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Selective lignin depolymerization *via* transfer hydrogenolysis using Pd/hydrotalcite catalysts: model compounds to whole biomass†

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Cleavage of lignin ether bonds via transfer hydrogenolysis is a promising route to valorize lignin, thus processes that use mild reaction conditions and exploit renewable hydrogen donor solvents (rather than molecular hydrogen) are economically advantageous. Herein we demonstrate the efficient catalytic transfer hydrogenolysis and tandem decarbonylation of lignin model compounds possessing aromatic ether bonds (α -O-4, β -O-4 and 4-O-5 linkages), over transition metal-modified Pd hydrotalcite catalysts with ethanol as the hydrogen donor and solvent. Quantitative conversions and yields were attained for all model compounds, except for 4-O-5 models, which possess inherently strong sp² C-O bonds. The latter demonstrates the utility of Pd hydrotalcite catalysts for transfer hydrogenolysis of model compounds. This process was employed to achieve whole pine biomass delignification with 97% yield and a 22% phenolic monomer yield, with 64% selectivity for 4-(3-hydroxypropyl)-2-methoxyphenol.

Introduction

The need to transition the manufacture of platform chemicals from fossil to non-food competitive, renewable resources is emphasized in Green Chemistry principles,¹ the newly-defined term of "sustainable chemistry",² and is well-aligned with the UN Sustainable Development goals 7, 9 and 11.³ Lignin has received considerable interest as a non-food competitive, renewable resource that can yield high quality aromatics through selective depolymerization.⁴-6 Currently, ~100 million tons per year of lignin are produced as waste from paper and ethanol manufacture, and the vast majority (>98%) is incinerated for energy recovery,² which does not harness its potential as a source of chemical feedstocks. While there is a clear opportunity present, chemical manufacturing from lignin requires scalable, selective, energy-efficient, and clean depolymerization processes which are economically viable.

The asymmetric and non-repeating nature of the lignin biopolymer presents a multitude of ether linkages through which to effect depolymerization, although the diversity of β -O-4, α -O-4, and 4-O-5 ether linkages and associated monomeric units is itself a challenge for selective depolymerization under mild conditions. Such C-O linkages have lower bond

dissociation energies compared to C–C linkages (C–O: 210–290 kJ mol⁻¹ vs. C–C: 450–500 kJ mol⁻¹)⁸ and hence are the chemical functionality targeted in the depolymerization of native lignin. Cleavage of C–C linkages often requires harsh conditions that in turn may result in undesired degradation and aromatic ring hydrogenation.

Most depolymerization strategies rely on hydrogenolysis with exogenous hydrogen derived from fossil sources.9-25 Alternative strategies, such as oxidation, 26-31 hydrolysis, 32-38 hydrothermal,39-42 photo-43,44 and electrochemical45 transformation of lignin have also been explored, with transfer hydrogenolysis (TH) being particularly promising for selective depolymerization under relatively mild conditions without molecular H₂. TH is advantageous for minimizing energy input and avoiding undesired aromatic ring hydrogenation. Although some evidence suggests that hydrogen can be sourced directly from biomass during its valorization,46 the lack of a hydrogen donating solvent requires longer reaction times or is altogether insufficient for complete depolymerization. A homogenous Ru-Xantphos catalyst is reported to facilitate dehydrogenation of benzylic alcohol in β-O-4 compounds, but not γ-alcohols whose presence inhibits hydrogenolysis.47 Use of homogeneous catalysts for large-scale, continuous processing of whole biomass is problematic due to the challenges of product separation, catalyst reuse, waste minimization and expensive ligands, and is associated with poorer energy economy coefficients and environmental energy impact factor.48 The predominant hydrogen sources for TH of lignin model compounds are currently formic acid, formate salts and isopropyl alcohol - all derived from nonrenewable sources. 49,50 Shen and co-workers recently

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demonstrated an ethanol/isopropanol mix as solvent and hydrogen donor (H-donor) at very high temperature (270 °C) for the transfer hydrogenolysis of organosolv poplar lignin over high loading NiCu bimetallic catalysts, 51 an approach that does not address the challenge of lignin fractionation from holocelluose pulp associated with processing whole biomass. In contrast, the use of ethanol as a hydrogen donor for TH is highly desirable due to its availability from fermentation of renewable sugars and starches, as well as its relatively low cost and low toxicity compared to alternative donors. 52,53 Ethanol is reported as a hydrogen donor for the reductive catalytic fractionation (RCF) of birch, poplar, spruce, and pine under forcing conditions (e.g., 210 °C and 15 h).54 Aqueous ethanol was also used a solvent for the RCF of hybrid poplar clones in the presence of 0.36 M HCl (at 175 °C)⁵⁵ over commercial 5 wt% Pd/C catalysts, albeit hemicellulose (not ethanol) was proposed as the internal H donor. Ethanol is not only an excellent external H-donor for TH and activation of cationic metal pre-catalysts, but also has good dissolution and solvolytic properties for lignin, 55,56

providing multiple advantages for use in transfer hydrogenolysis.

Catalytic TH of lignin model compounds has been reported (Table 1 and Fig. S1†); however, tandem catalytic TH and decarbonylation over a single catalyst in a one-pot reaction is, to our knowledge, previously unreported. Optimizing the activity and selectivity of multifunctional catalysts capable of such tandem transformations is challenging, since the reaction steps can be interdependent. High selectivity requires control over the electronic and geometric properties of catalytically active species (and their spatial distribution), and mechanistic insight of underlying structure-activity relationships, including the role of catalyst supports.63 Efforts by our group and others to develop heterogeneous Pd catalysts on active, tunable supports for organic synthesis and biomass valorization have exploited cooperativity between strongly interacting support matrices and metals to regulate metal speciation, reactivity and stability. 64-71 MgO and γ -Al₂O₃ supported metal catalysts exhibit a range of acid-base properties:72-75 for example, Saad et al. reported that

Table 1 Precedents for transfer hydrogenolysis (TH) of β-O-4 lignin model compounds over supported catalysts

Substrate	Conditions	Products (yield/%)		Conversion/%	Whole lignin (monomer yield/%)	Renewable H-donor	Ref.
	Ni/Al₂O₃-600 i-PrOH, 100°C, 3h	OH + +	O OH +	100	Yes (13) ^a	No	57
	Raney Ni i-PrOH, 80°C, 3h	100% 87% +	<1% 12% OH	100	No	No	58
	PdFe/Fe ₂ O ₃ i-PrOH, N ₂ (10 bar), 240°C, 1.5h	+	OH	100	No	No	59
OH	Pd/C EtOAc/H ₂ O, 0.4 eq. H ₂ , 0.05-0.08 eq. NaBH ₄ , 80°C, 3h	0		100	No	No	60
OH	Ni _{40%} -Ru _{5%} /Al ₂ O ₃ iPrOH, 150 °C, 1h	95% OH + + + +	92% OH OH 8% 23%	59	No	No	61
HO HO O HO	Pd-Fe ₂ O ₃ (40 wt. %) HCOONa (400 wt. %) Ethanol, 150 °C, 6h	но 60%	+ OH OH	100	Yes (20)	Yes	62
O HO O	Pd-CuHT EtOH, 150°C, 6h	90%	HO 93%	98	Yes (35) ^a	Yes	This work

^a Number of ether linkages between monolignols, and hence theoretical monomer yield (typically 30–40%), will depend on the wood type. See ESI for details on the quantification of ether linkages and associated theoretical monomer yield.

