REVIEW ARTICLE



Lignin pyrolysis reactions

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Abstract Lignin, an aromatic constituent of woody biomass, is a potential renewable aromatic feedstock for a sustainable future carbon economy. Pyrolysis-based technologies, such as fast pyrolysis and gasification, are promising methods for converting lignin into biochemicals, biomaterials, and biofuels. A better understanding of the molecular mechanisms involved in lignin pyrolysis/gasification would guide the development of the controlled pyrolysis and gasification systems to overcome issues with low product selectivity, an intrinsic drawback of current pyrolysis-based technologies. This review article summaries the state-of-the-art research into molecular mechanisms of lignin pyrolysis and gasification. This information should also be useful for understanding the influence of high temperature heat treatments on the properties of wood.

Keywords Lignin · Pyrolysis · Molecular mechanism · Homolysis · Heterolysis

Introduction

Many renewable energy options are available including photovoltaic, wind, and geothermal electricity generation. However, biomass resources are the only renewable source of carbon for use in chemicals and materials. Large amounts of the earth's biomass resources are accumulated in forest

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ecosystems, with an annual turnover of approximately 6% through photosynthesis [1]. Lignin, which accounts for 20–35% of wood (by weight), is an important renewable aromatic feedstock along with cellulose and hemicellulose, the polysaccharide components. Thus, it is important to establish efficient lignin utilization technologies.

Pyrolysis is defined as thermal degradation under a limited amount of oxygen and pyrolysis-based technologies show great promise for converting lignin and other wood components into biochemicals, biomaterials, and biofuels. For example, fast pyrolysis of wood, with a short residence time for volatile intermediates in the heating zone, produces good yields of an organic liquid that contains useful lignin-derived aromatic chemicals, which may then be separated [2, 3]. However, due to the low product selectivity, the necessary separation processes are tedious and costly, which limits the practical use of pyrolysis methods. Gasification provides an option to produce synthetic petroleum from wood via the conversion of syngas (CO and H₂) on Fischer–Tropsch catalysts. However, problems arising from the formation of tar and coke (solid carbonized substances converted from tar) are major drawbacks for establishing reliable wood gasification systems [4-6]. These substances cause pipeline clogging, catalyst deactivation and other issues. Lignin is considered to produce aromatic tar and coke during gasification. A better understanding of the molecular mechanisms involved in wood pyrolysis and gasification may guide the development of more efficient and reliable systems by addressing the aforementioned drawbacks.

Although it is not easy to define classes of pyrolysis and gasification reactions, such reactions may be generally divided into two stages, i.e., primary pyrolysis and secondary pyrolysis reactions [7]. Wood constituent polymers decompose into volatile and char (solid carbonized



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material) intermediates in the primary pyrolysis stage, which are then degraded further in the secondary pyrolysis reaction stage. Accordingly, this review article also follows this classification.

Effect of temperature on pyrolysis products

Lignin is a polymer of phenylpropane units containing three different aromatic ring substitution patterns: p-hydroxyphenyl (H), guaiacyl (4-hydroxy-3-methoxyphenyl, G) and syringyl (3,5-dimethoxy-4-hydroxyphenyl, S) depending on the wood species [8]. Softwoods contain a greater proportion of G units and smaller amounts of the H units, whereas hardwoods consist of mainly G and S units. These monomers are linked together through various kinds of ether linkages (C-O) and condensed (C-C) bonds, which leads to a heterogeneous chemical structure for lignins. Owing to these heterogeneous features, the behavior of lignins during pyrolysis reactions depends on the pyrolysis temperature. As illustrated in Fig. 1 [9], thermogravimetric (TG) analysis of lignin shows several derivative TG (DTG) peaks at 350 °C from primary pyrolysis reactions and at 400–450 °C (methoxyl group-related reactions) and at 550-600 °C (gasification of catechols (1,2-dihydroxybenzenes)) from secondary pyrolysis reactions. This is in contrast to the pyrolysis of cellulose, a homogeneous polymer of D-glucose units connected via β -1 \rightarrow 4 linkages, which exhibits only one sharp DTG peak around 350 °C.

The chemical structures of the pyrolysis products of lignins have been evaluated by gas chromatography/mass spectrometry (GC/MS) [10–12], nuclear magnetic resonance (NMR) [12–18] and infrared (IR) spectroscopic [15, 19–21] analyses, along with pyrolysis directly coupled with GC/MS [22–35] and IR [19, 36–39] (Py-GC/MS, Py-IR). Aromatic methoxy groups are stable during the primary pyrolysis stage and become very reactive in the temperature

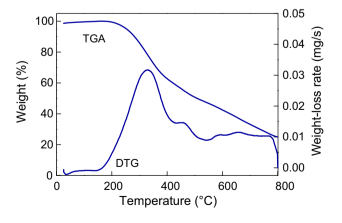


Fig. 1 Thermogravimetric analysis of milled wood lignin isolated from Japanese cedar (*Cryptmeria japonica*) [9]



range of 400–450 °C. Thus, the aromatic compounds produced during the primary pyrolysis stage are predominantly 4-substituted guaiacols (2-methoxyphenols) from G-lignins (Fig. 2) and 4-substituted syringols (2,6-dimethoxyphenols) from S-lignins. The majority of side-chains are unsaturated alkyl groups (i.e., alkyl groups containing>C=C<, >C=O and conjugated>C=O bonds) along with a smaller amount of saturated alkyls groups. The major volatile products from G-lignins in this stage include coniferyl alcohol (R: $-CH=CH-CH_2OH$), coniferyl aldehyde (R: -CH=CH-CHO), isoeugenol (R: -CH=CH-CHO), acetovanillone (R: $-CO-CH_3$), and dihydroconiferyl alcohol (R: $-CH_2-CH_2-CH_3-CH_3$).

