. Laccases acts along with lignin peroxidase and manganese peroxidase leading to complete degradation of lignin. It catalyzes the oxidation of phenolic units in lignin and phenolic compounds and aromatic amines to radicals. The capability of laccase in lignocelluloses degradation is improved by phenolic components such as 3-hydroxyanthranilic acid, 3-ethylthiazoline-6-sulfonate which will act as redox mediators. Without the role of redox mediators laccases have a limited effect.

Other enzymes like aryl alcohol dehydrogenase, cellobiose, aromatic acid reductase, vanillate hydroxylase, dioxygenase and catalase are regarded to contribute an important function in lignin reduction. The number of research investigations on the soft-rot fungi enzyme system and their degradation of lignin is low. Peroxidases and laccase provide by F. oxysporum, Xylaria sp., and Altenaria sp., Botrytis cinerea, Myceliophthora thermophila, Chaetomium thermophilium and Paecilomyces farinosus. Although, laccase provide by soft-rot fungi in a low proportion compared to white-rot fungi.

2. Fungal extracellular ligninases

Fungi degrade lignin by secreting enzymes collectively termed “ligninases”. Ligninases can be classified as either phenol oxidases (laccase) or heme peroxidases [lignin peroxidase (LiP), man- ganese peroxidase (MnP) and versatile peroxidase (VP)]. In general, laccases use molecular oxygen as electron acceptors while peroxidases use hydrogen peroxide as a co-substrate. White-rot fungi variously se-crete one or more of the lignin-modifying en-zymes (LMEs) in addition to other compounds necessary for effective lignin degradation. It has been shown that P. chrysosporium produces several LiP and MnP isoenzymes but no laccase. Correspondingly, the genome of P. chrysospori-um contains ten LiP and five MnP genes. In addition, H2O2-generating enzyme, glyoxal oxidase (GLOX) has been found in P. chrysosporium cultures. White-rot basidiomycetes, such as Coriolus versicolor, P. chrysospo-rium and T. versicolor, have been found to be the most efficient lignin-degrading microorganism studied. Although LiP is able to oxidize the non-phenolic part of lignin (which forms 80-90% of lignin composition), it is absent from many lignin degrading fungi. In addition, electron microscopy studies of the early stages of the fungal degradation of wood have shown that oxidative ligninolytic enzymes are too large to penetrate into the wood cell wall micropores. Thus, it has been suggested that prior to the enzymatic attack, low-molecular weight dif- fusible reactive oxidative compounds must initiate changes to the lignin structure (as dis- cussed below).

Phenol oxidases (laccases) (benzenediol:oxygen oxidoreductases)

3. Lignin Peroxidase (LiP)

Lignin peroxidase (LiP, EC 1.11.1.14) is a glycosylated enzyme containing heme protein with an iron protoporphyrin prosthetic group that requires hydrogen peroxide (H2O2) to catalyze the oxidation of non-phenolic lignin units and mineralize the recalcitrant aromatic compounds. Lignin oxidation takes place via electron transfer, non-catalytic cleavages of various bonds, and aromatic ring opening. The catalytic cycle of LiP consists of one oxidation and two reduction steps as follows:

Step 1 Two-electron oxidation of the resting (native) ferric enzyme ([LiP]-Fe(III)) by H2O2 to form the Compound I oxo-ferryl intermediate [Fe(IV)];
Step 2 Reduction of Compound I by the non-phenolic aromatic reducing substrate (A) to form Compound II by gaining one electron;

Step 3 Finally, the oxidation cycle ends when Compound II is returned to the resting ferric state with a gain of one more electron from the reducing substrate A.

LiPs have a high redox potential (1.2 V at pH 3.0) as compared with other peroxidases and can oxidize phenolic and nonphenolic structures of lignin directly without a mediator.

4. Manganese Peroxidase (MnP)
Manganese (Mn) is essential for the formation of MnP. The enzyme MnP plays an important role during the initial stages of lignin degradation. Compared to laccase, MnP causes greater degradation of phenolic lignin due to its higher redox potential with the eventual release of carbon dioxide. MnP is mainly produced by a broad species of white-rot basidiomycetes such as Phanerochaete chrysosporium.

