Ángel T. Martínez1*
Mariela Speranza1
Francisco J. Ruiz-Dueñas1
Patricia Ferreira1
Susana Camarero1
Francisco Guillén1,3
María J. Martínez1
Ana Gutiérrez2
José C. del Río2

1Biological Research Center, CSIC, Madrid, Spain
2Institute of Natural Resources and Agrobiology of Sevilla, CSIC, Spain
3Present address: University of Alcalá, Madrid, Spain

Received 24 March 2005
Accepted 15 May 2005
*Corresponding author:
A.T. Martínez
Centro de Investigaciones Biológicas, CSIC
Ramiro de Maeztu, 9
28040 Madrid, Spain
Tel. +34-918373112. Fax +34-925360432
E-mail: ATMartinez@cib.csic.es

Summary. Wood is the main renewable material on Earth and is largely used as building material and in paper-pulp manufacturing. This review describes the composition of lignocellulosic materials, the different processes by which fungi are able to alter wood, including decay patterns caused by white, brown, and soft-rot fungi, and fungal staining of wood. The chemical, enzymatic, and molecular aspects of the fungal attack of lignin, which represents the key step in wood decay, are also discussed. Modern analytical techniques to investigate fungal degradation and modification of the lignin polymer are reviewed, as are the different oxidative enzymes (oxidoreductases) involved in lignin degradation. These include laccases, high redox potential ligninolytic peroxidases (lignin peroxidase, manganese peroxidase, and versatile peroxidase), and oxidases. Special emphasis is given to the reactions catalyzed, their synergistic action on lignin, and the structural bases for their unique catalytic properties. Broadening our knowledge of lignocellulose biodegradation processes should contribute to better control of wood-decaying fungi, as well as to the development of new biocatalysts of industrial interest based on these organisms and their enzymes. [Int Microbiol 2005; 8(3):195-204]

Key words: wood-rotting fungi · lignin · analytical pyrolysis · oxidoreductases · catalytic mechanisms

Biodegradation of lignocellulosics: microbial, chemical, and enzymatic aspects of the fungal attack of lignin

Lignocellulosic materials
Forests represent approximately 27% of the world’s land area, and wood is the predominant commercial product from forests. Global wood consumption is around 3500 million m³/year, and has increased over 65% since 1960. More than half of this consumption is for fuel. The remainder of the global roundwood consumption is largely for pulp and paper products, building materials, and other wood in service.

Wood and other lignocellulosic materials are formed by three main polymeric constituents, cellulose, lignin, and hemicelluloses [25]. Cellulose is a linear and highly ordered (often crystalline) polymer of cellobiose (α-D-glucopyranosyl-β-1,4-D-glucopyranose) that represents over 50% of wood weight. By contrast, lignin is a three-dimensional network built up of dimethoxylated (syringyl, S), monomethoxylated (guaiacyl, G) and non-methoxylated (p-hydroxyphenyl, H) phenylpropanoid units, derived from the corresponding p-hydroxybenzyl alcohols, which give rise to a variety of subunits including different ether and C—C bonds. Acetylated lignin units have been recently identified in non-woody plants using analytical pyrolysis [7].

Lignin is highly resistant towards chemical and biological degradation, and confers mechanical resistance to wood. The highest concentration of this recalcitrant polymer is found in the middle lamella, where it acts as a cement between wood fibers, but it is also present in the layers of the cell wall (especially the secondary cell-wall), forming, together with hemicelluloses, an amorphous matrix in which the cellulose fibrils are embedded and protected against biodegradation [14]. Lignin composition in terms of the
H:G:S ratio varies between different vascular plant groups. Woody gymnosperms (softwoods) have the highest lignin content, and their lignin is made up mostly of G units. By contrast, lignin of woody angiosperms (hardwoods) consists of S and G units, and that from non-woody angiosperms contains also H units. Lignin composition between the different wood tissues and cell-wall layers also varies. For example, middle-lamella lignin typically has a lower S/G ratio than lignin from the secondary wall.

