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Computational evidence for kinetically controlled radical coupling during lignification

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ABSTRACT: Lignin is an alkyl-aromatic biopolymer that, despite its abundance, is underutilized as a renewable feedstock because of its highly complex structure. An approach to overcome this challenge that has gained prominence in recent years leverages the plasticity and malleability of lignin biosynthesis to tune lignin structure *in planta* through genetic approaches. An improved understanding of lignin biosynthesis can thus provide fundamental insights critical for the development of effective tailoring and valorization strategies. Although it is widely accepted that lignin monomers and growing chains are oxidized enzymatically into radicals that then undergo kinetically-controlled coupling *in planta*, direct experimental evidence has been scarce because of the difficulty of exactly replicating *in planta* lignification conditions. Here, we computationally investigate a set of radical reactions representative of lignin biosynthesis. We show that, contrary to the widely held notion that radical coupling reactions should be barrierless and dynamically controlled, the computed activation energies can be qualitatively consistent with key structural observations made empirically for native lignin in a variety of biomass types. We also rationalize the origins of regioselectivity in coupling reactions through structural and activation strain analyses. Our findings lay the groundwork for more detailed multiscale simulations of the lignification process.

Keywords: lignin biosynthesis, lignin structure, density functional theory, polymerization, radical coupling

Introduction

Lignin is an abundant alkyl-aromatic biopolymer constituting up to 30% by weight of biomass¹ that strengthens cell walls, facilitates water transport, and inhibits microbial attack.² It is composed mainly of phenylpropanoid building blocks (i.e., monolignols) that give rise to eight known C-O or C-C linkages of widely varying strengths and characteristics.²⁻⁴ Lignin offers tremendous potential value as a renewable resource⁵ for bulk chemical⁶⁻⁸, specialty chemical⁹, and functional polymer¹⁰ production, particularly if it can be efficiently depolymerized into monomeric units rather than lower-value oligomeric fractions or pyrolysis oil.¹¹ This is a formidable challenge primarily because of the high degree of structural complexity²⁻⁴ that limits achievable monomer yields of selective depolymerization strategies¹²⁻¹³, notably those targeting the most abundant and labile β -O-4 (alkyl aryl ether) linkages.¹⁴ Based on statistical arguments alone, monomer yields should increase with the proportion of β -O-4 linkages¹⁵⁻¹⁶, which should in turn be related to the method of lignin extraction⁵ and the relative proportions of monomeric units with different propensities to form β -O-4 bonds.⁴ However, recent experiments comparing monomer yields across various native¹⁷ and extracted¹⁸ lignin substrates have revealed surprising trends suggesting that post-depolymerization monomer yields are heavily influenced by additional, still unknown, factors beyond what is expected from a thermodynamically-controlled polymerization given a particular monolignol distribution in native lignin. Hence, a detailed understanding of lignin structure beyond the capabilities of modern analytical methods is critical for rationalizing these discrepancies and informing the development of more efficient depolymerization processes.

Plants produce lignin by polymerizing a small slate of monolignols that are biosynthesized in the cytoplasm²⁻⁴ and transported to the cell wall through mechanisms that are not yet well understood.¹⁹ In the cell wall, laccase and peroxidase enzymes facilitate the oxidation of the monolignols and the growing oligomeric chains into phenoxyl radicals. It is widely accepted that chain growth occurs by kinetically-controlled radical coupling, followed by aromaticity-restoring tautomerization and/or hydration, as demonstrated by the lack of optical activity in lignin²⁰ and the ease of incorporating nonstandard monolignols into growing chains.²¹ While empirical structural models have been developed²²⁻²⁷ and successfully used to predict product distributions from lignin pyrolysis²⁸, their usefulness is limited by the accuracy of the underlying experimentally measured analytical properties that they are designed to reproduce. Inconsistencies among experimental data sources have necessitated empirical parameter tuning in some of these models, which limits predictive capabilities.²⁷ The development of predictive, first-principles structural models²⁹ has so far been hindered by the complexity of lignin biosynthesis, which involves a complex, spatiotemporally controlled interplay of metabolic pathways, monomer transport, and enzymatic and non-enzymatic polymerization reaction steps.³⁰ Due to challenges in studying lignification *in planta* at the molecular scale, conclusive evidence for kinetically controlled radical coupling has been scarce in the open literature. *In vitro* polymerization experiments³¹⁻³⁶ have shed light on lignin polymerization mechanisms but are inherently limited by the historical difficulty of quantifying complex product mixtures and the inability to fully replicate *in planta* lignification conditions.