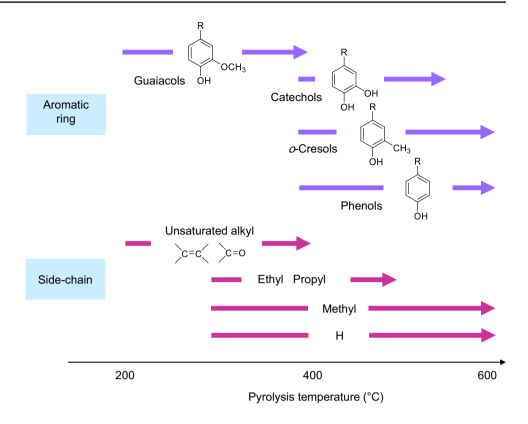
When the pyrolysis temperature is increased to 400-450 °C, secondary pyrolysis reactions take place and guaiacols/syringols rapidly transition to catechols (2-hydroxyphenols)/pyrogallols (2,3-dihydroxyphenols) and o-cresols (2-methylphenols)/xylenols (dimethylphenols) along with phenols (Fig. 2). In this temperature range, cracking of side-chain C-C bonds occurs, which increases the yields of monomers. The reaction products transition from unsaturated to saturated alkyls side chains (methyl, ethyl, propyl, 3-hydroxypropyl and others) and non-substituted (-H) types. Coke and polycyclic aromatic hydrocarbons (PAHs) formation also starts. Around 550 °C, catechols and pyrogallols disappear and non-condensable gas yields (particularly CO) significantly increase. At temperatures>700 °C the formation of PAHs is enhanced. Phenols and o-cresols are relatively stable at such high temperatures; hence, these compounds are observed along with PAHs even during high-temperature pyrolysis.

Primary pyrolysis reactions (200–400 °C)

Primary pyrolysis reactions of lignins occur over a wide temperature range of 200–400 °C with a DTG peak around 350 °C. As noted, lignins are heterogeneous polymers that arise from the inclusion of various types of linkages between phenylpropane units. As an example, spruce (a softwood) milled wood lignin (MWL) is reported to include 63–67% ether linkages (including 48% β -ether and 11.5–15% α -ether linkages) and 30–35% condensed linkages (including 9.5–11% 5–5′ (biphenyl) and 9–12% β -aryl] [8]. The proportions of ether linkages in hardwood lignins are normally higher than those of the softwood lignins, as can be seen from the 60% β -ether linkage proportion in birch (a hardwood) MWL [8]. Accordingly, the roles of these linkages during pyrolysis are important for understanding the primary pyrolysis reactions of lignins.

The use of model dimers that represent lignin ether and condensed type linkages is an effective means for

Fig. 2 Effect of pyrolysis temperature on aromatic substitution pattern and side-chain structure of the products from G-type lignin



understanding the pyrolytic reactions of lignins. Hence, various model dimers including β -ether [20, 27, 28, 34, 40–53], α -ether [20, 30, 34, 45, 47, 49, 51, 54], β -aryl [20, 31, 32, 47, 54], β - β [33], and biphenyl [20, 47, 54] types, trimers [34, 49, 51] and oligomers/polymers such as dehydrogenative polymerization products of cinnamyl alcohols [35] and β -ether type polymers [55–57] have been used in pyrolysis studies. Phenolic (Ph) and non-phenolic (Non-Ph) model compounds are commonly used to represent the ends and repeating units of lignin macromolecules, respectively.

Early studies conducted by the Domburg group [54, 58] reported the thermal stability of various model compounds, which were representative of the linkages between phenylpropane-units and side-chains. Analyses of the pyrolysis products of dimers along with NMR analysis of the pyrolyzed lignins revealed that α - and β -ether bonds are readily cleaved during the primary pyrolysis stage; however, condensed (C-C) type linkages are stable during depolymerization of lignin macromolecules [13, 20, 47]. The Ph β -aryl type dimers give stilbenes selectively, but this reaction does not lead to depolymerization of lignins [20, 47]. Normally, Ph dimers are more reactive than corresponding Non-Ph ones [20, 40, 41, 47, 49, 53]. Compared with hardwood lignins, softwood lignins contain a greater proportion of condensed type linkages. Thus, the differences in reactivity of these components can explain the observation that pyrolysis of softwood lignins tend to generate larger amounts of residue than hardwood lignins [59, 60].

Bond dissociation energy

Homolysis of C–C and C–O bonds to form a pair of radicals is one possible reaction that can explain pyrolytic lignin depolymerization. Recently, many papers have reported theoretical calculations of the bond dissociation energies (BDEs) for linkages in lignin related compounds [61–70]. These include calculations of α -O-4, β -O-4, 4-O-5, β -aryl, phenylcoumaran, pinoresinol, dibenzodioxocin type dimers. As summarized in Fig. 3 [63–66, 68, 71], α -O-4 type linkages had the weakest bond energies, with BDEs in the range of 40–60 kcal mol⁻¹, while the other bond types exhibited BDEs higher than 60 kcal mol⁻¹. The activation energy (E_a) for homolysis is considered to be slightly greater than the BDE; thus, the E_a for homolysis of these bonds can be expected to be greater than 60 kcal mol⁻¹, except for the α -O-4 type linkages.