The third structural component, hemicelluloses (polymers), has an intermediate degree of complexity and is made up of different pentose and hexose residues, which are often acetylated, and generally form branched chains. Typically, hemicelluloses in softwood are glucomannans, whereas those in hardwoods are mainly xylans together with variable percentages of galactose, arabinose, rhamnose and methylglucuronic acid units, and acetyl groups. Other non-structural components of wood include compounds extractable with organic solvents (the so-called extractives) which can be either polar (e.g. phenols and tannins) or apolar (e.g. fats and sterols), water-soluble compounds (e.g. sugars and starch), as well as proteins and ashes. These components together generally represent less than 5% of the dry weight of wood but can reach 20% in some softwoods (e.g. in some Cupressaceae)\[14\].

The above constituents form the three main types of wood tissue elements, namely fibers, vessels, and parenchyma cells (Fig. 1A,B). In gymnosperms, wood tissues have a relatively simpler structure than in angiosperms; they consist of 90−95% tracheid cells (softwood fibers) and low amounts of parenchyma, which includes the specialized resin channels in conifers. Parenchymatic rays, which contain phenolic and lipophilic extractives and water-soluble compounds as storage material, have a radial arrangement in wood. Vessels are large cells with a longitudinal arrangement, and they are responsible for the transport of water and nutrients along the plant stem. Finally, fibers, which are also longitudinally arranged, represent most of the wood volume and are characterized by their thick cell walls, which provide support to the tree.

**Wood biodegradation**

Lignocellulose degradation is a central step for carbon recycling in land ecosystems. Moreover, fungal decay of wood in

![Fig. 1. Wood anatomy and fungal degradation. In vitro degradation of Eucalyptus globulus wood by Inocutis jamaicensis (using the soil-block test). Images obtained using low-temperature scanning-electron microscopy. (A, B): Transversal and tangential sections, respectively, of a hardwood showing large vessels (v), parenchymatic rays (r) and fibers (f) (initial degradation of wood with hyphae inside vessels in shown in B). (C): Simultaneous white-rot decay characterized by strong degradation of all cell-wall components, including abundant mycelium (with extracellular mucilage) inside a vessel. (D): White-rot selective delignification characterized by preferential degradation of lignin and separation of fibers due to destruction of the middle lamella. Black arrowheads in B indicate hyphae inside vessels, as shown in C. White arrowheads in C and D show cell-wall degradation and fiber separation, respectively. Bars: 500 µm in A and B, 50 µm in C, and 20 µm in D.](Image)
service results in billion-euro losses. Basidiomycetes are the main wood rotters due to their ability to degrade or modify lignin, an enzymatic process that originated in the Upper Devonian period in parallel with the evolution of vascular plants [12]. Wood-rotting basidiomycetes are classified as white-rot and brown-rot fungi based mainly on macroscopic aspects [42,49]. Table 1 summarizes the characteristics of wood attack by several types of fungi. Basidiomycetes can overcome difficulties in wood decay, including the low nitrogen content of wood and the presence of toxic and antibiotic compounds. Extracellular oxidative enzymes (oxidoreductases) secreted by fungi are involved in degradation of cell-wall components (see below). White-rot basidiomycetes, the most frequent wood-rotting organisms, are characterized by their ability to degrade lignin, hemicelluloses, and cellulose, often giving rise to a cellulose-enriched white material. Due to the ability of white-rot basidiomycetes to degrade lignin selectively or simultaneously with cellulose, two white-rot patterns have been described in different types of wood, namely selective delignification, also called sequential decay, and simultaneous rot [35] (Fig. 1C,D).

Table 1. Anatomical, chemical features of different types of wood decaying and staining fungi

<table>
<thead>
<tr>
<th>Decay aspect and consistency</th>
<th>White rot</th>
<th>Brown rot</th>
<th>Soft rot</th>
<th>Stain fungi</th>
</tr>
</thead>
<tbody>
<tr>
<td>Bleached appearance, lighter in color than sound wood, moist, soft, spongy, strength loss after advanced decay.</td>
<td>Brown, dry, crumbly, powdery, brittle consistency, breaks up like cubes, drastic loss of strength at initial stage of decay. Very uniform ontogeny of wood decay.</td>
<td>Soft consistency in wet environments. Brown and crumbly in dry environments. Generally uniform ontogeny of wood decay.</td>
<td>Discoloration sapwood areas (specks, spots and patches), blue (softwood), black (hardwood), red, or other colors. Discoloration due to colored hypha, or physiological response of tree against damage.</td>
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<tr>
<th>Host (wood-type)</th>
<th>Simultaneous rot</th>
<th>Selective delignification</th>
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</thead>
<tbody>
<tr>
<td>Hardwood, rarely softwood</td>
<td>Hardwood and softwood</td>
<td>Softwoods; seldom hardwoods. Forest ecosystems and wood in service.</td>
</tr>
</tbody>
</table>