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Pt/MgO displays strong basicity, whereas Pt/Al₂O₃ displays moderate acidity; ⁷¹ similar observations were made by Groppo et al. for Pd analogues.76 Layered double hydroxides (LDHs) possess tunable acidity and basicity and can support a number of transition metals, yielding supported catalysts that facilitate a multitude of catalytic transformations. LDHs comprise layers of metal hydroxides sandwiched between layers of weakly bound anions (and often water). Hydrotalcites (HTs) are subset LDHs, with the formula $[M_{1-x}^{2+}M_x^{3+}(OH)_2]^{x+}(A^{n-})_{x/n}\cdot mH_2O$, wherein M^{2+} and M^{3+} are commonly Mg^{2+} and Al^{3+} (or interchangeable alkali earth and transition metal cations).77 Although HTs exhibit solid basicity approaching that of MgO,78 their variable cationic charge means that both O2- Lewis base and Al3+ Lewis acid centres are accessible to substrates. 79,80 Catalytic applications of Pd/LDH (or HT) include aldehyde decarbonylation,66 aldol condensation64 and acceptorless alcohol and amine dehydrogenation, 65,81 ole-

Here we demonstrate the feasibility of the one-pot, base-free cleavage of α -O-4, β -O-4 and 4-0-5 linkers of lignin with ethanol as H-donor over multifunctional Pd catalysts prepared by continuous flow synthesis (Fig. S2†), and featuring supports possessing different acid–base character; 83,84 MgO, of primarily basic character; HT, exhibiting Lewis acidity and basicity; and γ -Al $_2$ O $_3$, of primarily Lewis-acidic character. The same catalyst promotes TH, hydrogenolysis and decarbonylation. Novel aspects include: (i) efficient transfer hydrogenolysis of α -O-4, β -O-4 and 4-O-5 model compounds using ethanol as renewable hydrogen source; (ii) demonstration of a multifunctional catalyst for the tandem transfer hydrogenolysis and decarbonylation of β -O-4 motifs; and (iii) whole biomass valorization to phenolic derivatives using only ethanol as a hydrogen donor.

fination of carbonyls via aldol-decarbonylative coupling64 and

deoxygenative olefination of alcohols,82 among others.

Results and discussion

Catalyst characterization

Pd/γ-Al₂O₃, Pd-MgO and pure and metal-modified Pd/HT catalysts were synthesized and characterized to: (i) assess their catalytic activity for the TH of lignin ether linkages to phenolic platform chemicals; and (ii) establish structure-reactivity relationships that identify physicochemical properties desirable for active and selective transformations. A Pd loading between 2-5 wt% was targeted for Pd/γ-Al₂O₃ and Pd-MgO, with that of (metal-modified) Pd/HT \sim 5 wt%. Both oxide supported Pd catalysts were synthesized by wet impregnation of the supports with Pd(NO₃)₂·xH₂O. MgO was prepared by a previously reported batch co-precipitation method.85 Pd/HT and transition metal-modified analogues (Table 3, A-G) were synthesized by a mesoscale, continuous flow precipitation method, adapted from our previous report.86 Compositions of A-G were highly reproducible (standard deviations <5% of molar metal content), with ICP-OES elemental analysis showing actual metal compositions was within 5% of targeted values (Table S1†). A decrease in the M²⁺: M³⁺ ratio from the ideal value of 3.0 was observed for a few of the metal-modified Pd/HT samples, which is likely a consequence of the larger ionic radius

of +2 dopant cations compared to that of Mg²⁺, or +3 dopant cations compared to the ionic radius of Al³⁺.

All Pd/HT samples exhibited a single crystalline phase from powder X-ray diffraction (XRD), associated with the hydrotalcite phase, characterized by (003), (006), (009), (015), (018), (110) and (113) reflections (Fig. S3†). No Pd or PdO crystalline phases (Pd: $2\theta = 40.5^{\circ}$ (111), 46.7° (200), 67.5° (220); PdO: 33.7° (101), 41.8° (110), 54.6° (112), 60.0° (103), 60.6° (200)), were observed in Pd/ g-Al₂O₃ and the Pd/HT samples, even at 5 mol% Pd, suggesting that palladium is present either as low-nuclearity species of <2 nm particles, which are either dispersed over the HT support or incorporated as dopants within the HT cation layers. In contrast, Pd/MgO (Fig. S4†) exhibits weak reflections attributable to PdO. The (003), (006), (009) reflections of the HT are sensitive to the interlayer spacing (crystallographic parameter c), while the (110) and (113) reflections are related to the average metal-metal distance within the cation layers (crystallographic parameter a). The consistent values of a and c parameters for all doped HTs suggest that a common, crystalline HT structure was obtained in each case. Given that lattice expansion, reflected in parameter c (ESI Table S1†), does not vary significantly (range: 23.23-23.98 nm), there is no direct evidence that Pd was incorporated into the support structure to a significant extent. HT crystallite sizes, determined by applying the Scherrer equation, were insensitive to presence of Pd or other transition metal additives (10.8 \pm 0.8 nm).

Fourier transform infrared (FTIR) spectra were consistent with a hydrotalcite phase, exhibiting characteristic bands for carbonate anions (1350-1370 cm⁻¹) and interlayer water (\sim 1600–1700 cm⁻¹) (Fig. S5†). Surface areas and pore size distributions for A-G were determined by N2 porosimetry (Fig. S6 and S7†) and subsequent application of the Brunauer-Emmett-Teller (BET) and Barrett, Joyner and Halenda (BJH) models (Fig. S8 and S9†), respectively (Table S2†). Surprisingly, the bimetal modified HTs all possessed higher surface areas that the parent (148 m^2 g^{-1}) or single metal modified (53–126 m² g⁻¹) HTs. We infer that the introduction of a second transition metal during the flow synthesis promotes the formation of highly dispersed Pd,M-containing clusters/nanoparticles with surface areas of 152–385 m 2 g $^{-1}$. We previously reported that pore volume decreases with increasing Pd loading,65 which is consistent with blockage of micropore entrances to interlayer spaces at higher loadings. However, the effects of transition metal doping on surface area and pore size are not fully elucidated. Thermogravimetric analysis (TGA) of all Pd/HTs revealed decomposition profiles consistent with a MgAl-HT.87 Three endothermic transitions were observed (Fig. S10†): at ~100 °C due to the loss of physisorbed water; between 200-250 °C due to loss of interlayer water; and between 375-400 °C due to dehydroxylation and the loss of interlayer carbonate anions.77 These transitions occur within a relatively narrow temperature range for A-G, although a shift to lower temperatures is observed for metal-modified Pd/HTs relative to Pd-free HTs. This shift was most prominent for the high-temperature dehydroxylation/ decarbonation, indicating that Pd likely destabilizes cation substitution and causes corresponding structural disorder.88