In this regard, first-principles calculations can provide valuable, complementary insight. For instance, extensive study of the thermochemical properties of lignin linkages³⁷⁻⁴⁵ has helped to identify the labile β -O-4 linkage as a target for depolymerization processes. However, with few exceptions⁴⁶, computational studies of lignin polymerization have so far been limited to the thermodynamics⁴⁷⁻⁴⁸ of monolignol coupling reactions or the kinetics⁴⁹⁻⁵⁰ of β -O-4 coupling,

neither of which provide a complete picture of the chemical reactions occurring during lignification. We note that during radical coupling, spin delocalization throughout the conjugated carbon framework of each radical gives rise to a multitude of reactive sites⁴⁸ and possible coupling products. As such, the relative proportions and sequences of observed linkages would then be governed by a combination of the relative propensities of radical coupling at each reactive site and the relative concentrations of reactive intermediates⁴. Therefore, adequate first-principles models of lignin structure²⁹ can be developed by quantifying radical coupling and product formation rates. In this work, we take a step towards such a first-principles structural model of the lignification process by computing the activation barriers of a representative set of radical formation and coupling reactions and showing that they generally agree with qualitative experimental observations of lignin structure. Our work thus provides a foundation for more extensive multiscale simulations of the lignification process.

Computational Methods

The computational methods are described fully in the Supporting Information (SI). Briefly, all density functional theory (DFT) calculations were performed using ORCA 4.0.1.⁵¹ Geometries were optimized at the B3LYP⁵²⁻⁵⁴-D3⁵⁵/def2-SV(P)⁵⁶ level of theory, and single-point energies were calculated at the M06-2X⁵⁷/def2-TZVP⁵⁶ level of theory, which has been shown to perform well for the thermochemistry of lignin models.³⁹ Although DFT-calculated reaction energetics can be dependent on the functional used, our qualitative conclusions remain consistent across several functionals and coupled-cluster theory as they are based on relative reaction rates that are usually more invariant to functional choice (SI Tables S1–S4 and *vide infra*). Solvation contributions were treated implicitly with the CPCM model⁵⁸, and the choice of water ($\epsilon = 80.4$) as the solvent was motivated by the aqueous environments of *in planta* and *in vitro* polymerizations. Lignin polymerization is unique among biological reactions in that enzymes are thought not to be involved in the radical coupling and transfer steps that govern lignin structure. As one of the goals of this work was to investigate if such a model is consistent with experimental observations of lignin structure, we have accordingly modeled these steps in the absence of protein interactions. Considering further the lack of significant charge separation along the reaction coordinates, we believe an implicit solvent approach to be adequate for understanding the intrinsic selectivities of these steps as they occur in the developing plant cell wall. As rigorous conformational analyses⁵⁹ of large and flexible oligomers are computationally intractable, we instead identified a single lowest-energy conformer for each reaction by DFT optimizing structures obtained from force field-based searches, as in prior studies.^{39, 48} Future work will focus on explicit consideration of dynamic, solvation, and confinement effects for predicting the structure of growing lignin chains. All calculations were spin-unrestricted, and transition states were located by scanning the singlet and broken-symmetry spin surfaces along the radical coupling reaction coordinates, followed by eigenvector following optimizations from the energy maxima. Although necessary for accurate absolute rate predictions, entropic contributions to activation energies were not included owing to the difficulty of accurate estimation in solution.⁶⁰ We nevertheless note that all coupling reactions will have a positive entropy of activation that further contributes to their barriers in solution.

