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### **Green Chemistry**

#### COMMUNICATION



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# Flow-through solvolysis enables production of native-like lignin from biomass<sup>‡</sup>

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The inherent reactivity of lignin in conventional biomass processing commonly prevents isolation of native lignin and limits monomer yields from catalytic depolymerization strategies that target aryl-ether bonds. Here we show that flow-through solvolysis with methanol at 225°C produces native-like lignin from poplar, enabling the study of intrinsic lignin properties and evaluation of steady-state lignin depolymerization processes.

Lignin valorization is important for the economics and sustainability of biomass conversion,<sup>1, 2</sup> and catalytic deconstruction to valuable monomers is a common strategy to that end. The abundant aryl-ether linkage in lignin is the most frequent target of catalysis, but condensation reactions ultimately limit monomer yields from C–O bond cleavage.<sup>3</sup> To address this challenge, the lignin-first biorefining approach catalytically passivates reactive intermediates or uses stoichiometric reagents to functionalize the  $\beta$ -O-4 linkage.<sup>3, 4</sup> Lignin-first strategies typically yield a narrow slate of aromatic monomers, bounded by the aryl-ether content of the lignin.<sup>5</sup>

Many lignin-first biorefining studies employ reductive catalytic fractionation (RCF) in batch reactors where biomass, a reducing catalyst, and a hydrogen donor are combined in polar protic solvents. Given the typical reaction times and temperatures of RCF processes, a catalyst must be present during reaction to prevent condensation.<sup>6, 7</sup> However, physical mixing of the biomass and catalyst complicates kinetics studies and post-reaction analyses.

Recently, several reactor configurations have been applied to physically separate the biomass and catalyst for RCF processes, including conducting solvolysis and hydrogenolysis as separate batch reactions or in tandem flow-through reactors, or by using catalyst baskets in batch reactors.<sup>8-13</sup> In batch solvolysis reactions, the typical multi-

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hour residence time without catalytic stabilization results in lower monomer yields.8, 12, 13 In flow-through RCF configurations (Figure 1A), the catalyst performance is not measured at steady-state because lignin solubilization varies temporally. To overcome this, Lan et al. recently applied a protection-group method<sup>14</sup> to produce isolated lignin for subsequent depolymerization in a flow reactor.<sup>15</sup> Wang et al. also demonstrated that aqueous formic acid extracts lignin from biomass, with an estimated 75% retention of aryl-ether linkages.<sup>16</sup> Generally, the ability to isolate nativelike lignin for lignin property studies and to evaluate steadystate catalytic lignin depolymerization activity with intact C-O linkages would be useful, but with these few exceptions, the ability to isolate native-like lignin in a solubilized form has not been widely reported to our knowledge. In addition, reducing solvent usage in RCF is critical for process economics,<sup>17</sup> and the ability to isolate a native-like lignin may facilitate solvent recycling and enable more efficient use of a downstream catalytic step, thus reducing the costs in an RCF process.

In the present study, we hypothesized that solubilized lignin from whole biomass at typical RCF conditions must be rapidly exposed to a catalyst and hydrogen, otherwise it will undergo condensation.<sup>6, 7</sup> To test this hypothesis, we conducted RCF reactions in either an *in situ* or *ex situ* mode with a multi-bed flow system (Figure 1) and in batch reactors. Throughout, 'in situ' denotes that the lignin is exposed to the catalyst directly after solvolysis (either in batch or flow), and 'ex situ' denotes that lignin is first isolated, then in a separate processing step, subjected to hydrogenolysis. Throughout, we used methanol as a solvent, hybrid poplar as the substrate (Table S1), a reaction time of 3 h, a reaction temperature of 225°C, and 15 wt% Ni/C as the catalyst. The methanol-to-biomass ratio used in all batch and flow reactions was ~93 L/kg. All lignin monomer yields are reported on a wt% basis with respect to the total lignin content. The materials and methods are detailed in the Electronic Supplementary Information (ESI).