Analysis of the surface palladium by X-ray photoelectron spectroscopy (XPS) indicated the presence of multiple chemical environments in all samples. High-resolution Pd 3d XP spectra were dominated by a doublet with a $3d_{5/2}$ binding energy (BE) of 336.7 eV and spin-orbit splitting of 5.3 eV, assigned to Pd²⁺ in PdO (Fig. 2). A weaker doublet with a 3d_{5/2} BE of 335.4 eV attributable to metallic Pd was also apparent in all samples, with an additional, very weak doublet at 338.5 eV BE which we have previously assigned to Pd4+ substituted at low concentrations into the cation layers. Surprisingly, the diverse supports and metal modifiers exert little influence on the Pd surface speciation, which is dominated by PdO (70-80% of palladium present), alongside a modest amount of Pd metal (10-20%) and trace Pd⁴⁺ (<10%, likely in an octahedral coordination bound to 6 oxygen anions within the hydrotalcite layers). This observation was confirmed by complementary Pd K-edge X-ray absorption near edge spectroscopy (XANES), for which leastsquares fitting identified Pd2+ in oxide clusters/nanoparticles as the dominant palladium phase (Fig. S11†). Mg 1s spectra of Pd/MgO, Pd/HT, and PdZn/HT shows that the presence of small amounts of surface magnesium carbonate species, which suggest presence of strongly basic sites (Fig. S12†). A highenergy XPS Mg 1s peak corresponding to surface carbonate was observed for Pd/MgO and PdZn/HT, suggesting presence of strong basic sites. In contrast, XPS Mg 1s of PdFe/HT and PdCu/ HT were dominated by a peak characteristic of the pure HT, suggesting much lower basicity.

The morphology of samples **B–G**, as well as Pd/Al_2O_3 and Pd/MgO, were imaged by transmission electron microscopy (TEM). Uniformly dispersed metal-containing nanoparticles of ~ 1 –3 nm were visible for all materials (Fig. 3), consistent with the absence of crystalline phases (other than those of the supports) by XRD. Our previous studies of Pd/HT catalysts with nominal Pd loadings spanning 0.1 to 5 wt% revealed the presence of PdO and Pd metal clusters and nanoparticles for loadings >1 wt%, ⁶⁶ in accordance with Fig. 3.

These characterizations are consistent with a structure of the metal-modified 5 wt% PdHTs (B-G) that consists of atomically dispersed Pd4+ species incorporated into the HT cationic layers, low nuclearity Pd²⁺ species incorporated into the cationic layer or dispersed over the surface of HT layers, and Pd⁰ agglomerates on the surface. Unfortunately, we were not able to collect EDX mapping images of sufficient resolution to support this hypothesis. Furthermore, determination of Pd dispersion through common experimental means such as CO chemisorption could not be employed as this approach requires that CO titrates only Pd metal and not the support. However, weakly acidic CO molecules can bind to hydrotalcites as they are solid bases, thus hindering accurate dispersion measurements. Consequently, catalytic activity is quantitated based on the total Pd content, consistent with previous reports⁸⁹⁻⁹¹ and hence reported TONs could underestimate the true catalyst performance.

Catalytic transfer hydrogenolysis of model compounds

Catalyst activity for cleavage of different lignin motifs was subsequently explored for the synthetic model substrates in Fig. 1 (possessing α -O-4 (L-1), β -O-4 (L-2 to L-4) and 4-O-5 (L-5) linkages) and analogs lacking methoxy functions. Note that although we were unable to directly detect free H₂ by GC-MS for any reactions below, hydrogen pop tests qualitatively evidenced *in situ* hydrogen formation from ethanol (as discussed later).

a-O-4; benzyl phenylether (L-1). α -O-4 model compounds such as benzyl phenylether are the simplest phenolic ethers, and consequently their reactivity is dominated by transfer hydrogenolysis (*versus* competing organic transformations) enabling rapid screening of potential metal/HT catalysts. L-1 was selected as the simplest model substrate for determining activity for TH of α -O-4 linkages with isopropanol, ethanol and glycerol as hydrogen donor solvents. 5 wt% Pd/HT afforded \sim 80% yield of hydrogenolysis products (equimolar **a** and **b**)

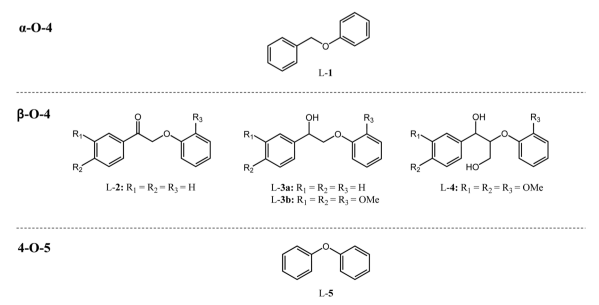


Fig. 1 Model compounds L-1–L-5 with α -O-4, β -O-4 and 4-O-5 motifs.

Table 2 Effect of solvent for the transfer hydrogenolysis of benzyl phenylether (L-1) (α-O-4 linkage)⁶

			Yield/%		
Entry	Solvent	Conversion/%	a	b	
1	Isopropanol	80	80	78	
2	Ethanol	62	62	61	
3	Glycerol: water (1:1)	12	10	10	
4	Isopropanol: water (1:1)	53	53	50	
5	Isopropanol: glycerol (1:1)	50	50	50	

^a Reaction conditions: 0.1 mmol benzyl phenyl ether (L-1), 3.0 mL solvent, 5 mol% Pd/HT catalyst (B), ~50 W microwave heating at 150 °C, 2 h.

with isopropanol in 2 h at 150 °C under microwave heating (Table 2, entry 1), but only 60% with ethanol (Table 2, entry 2); this is consistent with the more favorable kinetics and thermodynamics expected for the dehydrogenation of a secondary (isopropanol) versus primary (ethanol) alcohol. Neat glycerol was too viscous to use as a solvent and was therefore diluted 1:1 with deionized water, but nevertheless afforded low yields (Table 2, entry 3). To probe whether dilution with water or the low viscosity of glycerol contributed to this low yield, additional reactions were performed with 1:1 isopropanol/water (Table 2, entry 4), which lowered the yields of **a** and **b** by \sim 30% relative to the reaction with pure isopropanol. Similar modest yields were observed for reaction with a 1:1 isopropanol:glycerol solvent mixture (Table 2, entry 5), confirming that glycerol is not an effective hydrogen donor under these conditions. Note that the ability of glycerol to act as a hydrogen donor is strongly basedependent due to the need to convert the initial dehydrogenation product, dihydroxyacetone, into the more thermodynamically stable lactate product.92-94 Consequently, addition of base could improve the viability of glycerol as a hydrogen donor. Although isopropanol delivered superior yields of hydrogenolysis products compared to ethanol, the latter offers a renewable source of hydrogen donor from fermentation of cellulosic waste, and hence was selected for further studies.