Homolysis of the O-CH₃ bonds in lignin aromatic rings can occur by increasing the temperature to around 450 °C (i.e., the secondary pyrolysis reaction stage). This indicates that O-CH₃ bonds are stable during the primary pyrolysis stage (200–400 °C), which is also supported by results that show the primary pyrolysis products continue to bear their original aromatic methoxy groups. However, the BDE of the O-CH₃ bond in guaiacol has been measured as 56.3 kcal mol⁻¹ [71], which is lower than the above mentioned BDEs (>60 kcal mol⁻¹). These considerations indicate that direct homolysis of the C-O and C-C bonds



Fig. 3 Bond dissociation energies of various C-O and C-C bonds in lignin model compounds as evaluated by theoretical calculations

in lignin would not occur during the primary lignin pyrolysis stage except for α -O-4 type linkages. Thus, some effects that reduce the cleavage temperatures must be involved in these reactions. Many attempts have been made to model the weight-loss behaviors of lignin during pyrolysis; however, the measured $E_{\rm a}$ values arise from various pyrolysis reactions, including re-polymerization of the primary products, which reduce the weight-loss rate.

Homolysis and heterolysis mechanisms

Pyrolytic cleavage of C-O and C-C bonds can be considered in terms of both homolytic and heterolytic mechanisms, because these mechanisms give the same products as illustrated in Fig. 4 [an example of phenethyl phenyl ether]. The lack of reliable mechanistic evidence has given rise to a long standing controversy concerning which of the homolytic and heterolytic mechanisms take place. Experimental and theoretical investigations have recently been conducted to address this issue.

The theoretical approach provides useful information including $E_{\rm a}$ for the modelled reactions. Although these $E_{\rm a}$ values are a useful guide for understanding lignin pyrolysis reactions, a calculated reaction may not occur in reality if other competing reaction pathways with lower energy-barriers are available.

The effects of substituents on the cleavage reactivity provide more direct experimental evidence for a particular

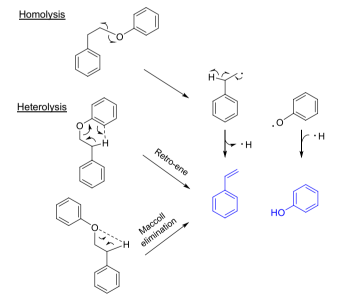
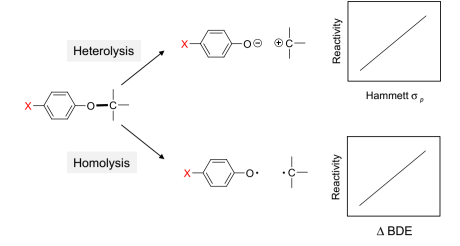


Fig. 4 Homolytic and heterolytic reactions for the conversion of phenethyl phenyl ether into styrene and phenol, as a model of the lignin β -ether structure

reaction pathway [49]. For example, substituents in the *para* positions of an aromatic ring have opposing effects on the heterolytic and homolytic cleavage of ether bonds (Fig. 5). Thus, the associated mechanisms are clearly indicated by plots of the cleavage reactivities against a



Fig. 5 Aromatic substituent effect as a tool for studying homolytic and heterolytic reactions of lignin ether bonds, based on parameters including Hammett σ_p and ΔBDE (bond dissociation energy)



Hammett's substituent constant (σ_p) and against the ΔBDE (the reduction in the BDE induced by the substituent).

For the heterolytic reaction pathway, the ether linkage is cleaved to form a para substituted phenolate anion. Thus, the reactivity of this pathway should increase as the electron-withdrawing ability of the substituent group increases, stabilizing the anion which results from bond cleavage. This property is quantified as σ_n . Conversely, homolysis of the ether bond forms a phenoxy radical and the reactivity depends on the stability of this species. When a stable radical is formed, cleavage of the ether linkages tends to proceed efficiently, because the increased stability of the radical reduces the BDE of the aromatic O-C bond. The stabilization obtained from substituents is, therefore, reflected in the ΔBDE value. When the phenoxy radical is stabilized by electron donating substituents, σ_p and ΔBDE values tend to increase in the opposite order to that of a heterolytic reaction.

Ether cleavage mechanisms

As discussed above, cleavage of α - and β -ether bonds plays an important role in lignin depolymerization during the primary pyrolysis stage. Hence, many studies have focused on the pyrolytic cleavage of the β -ether bond, which is the most abundant linkage type in lignin macromolecules.

Several concerted (heterolytic) mechanisms have been proposed for β -ether bond cleavage [42, 72–76]. Klein and Virk [42] proposed a six membered retro-ene mechanism based on an analysis of the kinetics for the formation of styrene and phenol from phenethyl phenyl ether. This system is representative of the simplest model compound without any aromatic ring substituents or side chains. In recent theoretical studies Elder and Beste [76] calculated E_a values for retro-ene and Maccoll elimination type concerted mechanisms (Fig. 4), with values of 62–68 kcal mol 1^{-1} and 56–58 kcal mol $^{-1}$, respectively. A similar range of values

have been reported by other researchers [73–75]. The Nimlos's research group [72] reported the formation of degradation products from the gas-phase pyrolysis of phenethyl phenyl ether and they concluded that the C–O homolysis occurred at high temperatures (>1000 °C), whereas the concerted retro-ene and Maccoll mechanisms were significant at lower temperatures (<600 °C).