The catalytic cycle of MnP is similar to that of LiP. Like LiPs, MnPs are also heme-containing glycoproteins which require H2O2 as an oxidant. Manganese acts as a mediator during MnP enzymatic activity. To begin with, MnP oxidizes Mn2+ to Mn3+. The enzymatically generated Mn3+ oxidant is freely diffusible and participates in the oxidation reaction as a redox couple.

In addition, organic acids such as lactate and malonate can chelate Mn3+ ion. The chelated Mn3+—organic acid complex oxidizes the phenolic compounds in lignin to phenoxy radicals. High levels of Mn can stimulate MnP enzymatic activity and enhance the degradation process of lignin in soils.

5. Versatile Peroxidase (VP)
Versatile peroxidase, as the name suggests, has catalytic properties of both LiP and MnP. VP was first purified from the genera of fungi Bjerkandera and was found to transform lignin even without an external mediator.

The VP enzyme possesses a hybrid molecular architecture with several binding sites including Mn2+ and is able to oxidize Mn2+ like MnP and LiP. However, unlike MnP, VP has the dual ability to oxidize Mn2+ in the independent oxidation of simple amines and phenolic monomers. VP can also oxidize a variety of substrates (with high and low redox potentials) including Mn2+, phenolic and non-phenolic lignin dimers, and aromatic alcohols.

6. Laccase (Lac)
Lac (EC 1.10.3.2, p-diphenol oxidase) is a copper-containing enzyme belonging to the oxidoreductase group which oxidizes a wide variety of organic and inorganic substances. Lac was extracted by Yoshida in 1883 for the first time from the Rhus vernicifera tree. In 1896, it was first showed to be present in fungi by Bertrand and Laborde. Thereafter, many Lacs have been identified from fungi. The white-rot fungus produces high levels of Lac and is very efficient at decolorizing dyes.

It was initially assumed that Lac could only oxidize phenolic compound, due to its lower redox potential (450–800 mV) as compare to Lips (>1 V). However, with the involvement of a mediator, a wide variety of substance can be oxidized using Lac.

Mediators are low molecular weight compounds that are easily oxidized by Lacs and subsequently reduced by the substrate. Due to its large size, the substance cannot reach the active site of the enzyme. A mediator, due to its small size, acts as a conveyer of an electron from the enzyme to the substrate. The mediator reaches the enzyme active site easily and gets oxidized to a more stable intermediate with a high redox potential. The oxidized mediator diffuses away from the enzyme and oxidizes more complex substrates before returning to its original state. The electrons taken by Lacs are finally transferred back to oxygen to form hydrogen peroxide.

Most of the enzymes are substrate specific, in contrast to Lac activity which oxidizes a variety of substrates like polyphenols, diphenols, benzenethiol, and aromatic amines.

7. Dye-Decolorizing Peroxidase (DyP)
The DyP enzyme is also a heme-based peroxidase that can cause lignin breakdown through a radical-mediated oxidation process. The DyPs are phylogenetically distinct from other peroxidases as they possess an α + β ferredoxin-like fold. However,
their oxidation mechanism is similar to VP and MnP. They are widely found in microorganisms and classified into four types: A, B, C, and D. Bacterial enzymes are predominantly found in type A to C, while type D is mostly clustered to fungal DyPs. All kinds of DyPs have peroxidase activities; however, they differ in substrate specificity values. In addition to lignin, DyPs can also oxidize synthetic dyes, non-phenolic methoxylated aromatics, Mn2, and high redox synthetic dyes such as anthraquinone and azo dyes.