Results and Discussion

The complex interlinked networks of sinapyl alcohol (S) and coniferyl alcohol (G) monomeric units found in the lignin structure of most biomass types begin from the simple dimerization of two monolignols. Accordingly, peroxidase-catalyzed bulk dehydrogenation polymerization (DHP) of S and G monomers, in which dimerization is the dominant process, has been extensively investigated *in vitro*. As illustrated in Fig. 1, the G monomer has 3 reactive sites (β , 5, and O) that give rise to 5 possible self-coupling products: β - β , β -O-4, β -5, 5-5, and 5-O-4, excluding O-O that yields an extremely unstable peroxide. The S monomer has 2 reactive sites (β and O; the 5 position is occupied by a methoxy group that blocks subsequent re-aromatization) that give rise to 2 possible self-coupling products: β - β and β -O-4. As a first test of the relevance of kinetically controlled radical coupling to lignification, we begin by comparing experimentally observed product distributions of these *in vitro* dimerization experiments to computed relative radical coupling activation energies. We focus on the initial radical coupling step as the direct radical coupling products, known as quinone methides, are unstable intermediates that readily undergo highly exothermic, and thus kinetically irrelevant⁴⁸, hydration and tautomerization steps to restore aromaticity.

We observe differences in activation energies among coupling products that are in general agreement with product distributions from *in vitro* dimerizations. Notably, G-G couplings involving the β position (β - β , β -5, and β -O-4) have comparable barriers (ca. 2-4 kcal/mol) that are much smaller than the corresponding barriers for 5-5 and 5-O-4 couplings (ca. 10 kcal/mol) (Fig. 1, top). Indeed, horseradish peroxidase (HRP)-catalyzed DHP of G monomers in aqueous solution yields a mixture of β - β , β -5, and β -O-4 dimers in comparable amounts.³⁴ The relative barriers of β - β , β -5, and β -O-4 couplings are somewhat sensitive to functional choice, but the 5-5 and 5-O-4 couplings are strongly disfavored across multiple levels of theory (SI Table S1). Further, the ca. 2 kcal/mol difference in activation energies between β - β , β -5, and β -O-4 couplings is within standard computational error, and dynamic effects that are outside the scope of this work may further contribute to the selectivities. Importantly, radical coupling thermodynamics alone do not accurately describe selectivity, greatly overpredicting the favorability of β - β linkages, thus necessitating explicit consideration of individual activation energies over simpler empirical relationships that relate thermodynamics to kinetics.⁶¹⁻⁶² Interestingly, addition of methoxy groups to the G units disfavors β -O-4 coupling but not β - β coupling, exhibiting a larger difference in barrier heights for the S-S coupling reactions (ca. 1.4 kcal/mol for β - β coupling vs. 7.2 kcal/mol for β -O-4 coupling) (Fig. 1, bottom). This is again consistent with experimental observations of primarily the β - β dimer in bulk DHP of S monomers.^{31, 34} While we recognize that the computed energetic differences are small, results obtained with other density functionals and coupled-cluster theory presented in SI Table S1 are also in full agreement with our observations.