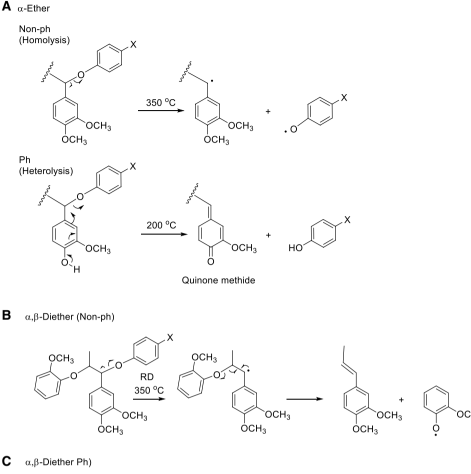
On the basis of the substituent effects at the *para* positions of the C_{α} -phenoxyl group (Fig. 5), the α -ether bonds in Ph and Non-Ph (methylated) dimers were confirmed to undergo heterolytic and homolytic cleavage, respectively (Fig. 6A) [49]. These results are consistent with the temperatures required for cleavage of this bond, which vary depending on the presence of Ph (200 °C) and Non-Ph (350 °C) type dimers [20]. These results suggested that a push and pull reaction to generate a quinone methide intermediate led to efficient heterolysis of the C_{α} -O bond in the Ph dimer. The low-temperature homolysis of the α -ether bond is consistent with the relatively low calculated BDE 40–60 kcal mol $^{-1}$, as described above.

Direct homolysis of the β -ether bond has been clearly indicated from substituent effects on the cleavage reactivities of Ph and Non-Ph C_{γ} -deoxy type dimers with *para* substituents for the C_{β} -phenoxy group (-H, -OCH₃, -Cl) as studied in a sealed reactor with tetralin that acts as a hydrogen donor [50]. However, the temperature (400 °C) required to cleave β -ether bonds in C_{γ} -deoxy type dimers could not explain the greater reactivity of guaiacylglycerol- β -guaiacyl ether, a Ph β -ether dimer bearing OH at C_{γ} which has an onset temperature of 250 °C [20].

The pyrolytic reactivities of Ph and Non-Ph α,β -diether type trimers that include various substituents at the *para* position of the C_{α} -phenoxy group have provided some insight into the cleavage mechanisms of β -ether bonds (Fig. 6b, c) [49]. The reactivities of the β -ether bonds depend solely on those of the α -ether bonds, demonstrating that cleavage of the α -ether bond acts as the



Fig. 6 Cleavage mechanisms of lignin ether bonds as suggested by the substituent effects on the reactivities of *para* substituted model compounds



Quinone methide

RD: rate-determning step

rate-determining step. A β -scission type reaction involving a C_{α} -radical, formed by the homolytic cleavage of the C_{α} -O bond, simultaneously cleaves the C_{β} -O bond homolytically (Fig. 6b). As noted, heterolysis of the α -ether bond in Ph compounds forms a quinone methide intermediate at 200 °C, which results in simultaneous homolytic cleavage of the β -ether bond in the same temperature range. This result is supported by the observation that the cleavage reactivity is unaffected by the addition of tetralin as a radical scavenger [50], along with the reduced reactivity of the C_{α} -deoxy dimer [45]. Low temperature homolysis of the C_{β} -O bond, via a quinone methide intermediate, has also been postulated under solvolysis delignification [77–79] and steam-explosion [80] conditions. Ponomarev [81]

calculated the BDE of the C_{β} –O bond in the quinone methide form to be 44.1 kcal mol⁻¹, a value that is much lower than that of the phenolic form (57.0 kcal mol⁻¹).

The reactivity of guaiacylglycerol- β -guaiacyl ether is lowered to the level of a Non-Ph dimer by elimination of the hydroxyl group from the C_{γ} , [45, 48]. This unexpected result is explained by the missing stabilization effect on the transition state of the quinone methide formation through hydrogen bonding between the C_{α} and C_{γ} hydroxyl groups. If the quinone methide mechanism occurred effectively during lignin pyrolysis, β -ether bonds would cleave simultaneously via the newly formed phenolic end groups. However, a careful comparison of the pyrolytic reactivities for model dimers, MWL, and methylated MWLs indicates that



cleavage of the β -ether linkages via the quinone methide mechanism does not proceed very efficiently in lignins, likely because of polymeric effects [20].

Radical chain mechanisms have been postulated for the cleavage of lignin β -ether linkages. The Brit and Beste research group have extensively studied hydrogen abstraction reactions from the $C_{\alpha}-H$ bond and subsequent cleavage of the β -ether bond using phenethyl phenyl ether and its derivatives [43, 44, 46].

The role of radical chain reactions is clear from the reactivities of the Ph and Non-Ph C_y-deoxy β-ether dimers, which vary substantially depending on the reactor type, (i.e., open-top and sealed reactors [50]. Although the dimers are recovered without undergoing pyrolysis reactions in open-top reactors, which allows volatile products to exit the heating zone, the reactivities of these compounds are dramatically increased in sealed reactors. Furthermore, the addition of tetralin effectively suppresses the reactivities in sealed reactors. From analysis of the products, two radical chain pathways (Fig. 7) have been suggested, which start from the C_{α} and phenoxy radicals formed via hydrogen abstraction of C_α-H and phenolic OH, respectively [51]. These reactions cleave β-ether bonds to form $C_\alpha = O$ and $C_\alpha = C_\beta$ type monomers via the β -scission type reaction of the C_{α} -radical and homolytic cleavage of the β-ether bond in the quinone methide intermediate, respectively. These pathways have been confirmed by the studies of kinetic deuterium isotope effects on product formation from regiospecifically deuterated dimers [53].