**Enzyme Assays**

The enzymes were extracted with 50 mM sodium acetate buffer (pH 5.5) supplemented with Tween 60 (0.1 g/L). The entire contents of each bioreactor was transferred to an Erlenmeyer flask and extracted with 500 mL of extracting solution. Five successive extractions were performed at 120 rpm for 4 h at 10 ± 1 °C. After the second extraction, the wood chips were stored overnight while soaking in the extraction solution at 4 °C. The crude extracts were recovered by filtration through a fine filter paper (Souza-Cruz et al. 2004). Enzyme activities were determined at 25 °C using a Helios gamma UV-Vis spectrophotometer (Thermo Fischer Scientific, California, USA). Manganese peroxidase activity was determined according to the modified method of Heinfling et al. (1998) by the formation of Mn3+ tartrate (ε238 = 6500 M/cm) from 0.10 mM MnSO4 using 100 mM tartrate buffer (pH 5) and 0.10 mM H2O2. Lignin peroxidase activity was monitored at pH 3.0 according to Tien and Kirk (1988), and the formation of veratraldehyde was monitored at 310 nm (ε310 = 9.3 mM−1 cm−1). Laccase was measured following the oxidation of 2.0 mM 2, 2′-azino-bis (3-ethylbenzthiazoline-6-sulphonic acid, ABTS) at 420 nm (Dias et al. 2004). Hemicylcellase activity was measured according to Ghose and Bisaria (1987). The liberated reducing sugars were quantified using dinitrosalicylic acid (DNS) reagent (Miller 1959).

**Fenton Chemistry in Lignin Degradation**

Brown-rot fungi hydrolyze and partially oxidize the lignocellulose component of the plant cell wall, in contrast to white-rot fungi which produce an array of extracellular lignin-degrading enzymes. During the oxidation process, a hydroxyl ion is generated via Fenton oxidation chemistry. The Haber–Weiss reaction is a specific example of the Fenton reaction. In Haber–Weiss reaction, ferrous salt and hydrogen peroxide react to produce a highly reactive hydroxyl...
free radical capable of oxidizing a wide variety of substrates. In a nutshell, Fenton reaction is a cyclic redox reaction. In the Fenton reaction, there is neither a breaking of old bonds nor the formation of new bonds. As a result, a highly reactive hydroxyl radical is produced. A detailed mechanism of the Fenton reaction has been explained using two extracellularly produced quinones, 2,5-dimethoxy-1,4-benzoquinone (2,5-DMBQ) and 4,5-dimethoxy-1,2-benzoquinone (4,5-DMBQ), in Gloeophyllum trabeum. The two hydroquinones have the ability to reduce ferric oxalate salt to ferrous oxalate. In the second step, ferrous oxalate reacts with hydrogen peroxide to generate highly reactive hydroxyl free radicals. This reactive hydroxyl free radical can react with many organic compounds including lignin. Quinone is then converted back to hydroquinone by fungi.

**Low Molecular Weight Compounds Involved in Lignin Degradation**

Electron microscopy studies have shown that enzymes such as peroxidase and laccase, due to their larger size than that of the pores of the cell wall, cannot reach to the native lignin molecule. Therefore, mediators are important parts of the ligninolytic enzyme system. Mediators help enzymes such as MnP and LiP to reach native lignin present in the wood. Many low molecular weight compounds have been identified which could play a fundamental role in the enzymatic ligninolytic systems of white-rot fungi. Lip and MnP can oxidise various lignin compounds and synthetic lignin only in the presence of a cofactor.

Lignin degradation conditions support both depolymerization and repolymerization, so it is important to optimize several factors, such as H₂O₂ and lignin concentration, O²⁻, and the presence of a suitable mediator.

**1. Manganese**

A high amount of manganese (10–100 mg/kg of dry wood) is present in the wood. Manganese deposits were found in wood decay resulting from white-rot fungi activity. Scanning electron microscopy revealed a 100-fold increase in manganese concentration as compared to delignified wood. Mn²⁺ precipitates in the form of MnO²⁻.

During lignin degradation, Mn³⁺ assists oxidation of various phenolic compounds whereas Mn²⁺ acts as a substrate and triggers MnP production. Mn²⁺ is a scavenger of the peroxide radical and it decreases the oxygen stress of cell, resulting in decreased LiP production, whereas it increases under an oxidizing atmosphere. Furthermore, Mn²⁺ induces MnsoD which further minimizes oxidative stress as well.

**2. Veratryl Alcohol**

Veratryl alcohol (VA) is synthesized from glucose. Its production starts in parallel with LiP production during the early phase of secondary metabolism. Mester et al. showed that manganese inhibits the production of VA in fungi Bjerkandera sp. strain BOSS5 and P. chrysosporium. In addition to de novo synthesis from glucose, several alternative pathways are present for VA production in Phlebia radiata. De novo synthesis of VA is repressed when products of lignin degradation are used as precursors. A is most likely the physiological substrate of LiP. Production of Lip and VA is triggered by the presence of secondary lignin metabolites. The introduction of VA to culture medium has been found to increase LiP. VA plays a key role in LiP-catalyzed oxidation reactions and LiP-mediated electron transfer reactions.