The family of Pd catalysts were subsequently screened for TH of benzyl phenyl ether (L-1) with ethanol as the hydrogen-donor solvent under microwave heating at 150 °C for 2 h, with a Pd loading of 5 mol% relative to the substrate (Table 3). Transition metal-free HT (A, entry 1), 3 wt% Pd-Al₂O₃ (entry 2) and a homogeneous Pd(OAc)₂ (entry 3) were inert under these conditions, while commercial 3 wt% Pd/MgO and 3 wt% Pd/C catalysts only achieved ~30% conversion to toluene and phenol (Table 3, entries 4–5). In contrast, Pd-containing HT catalysts afforded high conversions (81–100%) with Pd-HT, Pd-FeHT, Pd-CuHT and Pd-CoHT (entries 6,7, 9,10) promoting

quantitative conversion with full selectivity for toluene and phenol. To further differentiate the activity of the latter four, reaction time was decreased to 10 min, which allowed us to distinguish the following activity order: PdCu/HT (F) > PdFe/HT (C) > PdCo/HT(E) > Pd/HT(B) > PdZn/HT(G) > PdNi/HT(D). We note that PdFe/HT and PdCu/HT afforded >90% conversion in 10 min. We postulated that the superior activity of the Pd/HT catalysts relative to Pd/Al2O3, Pd/C and Pd/MgO is associated with their unique activity for alcohol dehydrogenation of ethanol. This was substantiated with substrate-free reactions in ethanol, where Pd/HT catalysts uniquely yielded dehydrogenation products (ethyl acetate and acetaldehyde). In conjunction with the similar Pd speciation observed for all catalysts (Fig. 2), these observations suggest that TH activity is primarily a function of the support acid-base properties, rather than electronic and geometric properties of the supported metals. The

Table 3 Catalytic activity for transfer hydrogenolysis of benzyl phenylether (L-1) $(\alpha$ -O-4 linkage)^a

Entry	Catalyst	Conversion/%		
1	HT (A)	0		
2	3 wt% Pd/Al ₂ O ₃	0		
3	$Pd(OAc)_2$	0		
4	3 wt% Pd/C	30		
5	3 wt% Pd/MgO	34		
6	5 wt% Pd/HT (B)*	100 (47)		
7	5 wt% PdFe/HT (C)*	100 (91)		
8	5 wt% PdNi/HT (D)*	81 (37)		
9	5 wt% PdCo/HT (E)*	100 (74)		
10	5 wt% PdCu/HT (F)*	100 (93)		
11	5 wt% PdZn/HT (G)*	88 (41)		

^a Reaction conditions: 0.1 mmol benzyl phenyl ether (**L-1**), 3.0 mL ethanol, 5 mol% catalyst, 50 W microwave heating at 150 $^{\circ}$ C, 2 h (or *10 min with resulting conversion in parentheses).

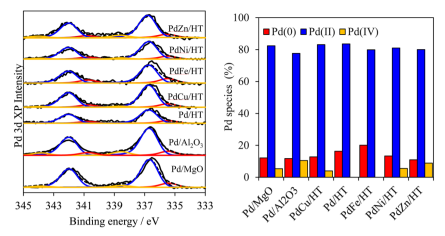


Fig. 2 Pd 3d XPs spectra of Pd/HTs (B-G), Pd-MgO, and Pd-Al₂O₃.

differences in activity between Pd/HTs modified with different transition metals can only be accounted for with EDX data that shows the metals are phase separated from Pd, alloyed with Pd, or in the HT cationic layers. Note that while $\alpha\text{-}O\text{-}4$ linkages are not common in lignin, the preceding results demonstrate the broader potential application of mono- and bimetallic Pd/HTs for organic synthesis, and insight into structure–reactivity relationships pertinent to the transfer hydrogenolysis of 2-phenoxy-1-phenethanol (L-3a) described below, which involves ether cleavage and subsequent benzyl alcohol dehydrogenation.

 β -O-4; 2-phenoxyacetophenone (L-2). The β -O-4 linkage is the most abundant in lignin and can undergo additional transformations prior to ether cleavage, including dehydrogenation or dehydration of the benzylic alcohol (see scheme in Table 4). Dehydrogenation weakens the ether bond,⁴⁴ and is presumed to occur first. To explore these competing processes, we first isolated the ether cleavage step by implementing 2-phenoxyacetophenone (L-2), the intermediate formed after dehydrogenation of the benzylic alcohol (Table S3†) and colloquially known as a 'C2-type' model compound due to the lack of

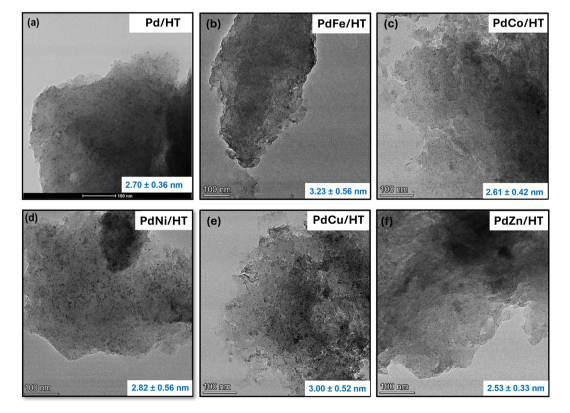


Fig. 3 TEM images of metal modified PdM/HT catalysts (a) Pd/HT, (b) PdFe/HT, (c) PdCo/HT, (d) PdNi/HT, (e) PdCu/HT, and (f) PdZn/HT. Corresponding particle size distribution (bottom right corner of images a–f) histograms are shown in Fig. S13.†

Table 4 Catalytic performance for transfer hydrogenolysis of 2-phenoxy-1-phenethanol (L-3a), and proposed two-step reaction pathway to phenol (b) and acetophenone (c) via 2-phenoxyacetophenone (L-2).^a

		Conversion (%)	Yield $(\%)^b$				
Entry	Catalyst		c	b	d	L-2	
1	HT (A)	0	0	0	0	1	
2	5% Pd/HT (B)	68	66	66	0	2	
3	5% PdFe/HT (C)	97	79	86	4	11	
4	5% PdNi/HT (D)	6	5	5	0	1	
5	5% PdCo/HT (E)	59	51	52	0	7	
6	5% PdCu/HT (F)	98	93	96	0	2	
7	5% PdZn/HT (G)	16	8	11	0	5	

^a Reaction conditions: 0.1 mmol 2-phenoxy-1-phenethanol, 3.0 mL ethanol, 5 mol% catalyst, 50 W microwave heating at 150 °C, 2 h. Note that hydrogen may derive from ethanol dehydrogenation or intramolecular dehydrogenation of L-3a. ^b Unaccounted for yield is a result of aldol condensation between acetaldehyde and acetophenone forming ethyl benzoate.