Pathway B occurs more effectively than pathway A, with onset temperatures of 260 and 360 °C, respectively [51]. This is explained by the number of radicals formed; three radicals are formed from the phenoxy radical intermediate in pathway B, which increases the radical concentration, whereas the number of radical species does not change in pathway A. Thus, this suggests an important role of the phenolic end groups as radical sensitizers. This proposal is supported by experimental results which have shown that the reactivity of pathway A of Non-Ph dimers significantly increases when Ph dimer is mixed into the system [53]. An α,β -diether trimer [1-(4-(3,4-dimethoxybenzoyloxy)-3-methoxyphenyl)-2-(2-methoxyphenoxy)-1-propanol], can be used to represent the benzyl ether derivative of the model dimer, and has an onset temperature of 320 °C. The reactivity of this species can be reasonably explained by the formation of a phenoxy radical intermediate of the dimer through homolysis of the benzyl ether (model α -ether bond) [51]. These cleavage temperatures coincide well with the DTG peak (350 °C), which corresponds to the primary pyrolysis of lignins.

Fig. 7 Proposed radical chain mechanisms for the cleavage of lignin ether bonds



Re-polymerization and side-chain conversion

The Ph β-ether dimer (guaiacylglycerol-β-guaiacyl ether) gives coniferyl alcohol in a 30.4 mol% yield, representing 60 mol% of the decomposition products of this dimer [47]. Cinnamyl alcohols, such as coniferyl alcohol, are the most important primary products in lignin pyrolysis. However, the contributions of cinnamyl alcohols in the pyrolyzates from wood and isolated lignins are much lower than expected from studies using model compounds. Coniferyl aldehyde, isoeugenol, dihydroconiferyl alcohol, 4-vinylguaiacol and vanillin are more important products (in the case of G-lignin pyrolysis). Only direct mass spectrometric analysis of the pyrolyzates from wood and lignin samples, without cooling, indicate significant contributions of coniferyl alcohol (MW: 180) and sinapyl alcohol (MW: 210) [7]. These apparently contradictory observations can be explained by the propensity of coniferyl alcohol and sinapyl alcohol to undergo secondary pyrolysis reactions, particularly polymerization reactions [82-85].

The relative evaporation/polymerization/side-chain conversion efficiency of *trans*-coniferyl alcohol has been studied by pyrolysis in an open-top reactor system (200–350 °C under nitrogen, heating period: 5 min) [85]. Evaporation of coniferyl alcohol started around 200–250 °C, and competed with secondary pyrolysis reactions, such as polymerization and side-chain conversion reactions. All these processes were enhanced at higher pyrolysis temperatures and completed at 300 °C, where approximately 70% of the polymerization products were formed along with evaporated coniferyl alcohol (15%) and side-chain conversion products (15%). Thus, at the DTG peak of 350 °C for lignin, secondary pyrolysis reactions, particularly re-polymerization reactions, reduce the coniferyl alcohol yield after it is formed, during the pyrolysis of G-type lignin.

It has been suggested that a quinone methide mechanism operates in the polymerization of coniferyl alcohol, based on the chemical structures of the dimers isolated from the pyrolysis of coniferyl alcohol in the presence of creosol (4-methylguaiacol) at 250 °C [84]. Hence, the polymerization reactivity of coniferyl alcohol is reduced by methylation of the phenolic hydroxyl group of the coniferyl alcohol. The stable methylated coniferyl alcohol is copolymerized with coniferyl alcohol during pyrolysis [85], which shows that coniferyl alcohol structures can add to the repeating phenylpropane units. The guaiacol derivatives with conjugated $C_{\alpha} = C_{\beta}$ side-chains are much more reactive than those with C_{α} -OR groups, which are the structures typically observed in natural lignins [84]. A radical chain vinyl condensation mechanism has been proposed to explain the polymerization of 4-vinylguaiacol and its methyl ether derivatives [84]. Thus, primary pyrolysis reactions that form side-chain double bonds promote condensation of natural lignins. Methylation of the phenolic hydroxyl groups of MWL prevents pyrolysis reactions that arise from the reactive phenolic end groups and considerably inhibits condensation reactions [20].

Competitive processes during primary pyrolysis

Lignin primary pyrolysis pathways (Fig. 8, shows G-type lignin pyrolysis) have been proposed based on the investigation of the influences of 1,2,3,10b-tetrahydrofluoranthene, a hydrogen donor, and diphenoxybenzene (DPB), an aprotic solvent [86, 87]. In DPB, the polymerization of coniferyl alcohol and sinapyl alcohol were effectively suppressed. Both DPB and H-donors were required for the effective formation of monomers from wood and MWLs, suggesting that lignin ether bonds undergo homolysis. The expected primary product of this reaction was a coniferyl alcohol radical that is stabilized by the H-donor. The H-donor also promotes side-chain reduction of the resulting coniferyl alcohol to form dihydroconiferyl alcohol and isoeugenol as the major monomers [86]. Under normal pyrolysis conditions, without the addition of any H-donors, the amount of H-donors is not sufficient to stabilize all the radical species formed through cleavage of ether linkages. Hence, primary radicals tend to undergo radical coupling reactions to form polymerization products.