**3. Oxalate**

White-rot fungi secrete oxalate as a major aliphatic organic acid. White-rot fungi decompose oxalate to carbon dioxide and formate as they cannot accumulate acid. This is further oxidized to a superoxidized form under aerobic conditions. This superoxide directly participates in the oxidation of lignin.

The enzymes oxaloacetate and glyoxylate oxidase are responsible for the biosynthesis of oxalate. LiP and MnP can decompose oxalate in the presence of VA or Mn²⁺.
Oxidation of oxalate is necessary; it can slow down the lignin mineralization by reducing VA$^+$ and Mn$^{2+}$.

2-Chloro-1,4-dimethoxybenzene

Chlorinated anisyl metabolites (CAMs) are physiologically involved in lignin degradation. CAMs act as a substrate for the aryl alcohol oxidase enzyme involved in extracellular peroxide production. White-rot fungi produce a wide variety of halogen metabolites. CAMs and chlorinated hydroquinone metabolites (CHMs) are the most common.

The CHM biosynthesis pathway involves the formation of 9 metabolites found in basidiomycetes. Among these, 9,2-chloro-1,4-dimethoxybenzene, 2,6-dichloro-1,4-dimethoxybenzene, tetrachloro-1,4-dimethoxybenzene, and tetrachloro-4-methoxyphenol were identified.

2-Chloro-1,4-dimethoxybenzene (2-Cl-1,4-DMB) acts as a substrate for LiP and is actively involved in lignin degradation and acts as a redox mediator. DMB free radicals are formed during LiP-catalyzed oxidation. These free radicals easily diffuse away from enzyme active site and are regarded as diffusible redox mediators.

2-Cl-1,4-DMB involvement in lignin degradation is still not clear as the formation of LiP and its biosynthesis do not coincide.

Lignin Component Changes

The decrease in lignin content differed between the two fungi-treated samples during different periods (Fig. 3). A lignin decrease of 6.6% (from 23.5% to 16.9%) and 5.1% (from 23.5% to 18.4%) was measured in the initial degradation period of *T. pubescens* C7571 and *T. versicolor* C6915, respectively. During this treatment period (week 4), LiP was the only enzyme whose activity was higher in *T. pubescens* C7571 than in *T. versicolor* C6915 (Fig. 2). This result was explained by the fact that LiP oxidizes non-phenolic lignin (Umezawa et al. 1987). Between week 4 and week 8, the decrease was approximately 2.7% for both types of white-rot fungi. Thus, there was a first phase (0 to 4 weeks) of growth, during which a rapid decrease in lignin was observed, and a second phase in which the values remained fairly stable. Additionally, during this treatment period, nearly all enzyme activity for *T. pubescens* C7571 was higher than that of *T. versicolor* C6915, except for that of hemicellulase. Hemicellulases are extremely important in the degradation of lignocellulosic biomass because of their lignin-carbohydrate complexes (LCCs) (Dinis et al. 2009). During week 16, *T. pubescens* C7571 showed higher activity for all three enzymes during the last treatment period (Fig. 2), thus yielding a low lignin content of 10.4%. A recent study also demonstrated the low enzyme producing capabilities of *T. versicolor* that laccase and MnP activity was only detectable and no LiP activity was detected (Kuhar et al. 2015). The results of enzyme activities as well as lignin component changes confirmed that the ability of *T. versicolor* to remove lignin was weaker than *T. pubescens* C7571. Chemical analyses in other studies have also indicated that this species is a non-selective type and that it removes all types of cell-wall constituents not only lignin (Bari et al. 2015). In summary, lignin degradation varied during different treatment periods; because *T. pubescens* C7571 showed a higher capacity for lignin degradation, it was considered for more specific studies.