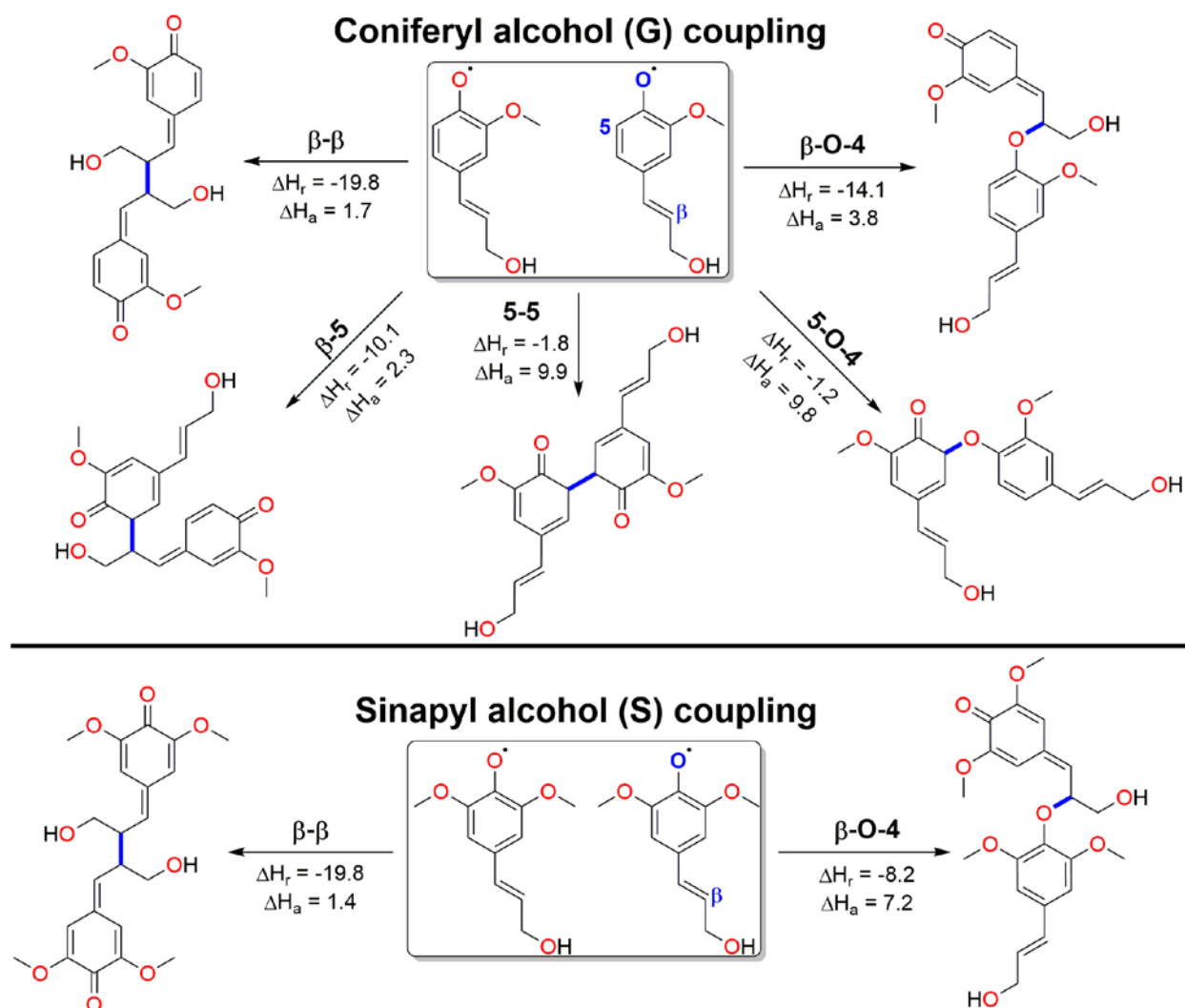


Figure 1. Monolignols and monolignol coupling reactions studied in this work. Each reaction is labeled with its reaction enthalpy (ΔH_r , kcal/mol) and activation enthalpy (ΔH_a , kcal/mol). Reactive positions of monomers and bonds formed in each coupling product are labeled in blue. Results obtained with other density functionals and coupled-cluster theory are presented in SI Table S1.