γ-functionality.95 Catalysts A-G were evaluated against 2-phenoxyacetophenone (L-2) under the preceding conditions employed for benzyl phenyl ether (L-1) (Table S3†). Similar reactivity trends were observed as for L-1: the parent HT was inactive, while PdFe/HT (C) and PdCu/HT (F) afforded highest activity (99% conversion and >95% selectivity). Despite the additional reaction pathways possible for this substrate, transfer hydrogenation of the ketone to the alcohol (expected to occur after ether cleavage) and/or aldol condensation of L-2 or acetophenone with itself (or in situ generated acetaldehyde), only a small decrease in selectivity to TH products was observed for L-2 relative to L-1. However, higher temperatures (170 °C) increase the propensity towards these additional reaction pathways and by-products, primarily the aldol product between acetophenone and acetaldehyde due to the large excess of acetaldehyde formed from the ethanol solvent, rather than selfcondensation of acetophenone. Hydrogenation of acetophenone to the secondary alcohol was a minor pathway. At 150 °C, these undesired products are identifiable but in negligible amounts for catalysts C, E, F, \sim 5% for B, and \sim 10-15% for D and G. The strong basicity of Pd/MgO and PdZn/HT, established based on the presence of a high-energy XPS Mg 1s peak corresponding to surface carbonate (Fig. S12†), accounts for the higher activity for undesired aldol reactions. In contrast, the

most active and selective Pd/HT, PdFe/HT and PdCu/HT (not shown) are dominated by a peak characteristic of the pure HT, and hence offer moderate base/acid character – a Goldilocks zone.

2-phenoxy-1-phenethanol (L-3a). 2-Phenoxy-1β-O-4; phenethanol (L-3a) is structurally akin to the native β-O-4 motif in lignin (Table 4) and hence was also studied. The benzylic hydroxyl group in L-3a introduces thermodynamic and kinetic complexity. Computational studies suggest that the ether bond dissociation energy (BDE) decreases by ~55 kJ mol⁻¹ after dehydrogenation to the ketone (L-2) (from 289 kJ mol⁻¹ to 234 kJ mol⁻¹),44 and hence that catalysts active for dehydrogenation should facilitate ether transfer hydrogenolysis.44 Note the benzylic hydroxy group is also a potential source of hydrogen equivalents, which could preclude the need for an external H-donor solvent. Such intramolecular hydrogen transfer reactivity is explored for substrate L-4. Considering the multistep transformation of L-3a, the parent HT (A) is inert for dehydrogenation of the benzylic alcohol and hence ether activation. Trends in the activity for ether TH are consistent with those for L-1 and L-2, with PdFe/HT (C) and PdCu/HT (F) being most active and selective to phenol (b) and acetophenone (c), albeit with slightly lower yields, and PdNi/HT (D) and PdZn/HT (G) the least active. For L-3a, the benzylic alcohol must first

undergo dehydrogenation to the carbonyl ketone. Previously we demonstrated that all Pd/HTs are capable of ethanol dehydrogenation, and are therefore expected to be similarly active towards the secondary benzylic alcohol in L-3a. However, the vast excess of ethanol (500 molar equivalents) relative to L-3a is expected to kinetically favor dehydrogenation of the ethanol solvent versus the substrate, likely accounting for the lower TH activity and yields obtained for L-3a (Table 4) versus L-2 (Table S3†). The lower selectivity of PdFe/HT $\it versus$ PdCu/HT to $\it b$ and $\it c$ cannot be ascribed to a deficiency in the former for the initial dehydrogenation step, as the Fe-modified Pd/HT produces significant L-2 intermediate and is thus attributed to intrinsically weaker activity for TH of the β-O-4 linkage. Timed reaction courses suggest that ether cleavage follows dehydrogenation of the benzylic hydroxy group, and is much faster than initial dehydrogenation, based on the low steady state concentration of L-2, and the detection of acetophenone (c) before benzyl alcohol (d).

The overall catalytic performance of transition metal modified Pd/HTs is summarized in Fig. 4, with PdCu/HT the most active for cleavage of all three linkages.

β-O-4; methoxy functionalized (L-3b). Further experiments were undertaken on model substrates more representative of lignin, such as 1-(3,4-dimethoxyphenyl)-2-(2methoxyphenoxy)ethanol (L-3b), to explore the effect of methoxy substitution (Fig. 5a). The most active 5 wt% PdCu/HT (F) catalyst exhibited highest activity for both dehydrogenation and TH (Table 4). Introduction of m,p-methoxy groups did not decrease selectivity compared to reactions with L-3a, but did decrease conversion to 60%. Computational modelling indicates that methoxy groups actually increases the thermodynamic driver for ether cleavage,96 which suggests the lower activity likely reflects slower kinetics. Varying the catalyst loading (Fig. 5b) had little impact on L-3b conversion, suggesting that these slower reaction rates arise from intrinsic limitations to ether bond activation (e.g. steric factors hindering

the correct adsorption configuration) rather than mass-transport limitations. High selectivity for products **e** and **f** (Fig. 5, ~90%) were obtained for 2.5 mol% PdCu/HT, and hence this catalyst loading was chosen to subsequently explore the impact of reaction temperature (Fig. 5c). Conversion of **L-3b** increased from 62% at 130 °C to 82% at 160 °C, although selectivity was highest at 150 °C, decreasing at higher temperature due to the hydrogenation of **e** and formation of aldol byproducts. As expected, **L-3b** conversion increased with reaction time (Fig. 5d), reaching 72% after 2 h at 150 °C, with (>90% selectivity for **e** and **f**). Near-quantitative conversion was obtained in only 6 h using 2.5 mol% PdCu/HT at 150 °C (yields of **e** and **f** of 93% and 95% respectively).

β-O-4; C3 type (L-4). Model compound L-4 was employed as a 'C3'-type compound, featuring a γ -hydroxy group, ⁹⁵ common in native lignin, which introduces additional potential dehydrogenation and decarbonylation pathways (Scheme 1). The PdCu/HT catalyst achieved near-quantitative conversion of L-4 (98%), and >90% selectivity to 3,4-dimethoxyacetophenone (e) and guaiacol (f) (Table 5, entry 1). The ability to directly transform L-4 into these desirable platform chemicals in a one-pot reaction is novel, previously necessitating multiple catalysts in multiple separate reactions. ^{44,97}

We hypothesize that the γ -hydroxy group undergoes dehydrogenation to an aldehyde, which is then eliminated via decarbonylation either before or after ether cleavage (dehydrogenation of α - and γ -hydroxy groups prior to ether cleavage is thermodynamically favorable). We have previously analyzed headspace gases of aldehyde decarbonylation reactions under similar conditions by GC-MS. Under air, such reactions evidenced CO₂ formation, likely from the oxidation of CO with residual oxygen. Control reactions without aldehyde substrates afford no CO₂, suggesting the source of CO₂ is decarbonylation and oxidation, and exclude the possibility that aldehydes are first oxidized to carboxylic acids and undergo subsequent