| Anatomical features | Cell wall attacked progressively from lumen. Erosion furrows associated with hyphae. | Lignin degradation in middle lamella and secondary wall. Middle lamella dissolved by diffusion mechanism (not in contact with hyphae), radial cavities in cell wall. | Degradation at a great distance from hyphae (diffusion mechanism). Entire cell wall attacked rapidly with cracks and clefts. | Cell wall attack in the proximity of hyphae starts from cell lumen. Logitudinal biconical cylindrical cavities in secondary wall (Type 1). Secondary wall erosions from cell lumen (Type 2). Facultative soft-rot decay by some basidiomycetes. |

| Causal agents | Basidiomycetes (e.g. T. versicolor, I. lactea, P. chrysosporium and Heterobasidium annosum) and some Ascomycetes (e.g. Xylaria hyphoxylon). | Basidiomycetes (e.g. Ganoderma australe, P. tremella, C. subvermispora, Pleurotus spp. and Phellinus pini). | Basidiomycetes exclusively (e.g. G. puteana, Gloeophyllum trabeum, Laetiporus sulphureus, Piptoporus betulinus, Postia placenta and Serpula lacrimans). | Ascomycetes (Chaetomium globosum, Ustulina deusta) and Deuteromycetes (Alternaria alternata, Thielavia terrestris, Paecilomyces spp.), and some bacteria. Some white (Inonotus hispidus) and brown-rot (Rigidoporus crocatus) basidiomycetes cause facultative soft-rot decay. |

Ascomycetes (e.g. Ophiostoma and Ceratocystis spp.) and Deuteromycetes (e.g. Aureobasidium pullulans, Phialaphora sp. and Trichoderma spp.)

Based on Eriksson et al [12], Schwarze et al. [42], and Zabel and Morrell [49].
Brown-rot fungi, which grow mainly on softwoods, represent only 7% of wood-rotting basidiomycetes. This group of basidiomycetes can degrade wood polysaccharides after only a partial modification of lignin, resulting in a brown material consisting of oxidized lignin, which represents a potential source of aromatic compounds for the stable organic matter fraction in forest soils.

Although only white-rot and brown-rot basidiomycetes can degrade wood extensively, some ascomycetes and their asexual states, the so-called deuteromycetes, can colonize wood in contact with soil. This results in a decrease in the mechanical properties of wood, giving rise to so-called soft-rot, a process that often involves bacteria (Table 1). Soft-rot fungi can degrade wood under extreme environmental conditions (high or low water potential) that prohibit the activity of other fungi. Moreover, some basidiomycetes also cause a soft-rot-type decay pattern. Finally, a limited number of ascomycetous fungi, called stain fungi, can colonize wood through parenchymatic rays and resin channels causing discoloration of softwood tissues but a very limited degradation, which mainly affects extractives and water-soluble materials.

### Chemical analysis of degraded wood

The main wood degradation patterns at advanced stages of decay can be identified macroscopically and microscopically, as described above. General changes in the chemical composition of wood can be also observed after fungal-induced decay (Table 1). However, a precise analysis of the degradation type requires chemical analysis of cellulose and lignin contents and of the modifications in the decayed wood. Lignin in wood is traditionally estimated by the Klason method, which is based on total acid hydrolysis of polysaccharides and gravimetric estimation of the lignin content (after deducing ashes and protein). This method, however, is time-consuming, and requires a considerable sample volume. In addition, its application to samples containing modified lignin (as in rotted wood samples) is problematic because it has been developed for lignin in sound (undecayed) wood. Estimation of Klason lignin is often combined with analysis of wood polysaccharide composition by gas chromatography (GC) of the monosaccharides present in the acid hydrolysate.