The notion of kinetically controlled radical coupling appears to contradict the widely held expectation that, excluding the positive free energy contribution from the negative entropy of reaction, all radical coupling reactions should be intrinsically barrierless⁶³ because the interaction of two singly occupied molecular orbitals (SOMOs) should produce a doubly occupied MO that is more stable than either SOMO. Although small radical coupling barriers can sometimes result from disruption of intermolecular forces in the reacting complex (RC)⁶³, this is not the case here as the relative monomer orientations are preserved along the bond-forming reaction coordinates (SI Fig. S1). We thus applied the distortion-interaction activation strain model (ASM)⁶⁴ to understand the regioselectivity accompanying these barriers that appear to be intrinsic to bond formation. ASM decomposes the relative electronic energy, ΔE , at any point along a reaction coordinate into a strain component, ΔE_{str} , corresponding to the energy required to distort the reacting monomeric fragments from their RC geometry to their current geometry, and an

interaction component, ΔE_{int} , corresponding to the stabilization when the fragments interact at this geometry, i.e.,

$$\Delta E = \Delta E_{\text{str}} + \Delta E_{\text{int}}$$

As ΔE_{str} is always positive and ΔE_{int} is always negative, a transition state (TS) will result only if their rates of change balance each other somewhere along the reaction coordinate⁶⁴, and this is indeed the case for all coupling positions (Table 1). Although reactivity differences can sometimes be rationalized solely by differences in the contributions of either component⁶⁵, we find that this is not the case here. For instance, the β - β reaction is most favorable overall despite its relatively large strain energy, and β -O-4 is much more favorable than 5-O-4 despite both reactions having comparable interaction energies. Rather, the observed regioselectivity of monolignol radical coupling appears to be controlled by a delicate balance between strain and interaction. Reactions at the β and 5 positions experience greater strain than reactions at the O position because pyramidalization of the C atoms disrupts π orbital conjugation and orbital overlap with adjacent atoms. At the β position, this increase in ΔE_{str} is compensated by a more favorable interaction energy arising from an increase in C 2p character of the SOMO that enhances spatial orbital overlap during bond formation (SI Fig. S2). The unusually low strain contribution to the β -5 reaction is attributable to a second hydrogen bond that enforces geometric similarity between the RC, TS and product (SI Fig. S1).

Table 1. Strain (ΔE_{str}) and interaction (ΔE_{int}) contributions in kcal/mol to the TS and product energies of radical coupling reactions. Plots of ΔE , ΔE_{int} and ΔE_{str} along the bond forming reaction coordinates are provided in SI Fig. S3.

Monomers	Linkage	Transition state		Product	
		ΔE_{int}	ΔE_{str}	ΔE_{int}	ΔE_{str}
G-G	β - β	-5.9	8.1	-82.7	60.4
G-G	β -5	-3.3	5.9	-65.7	53.6
G-G	5-5	-0.6	10.5	-55.3	51.7
G-G	β -O-4	-0.6	6.2	-53.3	39.0
G-G	5-O-4	-0.3	10.0	-43.4	40.6
S-S	β - β	-9.3	12.4	-85.9	63.9
S-S	β -O-4	-1.3	9.2	-54.3	44.9

Known discrepancies between the structures of synthetic DHP products and native lignin motivate consideration of chain growth reactions (i.e., cross-coupling between monolignol and oligolignol radicals) in addition to monolignol coupling reactions. Notably, there are too few β -O-4 linkages in most DHPs because monolignol dimerization and its dominant β - β / β -5 coupling modes are overrepresented⁴, as evidenced by the strong dependence of cross-coupling selectivity on the rate of monomer addition.³⁵ Furthermore, the strong preference for addition to the O position of the growing chain over the 5 position^{35, 66} suggests an intrinsic difference in regioselectivities of monolignol coupling and chain growth beyond the inability of the oligolignol (growing chain) radical to couple at its β position and explains why β -O-4 is the dominant linkage even in softwood (i.e., high G) lignins.⁴ Thus, to evaluate the ability of kinetically controlled radical coupling to qualitatively predict these reactivity differences, we investigated the coupling of a G radical to a representative G-G β - β resinol dimer (Fig. 2). In accordance with our expectations, we observe a slight preference for β -O-4 coupling over β -5 coupling (ca. 3.8 kcal/mol vs. 4.4 kcal/mol), with 5-