The relative oxidation/reduction efficiency of sidechains of coniferyl alcohol varies depending on the pyrolysis temperature. At relatively low temperatures (for example 250 °C), coniferyl aldehyde, an oxidation product, is the major monomer from lignin pyrolysis [24, 86]. In this case, the pyrolysis environment is expected to be radical conditions, which enhance hydrogen abstraction reactions [86]. The C_{γ} -hydrogen atom of coniferyl alcohol, which is at the conjugated allyl position, is the main site of H-abstraction; the abstraction of hydrogen then leads to production of coniferyl aldehyde. However, by increasing the pyrolysis temperature to >350 °C, the pyrolysis environment



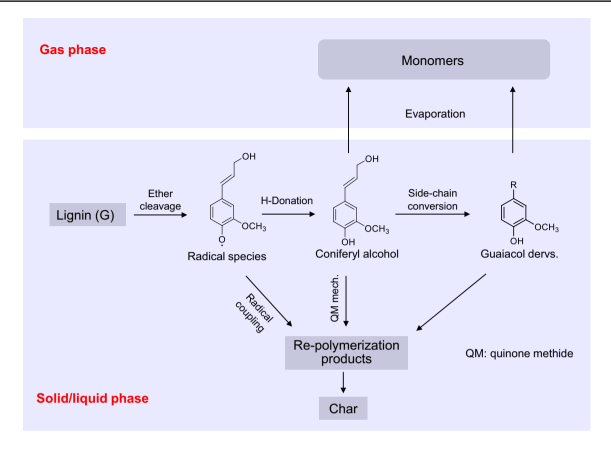


Fig. 8 Competitive pathways from coniferyl alcohol radical, a primary pyrolysis product, as proposed for the production of monomers and char during pyrolysis of G-type lignin at the primary pyrolysis stage (200–400 °C)

becomes richer in H-donor species (H-radicals), which results in increased monomer yields and the formation of side-chain reduction products such as dihydroconiferyl alcohol and isoeugenol.

Polymerization of the primary products by radical and quinone methide mechanisms can compete with side-chain conversion and evaporation processes [86]. Evaporation is the key step to recovering monomeric products. Otherwise, products polymerize and are finally converted to solid carbonized products. During the polymerization process, condensed (C–C) linkages are formed more preferably than ether (C–O) linkages based on the hard/soft, acid/base rule. The resulting polymerization products contain more C–C linkages, which are more resistant to further depolymerization.

The polymer effect also promotes polymerization and side-chain conversion of lignin primary pyrolysis intermediates [86]. Cleavage of ether linkages at the terminal endgroups of lignin macromolecules leads to direct formation of monomers, whereas the cleavage of ether linkages within the polymer, does not lead to immediate formation of monomers because of the polymeric nature of the intermediates formed. In these circumstances, pyrolyzates remain in the

heating zone and undergo polymerization and side-chain conversions. Thus, the monomer yield is reduced and the yield of char and side-chain conversion products increases.

Sinapyl alcohol, the corresponding primary product from S-type lignins, exhibits similar reactivity to that observed for the pyrolysis of coniferyl alcohol. However, sinapyl alcohol shows greater radical sensitivity at a relatively high pyrolysis temperature of 350 °C [87].

Secondary pyrolysis reactions (>400 °C)

Change in aromatic substitution pattern

As shown in Fig. 9, homolytic cleavage of the O–CH $_3$ bonds attached to the lignin aromatic rings occurs effectively at ~450 °C, associated with an *ipso* rearrangement starting from the phenoxy radical of guaiacol [88]. Intramolecular H-abstraction at the methyl group, by a phenoxy radical (reaction c) and subsequent 1,2-aryl migration (reaction e) have been proposed to account for this transformation [88, 89]. The 1,2-aryl migration product is further converted into o-quinone methide (reaction g),



Fig. 9 Pathways for the conversion of guaiacols, the primary pyrolysis products from G-type lignin, into catechols, o-cresols, phenols and coke at the secondary pyrolysis stage (>400 °C)

a key intermediate [90, 91], which is subsequently hydrogenated to *o*-cresol (reaction h). Demethoxylation occurs by α-scission of the formyl radical (reaction l) formed by H-abstraction from *o*-hydroxybenzaldehyde (reaction k) [88, 92]. Thus, syringols formed as primary pyrolysis products from S-type lignins are gradually converted into guaiacols and phenols [92]. Owing to these reactions, the aromatic substituents change from –OCH₃ to –OH, –CH₃, and –H groups. These reactions may proceed in both gas- and liquid/solid-phase reactions [9].

Concentrations of H-acceptors (radicals) and H-donors play important roles in these pathways. Catechol and methyl radicals formed by the homolysis reaction b are stabilized by the formation of catechol and methane, respectively, where two H-donors can donate hydrogen atoms to these radicals. Otherwise, these radicals are consumed by radical coupling reactions including methylation of aromatic rings by addition of the methyl radical to C-centered radicals that form as resonance structures of phenoxy radicals [92]. Methanol is also produced by coupling of methyl and hydroxyl radicals.

However, the formation of products via the OCH₃ rearrangement pathway requires both H-acceptors and H-donors. The numbers of H-acceptors/H-donors required for the formation of o-quinone methide and o-cresol from guaiacol are 1/1 and 1/3, respectively [93]. Accordingly, the selectivity for the homolysis and rearrangement pathways can be controlled by changing the concentrations of H-acceptors/H-donors in the pyrolysis environment [93, 94]. Asmadi et al. [93] observed that these reactions are promoted in the presence of coking reactions. On the basis of these results it was suggested that polyaromatization during charring reactions produces H-radicals, which act as H-acceptors and H-donors to promote the OCH3 rearrangement pathway. Smaller H-radicals are preferable for these reactions, because all the intermediates involved favor a linear transition state, which maximizes the interaction between the radical orbital and the vacant σ^* orbital of the bond to be cleaved.