Several modern spectroscopic and degradative methods have been used to analyze lignin or polysaccharides in wood [30]. Among them, pyrolysis coupled to gas chromatography-mass spectrometry (Py-GC/MS) has several great advantages as a technique enabling rapid analysis of small samples, yielding a precise identification of the proportion of H, G, and S lignin units [8]. Figure 2 shows a Py-GC/MS analysis of the fungal decay of a hardwood (*Eucalyptus globulus*). Two regions, corresponding to the polysaccharide- and lignin-derived compounds, can be distinguished in the pyrograms. Due to the higher stability of the degradation products of lignin (the main ones shown in Fig. 2D) than of polysaccharides, which yield several compounds that cannot be chromatographed, Py-GC/MS analyses result in overestimation of lignin. However, this method can be used to compare the changes in the relative proportion of lignin and polysaccharides in decaying wood. Moreover, the amounts of G-lignin and S-lignin (no H-lignin has been detected in eucalypt wood) can be separately estimated by Py-GC/MS analysis.

Figure 2A shows wood decay by the white-rot basidiomycete *Ceriporiopsis subvermispora*, which results in a relative decrease in the lignin peaks and a relative increase in the carbohydrate peaks. By contrast, the soft-rot deuteromycete *Paecilomyces* sp. decreases cellulose content to a larger extent than lignin content (Fig. 2B). Figure 3 shows wood degradation patterns by the basidiomycetes *Bjerkandera adusta*, *C. subvermispora*, *Coniophora puteana*, *Crepidotus variabilis*, *Fusarium oxysporum*, *Melanotus hepatochrous*, *Phanerochaete chrysosporium*, *Plebeia radiata*, and *Pleurotus pulmonarius* (Fig. 3A), the ascomycetes *Mollisia* sp, *Ophiostoma piliferum* (including an inoculum commercialized as *Cartapip*) and *Ophiostoma valdavianum* (Fig. 3B), and the deuteromycetes *Fusarium oxysporum*, *Kirramyces eucalypti* and *Paecilomyces* sp. (Fig. 3C), which can be differentiated based on their lignin/carbohydrate and S/G Py-GC/MS ratios. The highest cellulose enrichments are produced by the basidiomycetes *C. subvermispora* and *C. variabilis*, the former also causing the highest modification of lignin, as revealed by the S/G ratio. In addition, *Paecilomyces* sp. and other deuteromycetes produce an increase in the relative lignin content in wood (due to preferential removal of polysaccharides), whereas ascomycetes slightly modify wood composition, as revealed by the Py-GC/MS analysis.

### Enzymatic aspects

Lignin degradation and/or modification by basidiomycetes is the key step in lignocellulose decay. Therefore, the enzymes and mechanisms involved in lignin attack are described below. For a discussion of the subsequent steps in the degradation of wood polysaccharides, we recommend other reviews [36,43].