We first benchmarked the aryl-ether bond content in poplar lignin based on aromatic monomer yield from *in situ* batch and flow-through RCF reactions. From these experiments, we obtained monomer yields of  $36.8\pm0.2\%$  in a batch

reaction and 31.6 $\pm$ 1.3% in a flow-through RCF experiment (**Figure 2, Tables S1-S3**). For all flow-through reactions, we ran at temperature for 3 h, which resulted in a delignification extent of 63.1% $\pm$ 0.1% (duplicate). A control *in situ* batch reaction with an activated carbon support alone resulted in a monomer yield of 11.3 $\pm$ 0.3%.



Figure 1. Schematics of flow-through solvolysis and reductive catalytic fractionation (RCF) configurations used in the present study. (A) In situ flow-through solvolysis of biomass with methanol and hydrogenolysis with a Ni/C catalyst. (B) Ex situ flow-through solvolysis setup in flow, where catalysis is subsequently conducted in batch reactors or a flow reactor with isolated solvolysis liquor.

To compare the degree of lignin condensation without employing hydrogenolysis immediately upon extraction, we conducted an *ex situ* batch solvolysis reaction. The isolated lignin oil was then subjected to batch hydrogenolysis, where we obtained a monomer yield of  $18.6\pm0.2\%$  (**Table S2**). Relative to the *in situ* reactions, this result indicates that substantial lignin condensation (compared to  $36.8\pm0.2\%$ , above) occurs over 3 h without catalyst and hydrogen.

To examine condensation extents when lignin is rapidly quenched, we produced solvolysis liquor with methanol over multiple biomass beds at 225°C via ex situ flowthrough solvolysis (Figure 1B). We collected the solvolysis liquor in a knockout pot, pooled the liquor from 12 consecutive reactor beds (producing ~5.5 L total), and stored the sample at room temperature in a translucent LDPE bottle. After one week of storage at room temperature, we performed batch and flow-through hydrogenolysis on the solvolysis liquor, again using 15 wt% Ni/C at an equivalent catalyst to biomass ratio at 225°C. Much to our surprise, the monomer yields from these experiments were 35.2±0.0% and 32.3±0.0%, respectively, similar to the *in situ* RCF results (Figure 2, Table S2). A control ex situ flow-through experiment over the carbon support yielded 6.6±0.4% monomer yield.

The observation of similar monomer yields from *in situ* and *ex situ* RCF reactions suggests that flow-through solvolysis retains intact  $\beta$ -O-4 linkages in lignin. The *ex situ* liquor did not contain coniferyl alcohol, sinapyl alcohol, or any of the monomers shown in **Figure 2** at detectable levels,

suggesting that flow-through solvolysis alone does not produce monomers. To ascertain the presence of intact arylether linkages, we conducted 2D hetero-nuclear single quantum coherence (HSQC) NMR spectroscopy.<sup>18</sup> As shown in Figure 3, the HSQC NMR spectra of the ex situ flow-through solvolysis liquor exhibits intact aryl-ether linkages (blue). We note that partial methoxylation of the  $\alpha$ -C (in the  $\beta$ -O-4 linkage) is observed, suggesting that methanol protection is occurring at this position.<sup>8</sup> Conversely, both the in situ flow-through RCF oil and the ex situ flow-through RCF oil exhibit complete disappearance of aryl-ether linkages (Figure S2). Overall, the lignin extracted by ex situ flow-through solvolysis produces similar RCF monomer yields and NMR spectra compared to the original poplar, suggesting that the extracted lignin is a "native-like" substrate.