Coking and PAH formation

Coking of lignin-derived volatile intermediates occurs in several stages depending on the pyrolysis temperature. First, low-temperature coke formation (first stage coking) is observed along with the OCH₃ rearrangement pathway. It has been suggested that this coking reaction is closely related to the o-quinone methide intermediate (Fig. 9). Only guaiacols with methoxyl groups produced a significant amount of coke in the reactions of various compounds bearing guaiacol-, cresol-, catechol- and phenol-type aromatic nuclei (in a closed ampoule/N₂/ 600 °C/ 80 s, with a final temperature of 569 °C) and 2-ethoxyphenol produced 2,3-benzofuran instead of coke [95]. The 2,3-benzofuran formation can be reasonably explained by the reactivity of the o-quinone methide bearing an allyl moiety, which can be converted to 2,3-benzofuran by cyclization of the allyl radical intermediate formed by H-abstraction.

When catechol-type aromatic rings decompose into gas and other products at high temperatures (>550 °C), catechols/pyrogallols and cresols/xylenols tend to produce coke (second stage coking) [96]. The same o-quinone methide type intermediates can be considered for coking from o-cresols/xylenols, which can form by abstraction of phenolic and benzylic hydrogen atoms. The coke yield increases as the number of methyl groups in the molecules increase: o-cresol (6.1 wt%) <2,4-xylenol (12.8 wt%),



2,6-xylenol (9.2 wt%) <2,4,6-trimethylphenol (23.5 wt%) (in a closed ampoule/ $N_2/600\,^{\circ}$ C) [96]. Therefore, methoxyl and methyl groups are important structural elements for coke formation. Coking from catechols/pyrogallols may arise from reactive fragments, such as acetylene and propagyl radical, as will be described later.

Regarding the reactivities of G- and S-type lignins, the coking reactivity is greater for syringol than for guaiacol, as model lignin primary pyrolyzates. The additional OCH₃ group in syringol, effectively doubles the opportunity for coke formation [92]. Alternatively, the yields of GC/MS-detectable monomers from syringol are lower than those from guaiacol.

PAHs start to form at 600 °C and their production is accelerated at high temperatures > 700 °C [97–99]. Wornat et al. [100] reported 59 PAHs from pyrolysis of catechol at 1000 °C ranging in size up to eight aromatic rings (residence time 0.4 s). In these studies it was also pointed out that the PAH compositions of catechol pyrolysis resembled those of coal pyrolysis. PAH and soot formation during the combustion of hydrocarbon fuels is well understood, and acetylene, propagyl and cyclopentadienyl radicals are considered to be key precursors for the formation of the first aromatic ring [101, 102]. For example, addition of gaseous acetylene to radical sites followed by cyclization, combination of propagyl radicals, and coupling reactions of cyclopentadienyl radicals have been proposed to account for the construction of benzene rings. Ledesma et al. [103] reported propyne, propanediene and cyclopentadiene as stable forms of propagyl and cyclopentadienyl radicals along with acetylene (in maximum yields of ~1.6, 0.5, ~7, and ~20%, respectively) from the gas phase pyrolysis of catechol in the temperature range 600–1000 °C (residence time 0.4 s). These compounds are key intermediates for ring-growth reactions that produce PAHs. Ledesma et al. [104] measured E_a values for the formation of PAHs from catechol in the range of 50–110 kcal mol⁻¹. The values increased for structures with different numbers of rings in the order, 2-rings < 3-rings < 4-rings < 5-rings < 6-rings.

Elimination of aromatic substituents by replacement with hydrogen

Hydrogen-transfer reactions in aromatic rings during coal liquefaction in H-donor solvents, such as tetralin are reported to cleave the strong bonds between aromatic rings and aliphatic side-chains [105, 106]. Aromatic methyl substituents in lignin pyrolyzates, which are formed by OCH₃ rearrangements, methylation by methyl radicals, and homolysis of C–C bonds in ethyl and propyl side-chains, are also known to cleave in the secondary pyrolysis reaction stage [96, 107, 108]. For example, demethylation was reported to be the main reaction of

o-cresol and 2,6-xylenol at 600 °C (in an ampoule, 600 s), which produced 18.0 wt% and 18.2 wt% of the demethylation products, respectively, along with coke [96].

Demethylation reactions occur more selectively at the ortho and para positions to the phenolic hydroxyl group, as can be seen for pyrolysis of 2,3-xylenol, which gives *m*-cresol (32.6 wt%) rather than o-cresol (1.4 wt%) [96]. Demethylation is believed to proceed by replacement with hydrogen radicals (Fig. 10A). Thus, the higher reactivities of the o- and p-methyl groups should be explained by this mechanism. A radical coupling mechanism (Fig. 10b) [96] can explain the selectivity of this reaction, although further study is necessary to confirm this proposition. Coupling of phenoxy and hydrogen radicals proceeds only at the o- and p-positions, owing to the resonance structures of the intermediates, and the resulting cyclohexanedienone has a weak C-CH3 bond. The calculated BDE is reduced from 110.5 to 60.3 kcal mol⁻¹. This is a similar reaction type to that observed for low-temperature homolysis of β -ether linkages of quinone methide intermediates. The strong electron-withdrawing ability of the conjugated carbonyl moiety decreases the BDE of the C-CH₃ bond.