O-4 and 5-5 coupling remaining highly unfavorable. Although well within standard computational error, this preference is nevertheless consistent with the magnitude of the experimentally observed β -O-4 selectivity (e.g., a 1 kcal/mol difference in barriers corresponds to an 85:15 ratio of relative rates at 25°C) while also persisting across multiple levels of theory, albeit with variations in magnitude (SI Table S2).

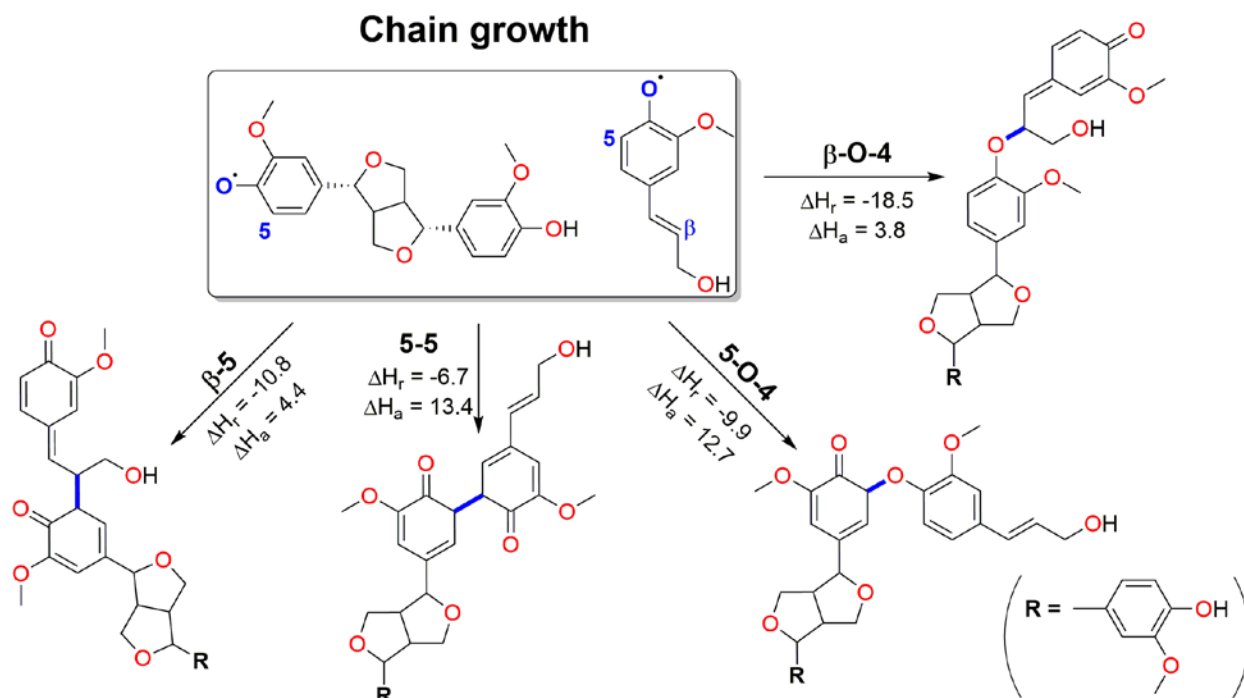


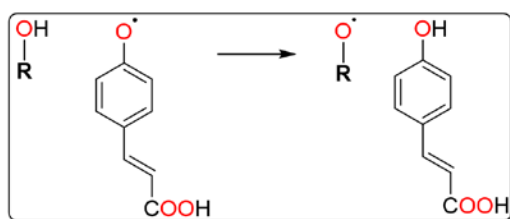
Figure 2. Representative chain growth reactions. Each reaction is labeled with its reaction enthalpy (ΔH_r , kcal/mol) and activation enthalpy (ΔH_a , kcal/mol). Reactive positions of coupling units and bonds formed in each coupling product are labeled in blue. Results obtained with other density functionals and coupled-cluster theory are presented in SI Table S2.