Laccases have been known for many years in plants, fungi, and insects, where they play a variety of roles, including synthesis of pigments, fruit-body morphogenesis, and detoxification [34]. Their production in fungal plate cultures was considered to be a characteristic unique to white-rot...
Fig. 2. Py-GC/MS analysis of the same hardwood degraded in vitro by the white-rot basidomycete \textit{C. subvermispora} (A) and the soft-rot deuteromycete \textit{Paecilomyces} sp. (B), and the corresponding control (C) showing carbohydrate derived (a–m) and lignin derived (1–26) compounds. (D) Chemical structure of the main G-type and S-type lignin marker compounds. \textit{E. globulus} wood chips were treated with the different fungi under solid-state-fermentation conditions. Adapted from del Río et al. [8]. Peak identification: \textit{a}, butanedial; \textit{b}, 1,4-pentadiene-3-one; \textit{c}, unknown; \textit{d}, 3-hydroxypropanal; \textit{e}, 3-furaldehyde; \textit{f}, (2H)-furane-2-one; \textit{g}, 2-furaldehyde; \textit{h}, hydroxymethylfuran; \textit{i}, (5H)-furane-2-one; \textit{j}, 2,5-dihydro-5-methylfuran-2-one; \textit{k}, 2,3-dihydro-5-methylfuran-2-one; \textit{l}, 4-hydroxy-5,6-dihydro-(2H)-pyran-2-one; \textit{m}, 2-hydroxy-3-methyl-2-cyclopenten-1-one; \textit{n}, guaiacol; \textit{o}, 2,3,4-dihydroxybenzaldehyde; \textit{p}, 4-methylguaiacol; \textit{q}, 4-ethylguaiacol; \textit{r}, 4-vinylguaiacol; \textit{s}, eugenol; \textit{t}, syringol; \textit{u}, cis-isoeugenol; \textit{v}, vanillin; \textit{w}, trans-isoeugenol; \textit{x}, 4-methylsyringol; \textit{y}, homovanillin; \textit{z}, acetoguaiacone; \textit{aa}, 4-ethoxyiseringol; \textit{ab}, guaiacylacetone; \textit{ac}, 4-vinlylsyringol; \textit{ad}, propiovannillone; \textit{ae}, 4-allylsyringol; \textit{af}, 4-propylsyringol; \textit{ag}, 4-allylsyringol; \textit{ah}, 4-propylsyringol; \textit{ai}, 4-allylsyringol; \textit{aj}, 4-propylsyringol; \textit{ak}, 4-propylsyringol; \textit{al}, 4-propylsyringol; \textit{am}, 4-propylsyringol; \textit{an}, 4-propylsyringol; \textit{ao}, 4-propylsyringol; \textit{ap}, 4-propylsyringol; \textit{aq}, 4-propylsyringol; \textit{ar}, 4-propylsyringol; \textit{as}, 4-propylsyringol; \textit{at}, 4-propylsyringol; \textit{au}, 4-propylsyringol; \textit{av}, 4-propylsyringol; \textit{aw}, 4-propylsyringol; \textit{ax}, 4-propylsyringol; \textit{ay}, 4-propylsyringol; \textit{az}, 4-propylsyringol.

Fig. 3. Hardwood biodegradation patterns: Py-GC/MS analysis of decayed wood (Fig. 2) permits the definition of degradation clusters corresponding to basidiomycetes (A), ascomycetes (B), and deuteromycetes (C) (an untreated control is also included). Adapted from del Río et al. [8].
basidiomycetes [27], although some brown-rot fungi produce laccase in liquid cultures [29]. These phenoloxidases have a low redox potential that allows direct oxidation only of phenolic lignin units, which often comprise less than 10% of the total polymer. The interest in laccases for biotechnological applications increased with the discovery of their ability to oxidize high redox potential substrates in the presence of synthetic mediators [3], which allows the degradation of xenobiotic compounds [40] and chlorine-free bleaching of paper pulp [4]. Natural mediators involved in lignin biodegradation remain to be identified, although some lignin-derived phenols could act as efficient laccase mediators [5].

Lignin peroxidase (LiP) and manganese peroxidase (MnP) were discovered in the mid-1980s in \textit{P. chrysosporium} and described as true ligninases because of their high redox potential [16,31]. LiP degrades non-phenolic lignin units (up to 90% of the polymer), whereas MnP generates Mn$^{3+}$, which acts as a diffusible oxidizer on phenolic or non-phenolic lignin units via lipid peroxidation reactions [26]. More recently, versatile peroxidase (VP) has been described in \textit{Pleurotus} [32,41] and other fungi as a third type of ligninolytic peroxidase that combines the catalytic properties of LiP, MnP, and plant/microbial peroxidases oxidizing phenolic compounds [24].

Other extracellular enzymes involved in wood lignin degradation are oxidases generating H$_2$O$_2$, and mycelium-associated dehydrogenases that reduce lignin-derived compounds. The former include the aryl-alcohol oxidase (AAO) described in \textit{Pleurotus eryngii} [18] and other fungi, and \textit{P. chrysosporium} glyoxal oxidase [28]. Fungal aryl-alcohol dehydrogenases (AAD) and quinone reductases (QR) are also involved in lignin degradation [20,21].