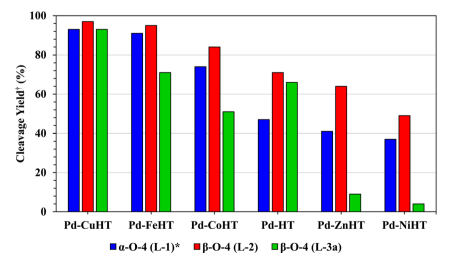
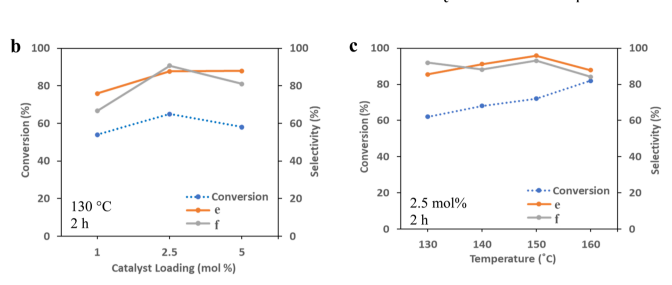


Fig. 4 Comparative yield of transfer hydrogenolysis products from L-1, L-2 and L-3a catalyzed by Pd/HTs. Reaction conditions: 0.1 mmol substrate, 3.0 mL ethanol, 5 mol% catalyst, 50 W microwave heating at 150 °C, 2 h (or *10 min for L-1). This data is a consolidation of Tables 3, S3†, and 4. †For L-1, cleavage yield includes toluene (a) and phenol (b); for L-2 and L-3a, cleavage yield includes acetophenone (c) and phenol (b).



^aHydrogen may arise from ethanol dehydrogenation or dehydrogenation of the L-3 substrate.

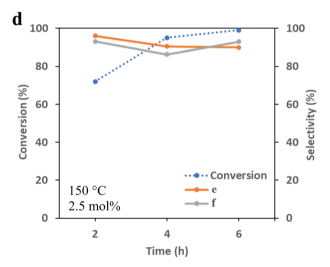


Fig. 5 (a) Reaction scheme for transfer hydrogenolysis of L-3b; and influence of (b) catalyst loading, (c) reaction temperature and (d) time on conversion and product selectivity. Reaction conditions: 0.1 mmol L-3b, 3.0 mL ethanol, PdCu/HT, 50 W microwave heating. ^aHydrogen may arise from ethanol dehydrogenation or dehydrogenation of the L-3 substrate.

decarboxylation (by showing carboxylic acids are inert under our catalytic conditions).

Dehydrogenation of the benzylic alcohol is likely more thermodynamically favorable than that of the primary γ -hydroxy. The product distribution suggests that both hydroxy groups are dehydrogenated prior to decarbonylation and ether cleavage, as product i was observed as an intermediate, while j and k were not. Note that i is the dehydrogenated product of 1-(3,4-dimethoxyphenyl)-2-(2-methoxyphenoxy)-1,3-propanediol

(L-3b), and thus likely follows the previously established pathways for its transfer hydrogenolysis (Table 4, starting from L-2). Dehydrogenation of benzylic alcohol produced from L-3a could also be followed by a retro-aldol cleavage^{98,99} to release formal-dehyde and L-2, which in turn could undergo ether cleavage to form acetophenone; although the formaldehyde by-product was not observed with only decarbonylation followed by ether cleavage observed. In comparison to the benzylic alcohol, dehydrogenation of the primary γ -hydroxy is less kinetically and

Scheme 1 Proposed reaction pathway for the multistep transformation of L-4

thermodynamically favored, but could likewise be followed by a retro-aldol cleavage (although we did not observed any products expected from such pathway either). The absence of a dehydrogenated γ -alcohol intermediate may thus reflect two possibilities: (i) once dehydrogenated, the γ -alcohol intermediate undergoes rapid decarbonylation and hence only attains a low steady state concentration; and/or (ii) ether cleavage occurs before decarbonylation, with subsequent rapid decarbonylation of the resulting intermediate ${\bf k}$ in Scheme 1.

To determine whether hydrogenolysis could be induced by intramolecular hydrogen from the α - and γ -hydroxy groups, additional reactions were performed in the aprotic solvents dioxane and p-xylene (Table 5). Although **L-4** conversion in p-xylene was comparable to that in ethanol (88%, Table 5, entry 2), the carbonyl intermediate **i** (**L-3b**) was the dominant product, arising from α - and γ -dehydrogenation and decarbonylation. Very low conversion was observed in dioxane (13%, Table 5, entry 3), with **i** (**L-3b**) again the dominant product. Hydrogen donor solvents are thus critical for cleavage of ether linkages in these systems, and intramolecular hydrogen transfer is insufficient.

4-O-5; diphenyl ether (L-5). This model compound features the 4-O-5 linkage found in lignin. Transfer hydrogenolysis of diphenyl ethers is extremely challenging due to the stronger sp² C–O bond compared to those in α -O-4 or β -O-4 motifs. We thus anticipated no activity for this linker with ethanol and tested the model compound with the more reactive hydrogen donor, isopropanol. Negligible conversion of L-5 was observed at 150 °C for all catalyst or solvent (Table 6). However, at 180 °C, Pd/HT (B) afforded 27% cleavage of L-5 with full selectivity to benzene (I) and phenol (a). Surprisingly, PdCu/HT (F), which had been more active than Pd/HT for all other model compounds, afforded only 7% conversion (Table 6). We postulate that stronger adsorption of L-5 on B than on F, but additional experiments and modelling are required to elucidate this reactivity trend. In the meantime, we conclude that PdCu/HT is selective for α -O-4 and β -O-4 over 4-O-5 linkers in ethanol.

On the role of copper

The superior activity of PdCu/HT likely reflects a synergy between the metals that promotes ethanol dehydrogenation. Effective transfer hydrogenolysis requires a high rate of ethanol Table 5 Transfer hydrogenolysis of L-4 model compound containing an additional γ-hydroxy group⁶

Entry			Yield/%			
	Solvent	Conversion/%	i	e	f	
1	EtOH	98	7	90	93	
2	<i>p</i> -Xylene	88	75	5	4	
3	Dioxane	13	11	2	1	

^a Reaction conditions: 0.1 mmol L-4, 3.0 mL solvent, 2.5 mol% PdCu/HT, 50 W microwave heating at 150 °C, 6 h.