Degradation of aromatic rings into gas

When the pyrolysis temperature is further increased to 550–600 °C, decomposition of catechols and pyrogarolls into non-condensable gases (mainly CO) occurs. Ledesma et al. [103] reported that gas-phase pyrolysis of catechol gives CO (approximately 50 wt%) and acetylene (20 wt%) at 800 °C (with residence time 0.4 s). The gas formation mechanism has also been explained as illustrated in Fig. 11. The mechanism is based on reports that suggest the phenoxy radical undergoes unimolecular decomposition into CO and a cyclopentadienyl

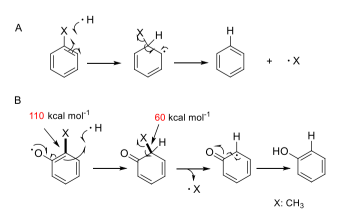


Fig. 10 Replacement of aromatic substituents by hydrogen radicals



radical. The catechol radical decomposes into CO and a cycrolentadienol-lyl radical, which then further decomposes to cyclopentadienone. The cyclopentadienone fragments decompose into CO and two acetylene molecules. Although the exact details of the formation mechanism are unknown, pyrogallols that are formed from the S-type lignins typically produce large amounts of CO₂ along with CO [96].

The gas formation reactivities of cresols/xylenols are lower than those of catechols/pyrogallols and the compositions of the gaseous products are very different for these two types of products [96]. Cresols/xylenols tend to produce $\mathrm{CH_4}$ and $\mathrm{H_2}$ rather than CO and $\mathrm{CO_2}$, along with demethylation products and coke at 600 °C in an ampoule. Methane is formed by H-donation to methyl radicals formed during demethylation and the coking process produces H-radicals during polyaromatization, which leads to the formation of $\mathrm{H_2}$. The $\mathrm{CH_4}$ yield is related to the number of methyl groups in cresols/xylenols, and the yield increases in the order, o-cresol (2.8 wt%)<2,4-xylenol (5.9 wt%) and 2,6-xylenol (6.0 wt%)<1,3,5-trimethylphenol (9.5 wt%). The $\mathrm{H_2}$ yield also increases in this order following the same order as the coke yield.

Lignin pyrolysis in cell walls

Wood cell walls, which have thicknesses of $1-10~\mu m$, are a heterogeneous layered material, in which cellulose microfibrils are filled with a matrix consisting of hemicellulose and lignin [109]. Accordingly, the heterogeneous nature of wood cell wall structures should be considered when assessing the pyrolysis reactions of wood constituent polymers. Lignins in the matrix will undergo pyrolysis alongside the pyrolysis of hemicellulose. The different chemical compositions of lignin and hemicellulose in softwoods and hardwoods may also affect the pyrolysis of these components in wood cell walls.

As discussed above, the majority of ether linkages between phenylpropane units are cleaved homolytically to form radical species, except for α -ether bonds at phenolic

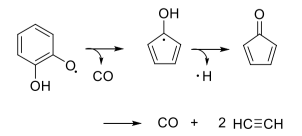


Fig. 11 Mechanism of the reaction of catechol radicals during pyrolysis of catechol



end groups. Conversely, the primary pyrolysis of cellulose and hemicellulose gives non-radical species, because heterolytic cleavage of glycosidic C–O bonds is the main primary pyrolysis reaction [110–112]. These different types of the primary pyrolysis products may interact with each other.

The radical chain reactivity of lignin in wood has been addressed by an *in situ* dimer probe method, which is based on changes in the reactivity of dimers in the presence of wood and its constituent polymers [113]. Strong interactions between lignin and wood polysaccharides have been indicated by this method, which also depend on the type of polysaccharide and the pyrolysis temperature [113]. Xylan activates the radical chain reactions of dimers, whereas glucomannan strongly inhibits non-phenolic dimers. Cellulose inhibits the radical chain reactions of lignins at temperatures below 300 °C, although these effects are minimal at temperatures above 350 °C, when cellulose rapidly decomposes.

Hydrogen donors may play important roles during primary pyrolysis of lignins in wood, because lignin primary pyrolysis occurs by homolysis mechanisms. Wood polysaccharides and their pyrolysis products may act as the H-donors to stabilize lignin-derived radicals, which results in an increase of the monomer yield. This efficiency is reported to be greater for pyrolysis of Japanese beech wood (a hardwood) than that of Japanese cedar wood (a softwood), probably due to the different roles of xylan and glucomannan pyrolysis [87].

Significant interactions have also been suggested in the secondary pyrolysis reaction stage. Levoglucosan, an important primary product from cellulose, efficiently polymerizes into polysaccharides by conversion of vapor into a molten substance under cooling [114, 115]. This polymerization reaction is suppressed in the presence of lignin-derived products [116, 117]. A stabilization mechanism has been proposed; proton donation to levoglucosan through intermolecular hydrogen bonding acts as an acid catalyst to promote the polymerization, which is effectively inhibited by the lignin-derived products through hydrogen bonding between aromatic π -electrons and OH groups of levoglucosan [112].

During the gas-phase secondary pyrolysis reaction stage, the monomer yields from Japanese cedar MWL increased from 5.3 to 20.2 wt% under the influence of cellulose pyrolysis, and the selectivity for catechol over cresols/xylenol products increased from 0.9 to 5.5 [94]. This can be reasonably explained by the action of cellulose-derived products as H-donors that stabilize the catechol radicals and suppress the concentration of guaiacol radicals, which start the OCH₃ rearrangement pathway (Fig. 9).

Conclusions and future prospects

A better understanding of lignin pyrolysis reactions is important for developing improved thermochemical conversion technologies to effectively utilize lignin for the production of biochemicals, biomaterials, and biofuels. This review article summarizes the current understanding of lignin pyrolysis reaction pathways and mechanisms. However, there remains much work still to be done to understand lignin pyrolysis and gasification behaviors. Recently, many theoretical studies have been conducted to estimate the BDEs of linkages in lignin and to help propose reaction mechanisms. However, there is a particular need to accumulate more experimental data to effectively support these calculation results.

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