As shown in Fig. 4, laccases or ligninolytic peroxidases (LiP, MnP, and VP) produced by white-rot fungi oxidize the lignin polymer, thereby generating aromatic radicals (a) [12]. These evolve in different non-enzymatic reactions, including C4-ether breakdown (b), aromatic ring cleavage (c), Cα-Cβ breakdown (d), and demethoxylation (e) [22]. The aromatic aldehydes released from Cα-Cβ breakdown of lignin, or synthesized de novo by fungi (f, g) [21] are the substrate for H$_2$O$_2$ generation by AAO in cyclic redox reactions involving also AAD [19]. Phenoxy radicals from C4-ether breakdown (b) can repolymerize on the lignin polymer (h) if they are not first reduced by oxidases to phenolic compounds (i), as reported for AAO [33]. The phenolic compounds formed can be again reoxidized by laccases or peroxidases (j). Phenoxy radicals can also be subjected to Cα-Cβ breakdown (k), yielding

![Fig. 4. A scheme for lignin biodegradation including enzymatic reactions and oxygen activation, (for explanation see text). Updated from Gutierrez and Martinez [22].](image-url)
p-quinones. Quinones from \( g \) and/or \( k \) contribute to oxygen activation in redox cycling reactions involving QR, laccases, and peroxidases \( (l, m) \) [20]. This results in reduction of the ferric iron present in wood \( (n) \), either by superoxide cation radical or directly by the semiquinone radicals, and its reoxidation with concomitant reduction of \( \text{H}_2\text{O}_2 \) to hydroxyl free radical \( (\text{OH}^-) \) [17]. The latter is a very strong oxidizer that can initiate the attack on lignin \( (p) \) in the initial stages of wood decay, when the small size of pores in the still-intact cell wall prevents the penetration of ligninolytic enzymes [13]. Then, lignin degradation proceeds by oxidative attack of the enzymes described above. In the final steps, simple products from lignin degradation enter the fungal hyphae and are incorporated into intracellular catabolic routes.

**Molecular aspects**

Due to their potential use as industrial biocatalysts, the catalytic mechanisms of lignin-degrading oxidoreductases (including peroxidases, oxidases, and laccases) have been extensively investigated and their molecular structures have been described (Fig. 5).

LiP and MnP were the second and third peroxidases whose crystal structure was solved [38,39,44], just 10 years after their discovery in *P. chrysosporium*. These peroxidases catalyze the oxidation of the recalcitrant non-phenolic lignin units by \( \text{H}_2\text{O}_2 \). This is possible because of the formation of a high redox potential oxo-ferryl intermediate during the reaction of the heme cofactor with \( \text{H}_2\text{O}_2 \). This two-electron reaction allows the activated enzyme to oxidize two substrate units, being reduced to the peroxidase resting state (which reacts again with peroxide). The catalytic cycle, consisting of the resting peroxidase and compounds I (two-electron oxidized form) and II (one-electron oxidized form), is common to other peroxidases. However, two aspects in their molecular structure provide ligninolytic peroxidases their unique catalytic properties: (i) a heme environment, conferring high redox potential to the oxo-ferryl complex; and (ii) the existence of specific binding sites (and mechanisms) for oxidation of their characteristic substrates, including non-phenolic aromatics in the cases of LiP, manganese iron in the case of MnP, and both types of compounds in the case of the new VP.

Similar heme environments in the above three peroxidases (located at the central region of the protein, Fig. 5A) have been evidenced by \(^1\text{H}-\text{NMR}, \) which allows the signals

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**Fig. 5.** Molecular structures of enzymes acting synergistically for lignin biodegradation. (A) Crystal structure of versatile peroxidase at 1.13 Å resolution including heme cofactor and both Mn\(^{2+} \) (right) and aromatic substrate (left) oxidation sites. (B) Homology molecular model of AAO, a flavoenzyme providing \( \text{H}_2\text{O}_2 \) to ligninolytic peroxidases [46]. (C) Crystal structure of active laccase at 1.90 Å resolution with its three catalytic copper.
of both the heme cofactor protons and several amino acid residues forming the heme pocket to be identified [1]. This is possible due to the paramagnetic effect caused by the cofactor iron, which displaces the signals of neighbor protons outside the region where most proton protons overlap. One of the main differences observed among peroxidases is the position of a protein iron ligand, the Ne of the side-chain of a histidine residue (the so-called proximal histidine). In ligninolytic peroxidases, this residue is displaced away from the heme iron, increasing its electron deficiency and increasing the redox potential of the oxo-ferryl complex [31].