The high propensity for chain growth in native lignin appears at first glance to be inconsistent with our calculations that reveal small differences between chain growth and monomer coupling barrier heights, but is in fact fully consistent with the longstanding hypothesis that the relative rates of monomer coupling and chain growth are governed by differences between the concentrations of monolignol and oligolignol radicals.^{4, 67} Indeed, chain growth and concomitant formation of labile β -O-4 linkages can be favored in DHP experiments by decreasing the rate of monolignol addition^{32, 35}, decreasing the peroxidase concentration⁶⁸, introducing alternative peroxidases with enhanced affinities for growing chains^{67, 69}, or increasing the solubility of growing chains³⁶, all of which promote the formation of oligolignol radicals over monolignol radicals. These observations highlight the importance of determining rates of radical formation, in addition to the rates of radical coupling, in first-principles kinetic models of lignification.

To this end, it was recently shown that, notwithstanding enzyme-binding effects, experimentally observed monolignol and dilignol reactivities toward HRP were correlated to the p orbital density of the phenolic oxygen.⁴⁶ Encouraged by this result, we computed activation energies for radical transfer from a p -coumaric acid radical to representative G and S monolignols and dilignols in order to further shed light on their relative propensities for radical formation (Fig.

3). Besides quantifying intrinsic dehydrogenation reactivities, these reactions also reasonably depict lignification in herbaceous feedstocks in which both growing chains as well as monolignols are oxidized primarily by radical transfer from *p*-coumarates, rather than directly by enzymes.⁷⁰ Our calculations confirm that radical transfers to monolignols are highly facile (Fig. 3 and SI Table S3). Analysis of TS geometries and electronic structures revealed that noncovalent interactions including π stacking favor asynchronous hydrogen atom transfer (HAT) reactions over stepwise proton-coupled electron transfers that would be more characteristic of phenoxyl radicals⁷¹ (SI Fig. S4). More importantly, we also found that radical transfers to growing chains, particularly those with G units, are less efficient than to monolignols, thus necessitating a slow monomer addition rate to achieve reasonable chain lengths in the absence of external factors that specifically promote the oxidation of growing chains. To this effect, oligolignol-specific peroxidases have been identified in hardwoods⁶⁹ and softwoods⁶⁷ that may help ease the requirement for very slow monomer addition, which would ostensibly also lead to very slow lignification rates. In herbaceous lignins, extensive ferulate cross-linking⁷² may also serve to reduce the number of difficult growing chain oxidations required for lignification. We further hypothesize that the recently noted inability of 5-O-4 and 5-5 linkages to undergo chain branching reactions in softwoods⁷³ can similarly be explained by an intrinsic lack of reactivity towards HAT reactions, and work along these lines is ongoing.

Radical transfer



-R	BDE	ΔH_a
	83.7	0.0
	79.8	-0.6
	85.4	5.2
	81.8	1.6

Figure 3. Representative radical transfer reactions. Each substrate is tabulated with its O-H bond dissociation energy (BDE, kcal/mol) and radical transfer activation enthalpy (ΔH_a , kcal/mol). The O-H BDEs do not correlate well with radical transfer kinetics, suggesting that monolignol reactivity is primarily kinetic in origin. Results obtained with other density functionals and coupled-cluster theory are presented in SI Table S3.