Table 6 Transfer hydrogenolysis of L-5 (4-O-5 linkage)^a

Entry	Catalyst	Solvent	Temperature/°C	Conversion ^b /%
1	Pd/HT	EtOH	150	2
2			180	2
3		iPrOH	150	22
4			180	27
5	PdCu/HT	EtOH	150	1
6			180	3
7		iPrOH	150	0
8			180	7

^a Reaction conditions: 0.1 mmol L-5, 3.0 mL solvent, 5 mol% catalyst, 50 W microwave heating, 5 h. ^b Full selectivity is observed for each entry, thus conversion is equal to the yield of a and l.

dehydrogenation, and a slow rate for the competing reaction between liberated H atoms to form (undesired) molecular $\rm H_2$. In the absence of substrate, Pd/HT is highly active for the ethanol

dehydrogenation to ethyl acetate and acetaldehyde, the former product requiring an interplay between metal and acid sites. However, literature reports evidence only modest activity for ethanol dehydrogenation over Cu/HT even at higher temperature than in this work, ¹⁰⁰ and exclusively produces acetaldehyde. Single crystal studies reveal that alloying Pd with Cu enables the spillover of H adatoms from the former to the latter, while coadsorbed CO inhibits H adatom combination and desorption over such single atom alloys that is otherwise rapid over pure Pd. ¹⁰¹ Reactively-formed CO released by the decarbonylation of intermediates **h** and **k** (Scheme 1) and/or **o** (Scheme 2a) is thus expected to suppress molecular H₂ formation over PdCu/HT *versus* Pd/HT. Future mechanistic studies will examine the sensitivity of ethanol dehydrogenation over Pd/HT and PdCu/HT to the presence of gas phase CO.

Catalyst recycling

A catalyst recyclability study for PdCu/HT (F) was conducted to explore stability. L-3a was implemented as the model substrate for this purpose, utilizing the same reaction conditions as employed in the catalyst screen (Table 4). Postreaction, the mixture was centrifuged for 5 min at 5000 rpm, the supernatant was decanted off and sampled for product analysis. Fresh substrate and ethanol were added and heated at 150 °C for 2 h.

Scheme 2 Proposed pathways from hydroconiferyl alcohol (o) to (a) ethyl-, and (b) propyl quaiacols.

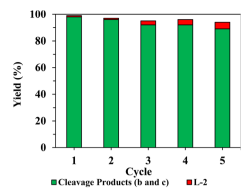


Fig. 6 Recyclability of PdCu/HT (F) for the transfer hydrogenolysis of 2-phenoxy-1-phenethanol (L-3a). Reaction conditions: substrate: 2-phenoxy-1-phenethanol (L-3a) (0.1 mmol), ethanol (3.0 mL), PdCu/HT (F) (2.5 mol%), 50 W microwave heating, 150 °C, 2 h.

This process was performed for five cycles. This method is beneficial as it does not necessitate filtration and washing, which can result in catalyst loss. Over five cycles, catalyst F retained >95% of its activity (Fig. 6), with a marginal decrease in selectivity for cleavage products due to the presence of unreacted dehydrogenated intermediate L-2. FTIR and PXRD of the used catalyst showed no change in catalyst phase or crystallite size (Fig. S19 and S20†), while TGA showed no evidence for retention of organic residues that might otherwise cause blocking of active-sites of micropores (Fig. S21†). Elemental analysis shows negligible loss of Cu or Pd (<5% of metal% by

mass) post-reaction as compared to the as-prepared catalyst. XPS data of similar post-reaction catalysts for similar applications showed a reduction of Pdⁿ⁺ to Pd⁰ after multiple cycles.⁶⁵ Note that TEM revealed a small increase in Pd nanoparticle size after reaction (2.95 vs. 2.98 nm) (Fig. 7), with a doubling of the standard deviation (0.52 vs. 0.98), indicative of limited particle sintering and loss of active surface area that likely accounts for the minimal loss of initial activity (<5%) after five reaction cycles. Even if long-term (slow) leaching Pd or Cu is observed in future studies, such hydrotalcites can be readily regenerated by dissolution in nitric acid and reprecipitation using Na₂CO₃ and NaOH.⁶⁵

Whole lignocellulose depolymerization

Application of Pd/HT catalysts for ether bond cleavage was extended to whole lignocellulose biomass (pine sawdust) through the RCF method, which circumvents the degradation of native lignin into a calcitrant structure observed during Kraft pulping. 102,103 RCF is also referred to as a 'lignin-first' approach, and has been extensively reviewed. 4,103–109 The RCF method was adapted for simultaneous: 106,107 (i) lignin extraction from whole lignocellulose using a polar-protic solvent (ethanol in this case); and (ii) selective depolymerization using Pd/CuHT (F) with ethanol as hydrogen donor. Lignin extraction and depolymerization were thus combined in a novel single step process (Fig. 8), contrasting with the conventional two-step extraction of technical lignin and subsequent selective depolymerization.

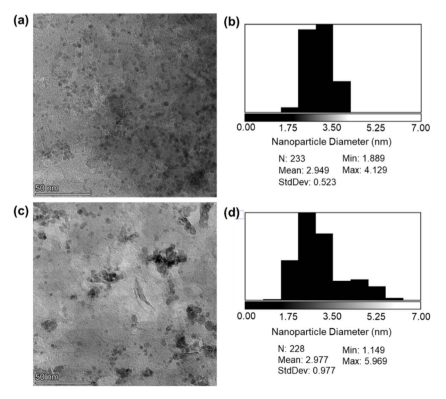


Fig. 7 TEM image of (a) fresh Pd-CuHT and (b) Pd nanoparticle size distribution of fresh 5 wt% PdCu/HT; (c) TEM image of used 5 wt% PdCu/HT after five reaction cycles for the TH of L-3a and (d) Pd nanoparticle size distribution of used 5 wt% PdCu/HT.

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Fig. 8 One-pot lignin extraction from whole biomass and integrated depolymerization in a sealed reactor.

Pine wood sawdust was selected as the biomass source, with a 'wet' lignin content of 32 wt% (see ESI eqn (1)† for detailed procedure on lignin percent determination). The theoretical maximum monomer yield for a certain substrate can be calculated by considering a simplified, linear lignin polymer in which each building block is linked to two other units by either C-C or ether bonds. Lignin depolymerization occurs by transfer hydrogenolysis of the ether bonds. For building blocks possessing two ether bonds, cleavage of both will result in a monomer. To determine the efficiency of lignin conversion to monomers, the maximum theoretical monomer yield for the whole biomass sample was first calculated by nitrobenzene oxidation (NBO), a depolymerization method for subsequent quantitative analysis of the β -ether content by NMR.¹¹⁰⁻¹¹² Note that NBO is not well-suited to determining guaiacyl/ hydrophenyl monomer yields of poplar, aspen, willow and palms lignins.112 The NBO method afforded a theoretical maximum monomer yield of 23 wt% (see ESI† for further details of ether content and theoretical monomer yield determination). This value compares favorably to a theoretical yield of 25 wt% β-

O-4 content in the lignin determined via the HSQC method.21,46,113

The one-pot reaction with pine sawdust was performed in a stainless-steel Parr 4560 stirred autoclave at 225 °C, which is sufficient for facilitating solubility and mass transfer of the whole pine biomass.114 Control reactions without catalyst (Table 7, entry 1) afford only delignification (62%, eqn (S2)†), i.e. lignin extraction from cellulose and hemicellulose, but without monomer and oligomer formation. This affirms the need for a catalyst to afford lignin depolymerization. Addition of 5 wt% Pd/CuHT (F) increased delignification to 97% (Table 7, entry 2), accompanied by a 22% monomer yield, approaching the theoretical maximum of 23% based on the NBO method discussed above (see ESI† for monomer yield determination, eqn (S3)†). The major monomer product was 4-(3-hydroxypropyl)-2methoxyphenol (hydroconiferyl alcohol, HA, o), a monolignol derived from the dominant guaiacol components of pine lignin,46 which was obtained with 64% selectivity using PdCu/ HT (14 wt% based on biomass). These values are greater than monophenolic yields of 10 wt% of initial wood mass obtained