Chemical Component Analysis and Molecular Weight of Extracted Lignin

To reveal the effect of white-rot fungi treatment on the structural changes of lignin, MWL1, MWL2, MWL3, and MWL4 from *T. pubescens* C7571-treated poplar were further investigated. The sugar analysis of the extracted lignin fractions is listed in Table 1. The lignin fraction from the untreated sample had a relatively higher amount of sugar (6.49%) than the treated samples. Comparatively, MWL4 obtained from treated samples during the last 16 weeks had the lowest amount of bound sugars (0.97%). This result might be caused by the destruction of linkages between lignin and carbohydrates by the hemicellulase in white-rot fungi (Dinis et al. 2009). In all cases, xylose was the predominant sugar (0.28 to 3.06%), followed by mannose, glucose, and galactose, with a small amount of arabinose. An increase in treatment time.
from 4 to 16 weeks resulted in a noticeable decrease in xylose content (from 0.92% to 0.28%).

The molecular weight distribution of the extracted lignin samples was analyzed using gel permeation chromatography (GPC). Changes in the molecular weights of lignin provide insights to lignin fragmentation and re-condensation reactions during bio-treatment. As shown in Table 2, all lignin fractions possessed narrow molecular weight distributions ($M_w/M_n$, lower than 1.5). The molecular weight of the MWL0 sample was $M_w$ 3585.6 g/mol and $M_n$ 2451.5 g/mol. When treated for 4 to 12 weeks, the extracted lignin had low molecular weights (2555.3 to 3226.4 g/mol and 1773.0 to 2250.0 g/mol). A significant reduction in molecular weight was observed in the lignin fraction MWL4 ($M_w$ 2555.3 g/mol and $M_n$ 1773.0 g/mol). The decrease in molecular weight indicated that lignin was fragmented during bio-treatment.

**FT-IR Spectra of the Extracted Lignin**

To better understand lignin structural changes, the extracted lignin fractions were characterized by FT-IR spectroscopy (Fig. 4). The spectra of samples from weeks 4, 8, 12, and 16 were rather similar. The band at 1024 cm$^{-1}$ was indicative of the aromatic C-H in-plane deformation. The strong band at 1219 cm$^{-1}$ was caused by C-C, C-O, and C=O stretching. Syringyl (S) and condensed guaiacyl (G) absorptions were clearly observed at 1329 cm$^{-1}$, whereas guaiacyl ring breathing with C-O stretching appeared at 1373 cm$^{-1}$. The bands at 1420, 1503, and 1591 cm$^{-1}$ corresponding to the aromatic skeletal vibrations, and the C-H deformation combined with the aromatic ring vibration at 1462 cm$^{-1}$ were present in these five spectra. The absorption at 1662 cm$^{-1}$ was attributed to the carbonyl stretching in conjugated p-substituted aryl ketones. A wide absorption band at 3452 cm$^{-1}$ originated from the OH stretching vibration in aromatic and aliphatic OH groups, whereas the bands at 2840, 2887, and 2937 cm$^{-1}$ arose from the C-H asymmetric and symmetrical vibrations in the methyl and methylene groups, respectively. The lignin degraded by white-rot fungi revealed an intensity at the 1688 to 1708 cm$^{-1}$ band was recorded in degraded lignin, which indicated unconjugated carbonyl groups. The results showed that during the bio-treatment of lignin, white-rot fungi generated conjugated and unconjugated C=O, which suggests that lignin biodegradation involved oxidation.