The small but significant fractions of 5-O-4 and 5-5 linkages (ca. 1%⁷³ and 4%⁷⁴, respectively) present in native softwood lignin pose a final test for kinetically controlled radical coupling, as their orders of magnitude are inconsistent with the differences in coupling activation energies calculated so far (e.g., $\Delta\Delta H_a$ of 6 kcal/mol corresponds to a <0.1% ratio of relative rates at 25°C). As DHP of model dilignols in the absence of monolignols does indeed yield such linkages³⁵, it has been hypothesized that they result from the coupling of two growing chains, which can each only react at the 5 or O positions.^{2, 4, 73} These couplings would be vanishingly unlikely in DHP experiments where monolignols can freely diffuse in solution together with growing chains, but may be facilitated by spatial confinement or mass-transfer regulation effects under *in planta* lignification conditions. However, the feasibility of this pathway would require much lower intrinsic barriers for the coupling of two growing chains that would be predicted from monolignol coupling or chain growth. Indeed, we observe low barrier heights (ca. 2-3 kcal/mol) for the 5-5 and 5-O-4 coupling of two pinoresinol units that are comparable to barrier heights for monolignol coupling at the β position, suggesting that growing chains can indeed easily couple if they are both oxidized and suitably oriented (Fig. 4 and SI Table S4).

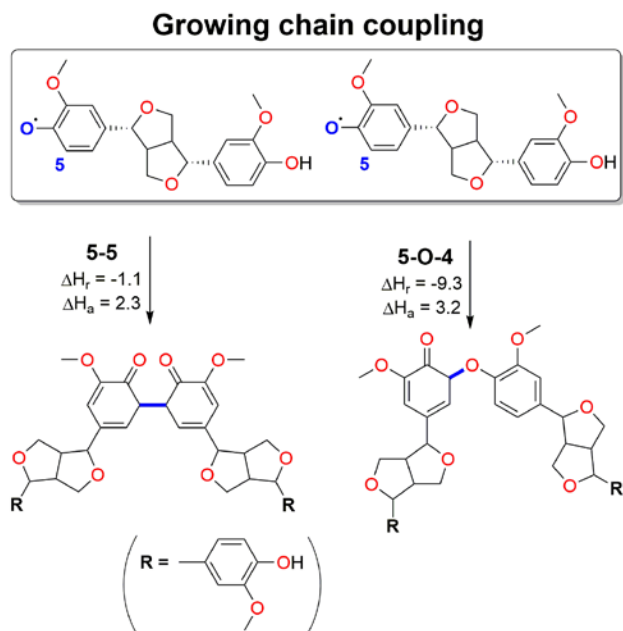


Figure 4. Representative chain coupling reactions. Each reaction is labeled with its reaction enthalpy (ΔH_r , kcal/mol) and activation enthalpy (ΔH_a , kcal/mol). Reactive positions of coupling units and bonds formed in each coupling product are labeled in blue. Results obtained with other density functionals and coupled-cluster theory are presented in SI Table S4.

4. Conclusions

In conclusion, we have computed activation energies for a representative set of radical coupling and transfer reactions and, contrary to expectations that radical couplings should be barrierless and dynamically controlled, demonstrated qualitative agreement with key structural observations of native lignin and synthetic dehydrogenation polymers. Overall, our findings provide computational evidence for the role of kinetically controlled radical coupling, in particular the differences between monolignol coupling, chain growth, and oligolignol coupling rates, in the lignification process. Ongoing work in our groups is focused on extending these findings towards first-principles lignin structural models based on *ab initio* dynamic simulations of the lignification process, and unraveling the effects of monolignol composition and biosynthetic parameters on lignin structure and depolymerization yields.

ASSOCIATED CONTENT

Supporting Information. Full computational details; energies and Cartesian coordinates of optimized structures; results with other density functionals and coupled cluster theory; illustrations of RC, TS and PC geometries; detailed activation strain analyses; MO analyses of radical coupling and transfer TSs. This material is available free of charge via the Internet at <http://pubs.acs.org>.

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Notes

The authors declare no competing financial interest.

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Synopsis: The activation energies of radical reactions representative of lignin biosynthesis are computed and shown to generally agree with experimental structural observations.

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Rationalizing lignin structure with kinetically controlled radical coupling