Table 7 Reductive catalytic fractionation of pine wood sawdust using PdM/HT catalysts and ethanol^a

Catalyst (5 mol %, 0.60 g) Whole Pine Lignocellulose Cellulose/Hemicellulose Pulp + Lignin Oil (Monomers + Dimers/Oligo EtOH (50.0 mL) 225 °C, 5h Parr Reactor, 400 rpm

		Phenolic yield ^b /wt%					
Entry	Catalyst	HA^c	Other monomers	Total monomer	Dimers	Delignification/wt%	
1	N/A	0.5	<1	<1	<1	62	
2	5 wt% Pd/CuHT	14	8	22	21	97	

^a Reaction conditions: 5.0 g dry pine sawdust (32 wt% lignin), 50 mL ethanol, 0.6 g catalyst, 225 °C, 5 h. ^b Depolymerization to monomer units estimated from proportion of ether linkages in lignin content and monomer yield is a percentage of the maximum lignin present in the dried biomass (not just that present in extracted lignin). See ESI for detailed explanation for calculations of lignin percentage, delignification, maximum theoretical monomer yield, and experimentally obtained monomer yield. ^c 5-Hydroxy coniferyl alcohol.

by RCF of polar clones in acidified aqueous ethanol, albeit over a different (Pd/C) catalyst.55 The balance of monolignols in the present work was accounted for by ethyl guaiacol and propyl guaiacol. Note that while the β -O-4 model compound L-4 afforded acetophenone and guaiacol, neither product was observed in one-pot reactions with pine sawdust, which instead favoured hydroconiferyl alcohol (HA, o, Scheme 1) and ethyl/ propyl guaiacols. To explain this difference in selectivity of model compounds versus whole biomass, we suggest that in model compounds, dehydrogenation of β -O-4 moieties at the α position forms g, which can undergo subsequent dehydrogenation and decarbonylation of the γ -OH to intermediate i, or transfer hydrogenolysis and further transfer hydrogenation to j (Scheme 1). In contrast, in whole biomass, the transfer hydrogenolysis is faster than γ -OH dehydrogenation, likely due to the more hindered γ-OH and stabilization of OH by H-bonding to other OH groups in lignin. The formation of ethyl guaiacol from biomass likely reflects the dehydrogenation of o to the corresponding aldehyde, followed by decarbonylation to ethyl guaiacol (Scheme 2a). The latter pathway is evidenced by a decrease in yield of coniferyl alcohol (HA, o in Scheme 2) with increasing reaction time and a concomitant increase in the yield of ethyl guaiacol (m, Scheme 2). Alternatively, propyl guaiacol could be produced through HA dehydration and hydrogenation (Scheme 2b). Further details on the mass balance and analysis of the lignin bio-oil are provided in Fig. S22, 23 and S24-S28† respectively.

In summary, the one-pot depolymerization of whole pine biomass in the presence of 5 wt% PdCu/HT promotes both delignification and lignin depolymerization, with a nearquantitative yield of hydroconiferyl alcohol, ethyl-, and propyl guaiacol monomers. Although these monomers differ from the products obtained from their representative model compounds, their formation involves the same transfer hydrogenolysis and decarbonylation pathways. Additional studies are underway to examine our catalyst performance towards organosolv lignin. Typical RCF processes use hydride donors as stoichiometric reducing agents. In our work, the only readily available sources of hydrogen are the ethanol solvent and alcohol groups in lignin. Evidence that ethanol is the primary hydrogen source is presented in Table 5, wherein replacing ethanol with nonhydrogen donating solvents, such as p-xylenes and dioxane, drastically lowered the yields of hydrogenation products in model compound studies (EtOH: 90%, p-xylene: 5%, dioxane: 2%). We thus conclude that hydrotalcite supported PdO clusters/nanoparticles are effective for RCF of lignin and the resulting one-pot synthesis of monolignols from woody biomass. The extent of delignification and attendant monomer yields from transfer hydrogenolysis over PdCu/HT are comparable to literature reports using molecular hydrogen.

Conclusions

Lignin model substrates of α -O-4 (L-1), β -O-4 (L-2 to L-4), and 4-O-5 (L-5) linkages (Fig. 1) were used to probe the catalytic activity of Pd-HTs. α -O-4 model compound benzyl phenylether (L-1) was used as substrate for an extensive screen of Pd/HT

catalysts to determine the efficiency for transfer hydrogenolysis using ethanol as hydrogen donor. The initial catalyst screen identified PdFe/HT (C) and PdCu/HT (F) as the most active catalyst for benzyl phenylether, achieving full conversion using ethanol in under 30 min at 150 °C. This is the first system effective with ethanol as H-donor, furthermore, no base is required which is critical for making the system potentially economically viable. Similarly, a screen for the reactions that cleave the β-O-4 linker (L-2, L-3a, L-3b, and L-4) identified the same catalyst, PdCu/HT (F), as the most active and selective catalyst for both C2 and C3 type β-O-4 compounds. Extensive further optimization of reaction conditions has been carried out, as well as structure-activity studies to elucidate how catalytic activity for the multi-step process can be rationally optimized. For the 4-O-5 linkage, biphenyl ether (L-5), the best catalyst B only afforded 27% conversion. This is not surprising given the challenging nature of transfer hydrogenolysis of biphenyl ethers. Further catalyst optimization is ongoing for this linkage. The efficacy of these catalysts for the reductive catalytic fractionation of whole pine lignocellulose was also studied. PdCu/HT (F) provided the best activity, giving 97% delignification and 22% monomer yield, close to the theoretical maximum where one compound (4-(3-hydroxypropyl)-2methoxyphenol) accounted for 64% of all monomers identified. Model compound studies demonstrated that Pd doped HTs can adequately cleave ether bonds via hydrogenolysis in the absence of exogenous hydrogen. Transition metal modified HTs are a viable option for RCF of whole lignocellulose using ethanol without external hydrogen or high pressures.

Data availability

The data supporting this article have been included as part of the ESI.†

Author contributions

A. M. V.-K. conceived the research concept. D. D., R. B. C. R. and A. M. V.-K. planned the research methods. A. M. V.-K. led the project supervision, A. F. L. and K. W. provided co-supervision. D. D. led the experimental work, aided by R. B. and C. R., and undertook the data analysis. D. D. prepared figures and drafted the manuscript and revisions. A. M. V.-K., A. F. L. and K. W. edited the manuscript draft and revisions.

Conflicts of interest

There are no conflicts to declare.

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