**2D HSQC NMR Analysis of the Extracted Lignin**

Two-dimensional 1H-13C NMR (2D NMR) provided important structural information and allowed for the resolution of otherwise overlapping resonances observed in either the 1H or 13C NMR spectra. To understand the detailed structural changes in the lignin fractions, they were characterized by 2D HSQC NMR. The HSQC NMR spectra of lignin showed two regions corresponding to the side chain (Fig. 5(a) and (b)) and aromatic 13C-1H correlations (Fig. 5(c) and (d)). The HSQC spectra of untreated wood lignin and treated lignin were similar, which meant that the main structure of lignin remained the same. In the side-chain regions of the HSQC spectra of these two lignin fractions, cross-signals of methoxyls ($\delta_C/\delta_H$ 56.0/3.75) and side chains in β-O-4′ aryl ether linkages were the most prominent. The $C_γ$-$H_γ$ correlations in β-O-4′ substructures were observed at $\delta_C/\delta_H$ 72.1/4.85 (structures A, A′, and A″ (Fig. 6). The $C_β$-$H_β$ correlations corresponding to the erythro and threo forms of the S-type β-O-4′ substructures were distinguished at $\delta_C/\delta_H$ 83.9/4.29 in structure A linked to the G/H lignin units and γ-acylated β-O-4′ aryl ether substructures (A′ and A″) linked to the S lignin unit. The $C_γ$-$H_γ$ correlations in structure A were observed at $\delta_C/\delta_H$ 59.5 to 59.7/3.41 to 3.64. In addition to the β-O-4′ ether substructures, β-β′ (resinol, B) and β-5′ (phenylcoumarin, C) linkages were observed. Strong signals for resinol substructures B were observed with $C_γ$-$H_γ$, $C_β$-$H_β$, and the double $C_γ$-$H_γ$ correlations at $\delta_C/\delta_H$ 85.0/4.70, 53.8/3.09, and 71.1/4.19/3.84, respectively. Phenyl coumarin substructures C were found in lower amounts. The signals for their $C_γ$-$H_γ$ and $Cβ$-$Hβ$ correlations were discovered at $\delta_C/\delta_H$ 87.0/5.49 and 53.1/3.47, respectively, whereas the $C_γ$-$H_γ$ correlations overlapped with other signals at approximately $\delta_C/\delta_H$ 62.5/3.73. In addition, $C_γ$-$H_γ$ correlations (at $\delta_C/\delta_H$ 61.4/4.10) in the p-hydroxycinnamyl alcohol end group (I) and various signals from the associated
carbohydrates and polysaccharide from fungi (δC/δH 65.0 to 77.1/2.9 to 4.2) were found in the side-chain regions of the lignin HSQC spectra.

In the aromatic regions of the HSQC spectra, cross-signals from the S- and G-lignin units were observed. The S-lignin units showed a prominent signal for the C₂₂.₆-H₂.₆, correlation at δC/δH 104.0/6.70, whereas the G units showed different correlations for C₂-H₂, C₅-H₅, and C₆-H₆, at δC/δH 111.1/6.99, 115.1/6.78, and 119.1/6.81, respectively. Signals corresponding to the C₂₂.₆-H₂.₆ correlations in Ca-oxidized S units (S′) (δC/δH 105.3/7.31) were present in all of the HSQC spectra of these two lignin fractions. The correlation for the C₂-H₂ in oxidized α-ketone structures G′ was observed in the spectra of MWL⁰ and became stronger in the spectra of MWL₄. The C₂₂.₆-H₂.₆ correlations of PB were observed as a strong signal at δC/δH 131.3/7.69. The signal at δC/δH 123.0/7.6 might be assigned to vanillic acid or its analogue (Huang et al. 1993; Ali et al. 2011), which was one of the lignin-degraded products.

The different structural features among the treated and untreated lignin fractions were quantitatively investigated (Fig. 5(e) and (f)). The percentages of lignin side chains involved in the primary substructures A-C and the lignin units were calculated from the corresponding HSQC spectra (Wen et al. 2013). As expected, the main substructures present in all lignin fractions were the β-O-4′ linked ones (A, A′, and A″), which ranged from 31.5/100 Ar to 76.4/100 Ar. The β-β′ resinol substructure (B) appeared to be the secondary major substructure, comprising 7.5/100 Ar to 18.0/100 Ar. The phenylcoumaran substructures (C) were calculated to be minor amounts, ranging from 1.9/100 Ar to 5.6/100 Ar. The content of β-O-4′ aryl ether in MWL⁰ was 76.4/100Ar, and it decreased to 31.5/100 Ar in MWL₄. These data suggested that the cleavage of the β-O-4′ aryl ether (depolymerization) is the predominant reaction during bio-treatment. This might be because of the high activity of LiP in T. pubescens C7571, which attacks the β-O-4′ linkage of lignin (Umezawa et al. 1987). In addition to β-O-4′ aryl ether linkage cleavage and carbon-carbon linkage degradation, the S/G ratio in lignin was another prominent structural alteration observed after bio-treatment. The S, G, and PB ratios decreased from 53.1/100 Ar, 46.9/100 Ar, and 4.5/100 Ar to 23.0/100 Ar, 17.9/100 Ar, and 0.6/100 Ar for MWL⁰ and MWL₄, respectively. Interestingly, the data also suggested that the G-type lignin was further degraded than the S-type, as revealed by a slightly elevated S/G ratio in MWL⁰ (1.13) and MWL₄ (1.29). Thus, G-type lignin was more easily degraded. Another possible reason is that the isolated ML was mainly extracted from the middle lamella of the cell wall (Whiting and Goring 1982), which is more easily penetrated by the hyphae of white-rot fungi. Because G-type lignin has less methoxyl than S-type lignin, the cross-signals of methoxyls (δC/δH 56.0/3.75) of MWL₄ were stronger.