In addition, recent studies have contributed to identification of the substrate binding sites in ligninolytic peroxidases. The aromatic substrate binding site and the manganese binding site were first identified in LiP and MnP [9,16,45], and then confirmed in the crystal structure of VP solved at atomic resolution (unpublished). These studies revealed that the novel catalytic properties of VP are due to its hybrid molecular architecture, as suggested several years before [6,41]. Mn²⁺ oxidation is produced at a binding site near the cofactor, where this cation is bound by the carboxylates of three acidic residues, which enables direct electron transfer to one of the heme propionates (Fig. 5A right). By contrast, veratral alcohol (and other lignin model substrates) are oxidized at the surface of the protein by a long-range electron transfer mechanism that initiates at an exposed tryptophan residue (Fig. 5A left). The rationale of the existence of this electron transfer mechanism is related to the fact that many LiP/VP aromatic substrates, including the lignin polymer, cannot penetrate inside the protein to transfer electrons directly to the cofactor. Therefore, these substrates are oxidized at the enzyme surface, and electrons are transferred to the heme by a protein pathway.

The H₂O₂ responsible for oxidative degradation of lignin is generated by extracellular fungal oxidases, which can reduce dioxygen to peroxide in a catalytic reaction. Flavin cofactors are generally involved in this reaction, as in the Pleurotus flavoenzyme AAO, although glyoxal oxidase from P. chrysosporium is a copper-containing enzyme [47]. Among flavoenzyme oxidases, fungal glucose oxidase has been crystallized [48], but this is an intracellular enzyme that is not involved in lignin degradation. However, its crystal structure has been used as a template to predict the molecular structure of AAO (shown in Fig. 5B, including the FAD cofactor) [46]. It has been shown that AAO is a unique oxidase due to both its spectroscopic characteristics (flavin intermediates and reactivity) and to the wide range of aromatic and aliphatic polysaturated primary alcohols (and even aldehydes) that it is able to oxidize [15]. The molecular structure of AAO includes two catalytically active histidines near the N5 of the flavin ring, which might help electron transfer to/from the cofactor by acting as bases in the oxidation of aromatic alcohols (which would proceed via a hydride transfer mechanism) and as acids in the reduction of oxygen to H₂O₂.

As noted above, laccases were the first ligninolytic enzymes to be investigated, and had been known in plants for many years. Nevertheless, the first molecular structure of a complete fungal laccase was published only in 2002. That year, the crystal structures of the laccases from the basidiomycete T. versicolor, and the ascomycete Melanocarpus albomyces were reported [2,23,37] (Fig. 5C). The first structure of a bacterial laccase was published one year later [11]. A previously reported laccase structure corresponded to an inactive form due to the loss of a copper ion during deglycosylation to obtain suitable crystals for X-ray diffraction [10]. The active site of laccases includes four copper ions. Type-I copper (right sphere in Fig. 5C) acts as electron acceptor from substituted phenols or amines (the typical laccase substrates); and type-II copper, which transfers the electrons to the final acceptor, dioxygen, which is reduced to water. The two type-III coppers act as intermediates in the electron transfer pathway that also includes one cysteine and two histidine protein residues. The molecular environment of laccase type-I copper seems to regulate the redox potential of the enzyme [37]. The fact that laccase can use atmospheric oxygen as the final electron acceptor represents a considerable advantage for industrial and environmental applications compared with peroxidases, which require a continuous supply of H₂O₂. Taking into account that the advantage of peroxidases is their higher redox potential, engineering the active site of laccases to obtain high redox potential variants would be of considerable biotechnological interest.

Most enzymes involved in wood lignin degradation (a multienzymatic process that includes, among others, peroxidases, oxidases, and laccases acting synergistically) have been identified, and the mechanisms of action of several of them have been established at a considerably precise level. These enzymes, however, cannot penetrate the compact structure of sound wood tissues due to their comparatively large molecular size. Therefore, small chemical oxidizers, including activated oxygen species and enzyme mediators, are probably involved in the initial steps of wood decay.