**Factors Influencing Biological Pre-treatment**

Biological pre-treatment is not only involved in generating any inhibitors and environmentally friendly methods, but also, it’s a quietly time-consuming method. To enhance it by choosing the most efficient strain and culture conditions can make the method more effective by decreasing the treatment time and carbohydrate loss. Important process factors affecting biological pre-treatment comprise the nature, component of biomass, and other factors such as variety of organisms engaged in incubation time and temperature, acidity (pH), inoculums concentration, moisture content and aeration rate.

The optimum temperature during biological pre-treatment varies with the type of microorganism employed. Most of the white rot ascomycetes fungi grow optimally around 39°C while the white rot basidiomycetes grow optimally around 25°C and 30°C. The metabolism of these fungi generates heat and develops temperature gradients in solid state media. The accumulated heat can destroy or inhibit fungal growth and metabolism. Various optimal temperature for biological pre-treatment of biomass is because of fungal physiology, fungal strain and type of substrate. Incubation time requested for microorganisms pre-treatment differs depending on the strain and component of the biomass utilized for pre-treatment. Long incubation time due to low delignification rate is one of the major barriers for large scale application of biological pre-treatment.

Acidity (pH) plays a prominent function in the cultivation of fungi and it is very complicated to
control it in a solid culture condition. Production of lignolytic gene is influenced by the initial pH of the medium. In the pH of 4 to 5 most of the white rot fungi can grow properly and also the substrate acidity decrease their growth. Inoculum concentration performs a significant function in biological pre-treatment. The time required for the colonization of the substrate is affected by the type and amount of inoculums. Spores are the commonly used inoculum. Larger quantity of inoculum leads to shorter time for colonization of the substrate.

High substrate concentrations have to be used for biological pre-treatment to make the process economically viable. Generation of inhibitor compounds increase by using high dry material that may unfavourably influence sugar yield reduction. Hence pre-treatment to be carried out with a compromised condition to minimize the generation as well as accumulation of inhibitory compounds. Initial moisture content is essential for the establishment of microbial growth in the biomass. Initial moisture content critically affects the fungal growth and enzyme production and significantly affects lignin degradation [63]. The production and pH of lignolytic enzymes mainly affected by aeration. Aeration has many functions which are including oxygenation, CO\textsubscript{2} removal, heat dispersion, humidity conservation and also dispersal of volatile combinations produced during metabolism. Since lignin degradation is an oxidative process, oxygen availability is important for ligninase activity of white rot fungi. High aeration could improve delignification rate and hence controlled aeration is essential for improvement of biological pre-treatment. Efficiency of manganese peroxidase is not considerably influenced by aeration.

**Conclusion**
A surveying of the literature tells us that very little is known about the degradation of lignin in soil. Lignin is the most important and common aromatic organic compound found in the plant cell wall, and becomes a major source of humic acid in soil. Degraded lignin fragments are building blocks of the humic compounds in soil. Hence, lignin degradation has received vast attention from various researchers. The most efficient organisms for lignin mineralizing are white-rot fungi. Enzymes from specialized bacteria and fungi have been surveyed in this study that can metabolize lignin and enable utilization of lignin-derived carbon as a resource. The study summarizes lignin biodegradation by various fungi and bacteria present in plant biomass and soils that are capable of producing ligninolytic enzymes such as LiP, MnP, VP, and DyP. Most of these enzymes are substrate specific, in contrast to Lac activity which oxidizes a variety of substrates like polyphenols, diphenols, benzenethiol and aromatic amines. Recent efforts to identify new lignin-degrading microbes and thorough study of their genomics, biochemistry, and proteomics will uncover the role of ligninolytic enzymes in the coming years.

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