Acknowledgements. These studies have been partially supported by ENCE (Spain), by Spanish projects AGL2002-393 and BIO2002-1166, by EU projects QLK5-99-1357 and QLK3-99-590, and by an EU/FORES(ENCE)-PTD/MEC(Uruguay) grant. Carmen Ascaso (CCMA, CSIC, Madrid) is acknowledged for low-temperature scanning-electron microscopy facilities. Klaus Piontek (ETH, Zurich) is acknowledged for solving the VP crystal structure. Lina Bettucci (Universidad de la República, Montevideo) is acknowledged for an Inocutis jamaicensis strain. M.S acknowledges MEC for a Postdoctoral Fellowship. F.J.R.-D. thanks CSIC for an I3P contract. A.G and S.C. thank MEC for their “Ramón y Cajal” contracts.
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**Biodegradación de la lignocelulosa: aspectos microbiológicos, químicos y enzimáticos del ataque fúngico a la lignina**

**Resumen.** La madera es el principal material renovable en la Tierra y es utilizada en gran parte como material de construcción y en la fabricación de celulosa. Esta revisión describe la composición de los materiales lignocelulósicos y diversos procesos de alteración de la madera por parte de hongos, como el deterioro causado por los llamados hongos de podredumbre blanca, de podredumbre parda y de podredumbre blanda y por los hongos cromógenos o manchadores de la madera. También se tratan los aspectos químicos, enzimáticos y moleculares del ataque a la lignina por los hongos, que es clave en el deterioro de la madera. Se describen las técnicas analíticas modernas para investigar la degradación y la modificación del polímero de la lignina causadas por hongos, así como las diversas enzimas oxidativas (oxidoreductasas) que intervienen en la degradación de la lignina. Entre éstas se encuentran lacasas, peroxidasas ligninolíticas de alto potencial redox (lignina-peroxidasa, manganeso-peroxidasa y peroxidasas versátiles) y las oxidasas. Se destacan las reacciones catalizadas, su acción sinérgica sobre la lignina, y las bases estructurales de sus exclusivas propiedades catalíticas. Un mejor conocimiento de los procesos de biodegradación de la lignocelulosa debería contribuir a un mejor control de los hongos descomponedores de la madera, así como al desarrollo de nuevos biocatalizadores de interés industrial basados en estos organismos y sus enzimas. [Int Microbiol 2005; 8(3):195-204]

**Palabras clave:** hongos descomponedores de madera · lignina · pirólisis analítica · oxidoreductasas · mecanismos catalíticos

**Biodegradação da lignocelulose: aspectos microbiológicos, químicos e enzimáticos do ataque fúngico à lignina**

**Resumo.** A madeira é o principal material renovável na Terra e é utilizada em grande parte como material de construção e na fabricação de celulose. Esta revisão descreve a composição dos materiais lignocelulósicos e diversos processos de alteração da madeira por parte de fungos, como a deterioração causada pelos chamados fungos de podridão branca, de podridão parda e de podridão mole e pelos fungos emboloradores ou manchadores da madeira. Também se tratam os aspectos químicos, enzimáticos e moleculares do ataque à lignina pelos fungos, que é chave na deterioração da madeira. Se descrescem as técnicas analíticas modernas para investigar a degradação e a modificação do polímero da lignina causadas por fungos, assim como as diversas enzimas oxidativas (oxidoreductasas) que intervêm na degradação da lignina. Entre estas se encontram lacasas, peroxidasas ligninolíticas de alto potencial redox (lignina-peroxidasa, manganeso-peroxidasa e peroxidasas versáteis) e as oxidasas. Se destacam as reações catalisadas, sua ação sinérgica sobre a lignina, e as bases estruturais de suas exclusivas propriedades catalíticas. Um melhor conhecimento dos processos de biodegradação da lignocelulose deveria contribuir para um melhor controle dos fungos descomponedores da madeira, assim como ao desenvolvimento de novos biocatalisadores de interesse industrial baseados nestes organismos e suas enzimas. [Int Microbiol 2005; 8(3):195-204]

**Palavras chaves:** fungos descomponedores de madeira · lignina · pirólise analítica · oxidoreductasas · mecanismos catalíticos