Figure 2. Monomer yield and selectivity data from in situ and ex situ RCF experiments in batch and flow-through modes. (A) Results from in situ batch and flow-through RCF experiments, including an in situ flow-through control reaction with biomass and carbon support in methanol. (B) Results from ex situ flow solvolysis and subsequent batch and flow-through hydrogenolysis reactions after 1 week of solvolysis liquor storage, as well as an ex situ control of solvolysis liquor run over carbon support only in flow. Product selectivity is shown for syringyl monomers in blue and guaiacyl monomers in green. These data are also provided in **Table S2**. All experiments were conducted in duplicate, and the error bars are the range for total monomer yield. Batch reactions: either 30 mL (23.76 g) of ex situ solvolysis liquor or 0.313 g poplar in 30 mL methanol, 0.05 g catalyst (either 15 wt% Ni/C or activated carbon support), 30 bar H<sub>2</sub> at 225°C, 3 h (exclusive of 30 min heating ramp). Flow reactions: 2 mL min<sup>-1</sup> methanol or ex situ solvolysis liquor, 5 g poplar (in situ) or no biomass (ex situ), 0.9 g catalyst (either 15 wt% Ni/C or activated carbon support), 1,600 psig, 200 sccm H<sub>2</sub>, 225°C, 3 h (exclusive of 1 h heating ramp for in situ runs).

We were additionally interested in the duration that the solvolysis liquor could be stored and to understand if the methanol could be removed from the solvolysis liquor without compromising lignin reactivity. These questions are relevant for downstream chemistries that require catalytic processing in different solvents, processing neat RCF oil, studies of lignin properties, and applications of native-like lignin in materials or other direct-use applications.



Figure 3. NMR spectra of solvolysis liquor demonstrates the production of native-like lignin. (A) 2D HSQC NMR spectrum of the native poplar biomass sample. (B) 2D HSQC NMR spectrum of the ex situ flow-through solvolysis liquor.

To address these questions, we conducted a 12-week experiment in which a fraction of the solvolysis liquor was maintained at room temperature in methanol, and duplicate ex situ batch RCF reactions were conducted after storage for 1 (vide supra), 2, 3, 5, 8, and 12 weeks. For a fraction of the solvolysis liquor, we used rotary evaporation to produce a methanol-free oil. At three time points over the same 12 weeks, we reconstituted a fraction of this solvent-less solvolysis liquor in methanol for conducting equivalent ex situ batch RCF reactions. Much to our delight, in both the stored samples in methanol and the reconstituted samples, the monomer yields are essentially invariant over the 12week period, with only a  $1.8\pm0.1\%$  and  $2.0\pm0.0\%$  decrease in monomer yields in the 8- and 12-week reconstituted samples, respectively, as shown in Figure 4 and Table S3. We also characterized the molecular weight distributions of the ex situ flow-through solvolysis liquor after batch hydrogenolysis experiments as-is and reconstituted after 1, 5, 8, and 12 weeks (Figure S3). As shown, the posthydrogenolysis molecular weight distributions are invariant as a function of storage time. Together, these data indicate that the solvolysis liquor is shelf-stable at least for 3 months.



Figure 4. Time-resolved study of ex situ batch RCF reactions on solvolysis liquor produced in flow and stored in methanol or reconstituted after solvent evaporation (denoted as RC). The monomer yield and selectivity for batch hydrogenolysis reactions of solvolysis liquor produced in flow. The as-made flow solvolysis liquor was tested at 1, 2, 3, 6, 8, and 12 weeks. Removing the methanol before aging was also tested at 5, 8, and 12 weeks. These data are also provided in **Table S3**. All experiments were conducted in duplicate, and the error bars are the range of the total monomer yield. Reaction conditions: 30 mL ex situ solvolysis liquor, 0.1 g 15 wt% Ni/C catalyst, 30 bar H<sub>2</sub> at room temperature, 225°C, 3 h (exclusive of 30 min heating ramp).

One application of the *ex situ* solvolysis liquor is the ability to conduct flow experiments where the catalyst subjected to a continuous feed of uniform lignin. As shown in **Figure 5**, the *ex situ* solvolysis liquor, including a reconstituted sample and two stored samples (4 and 7 weeks of storage in methanol), exhibits a linear increase in monomer yield (using the total lignin fed as the baseline for yield), while the nonlinear increase apparent for the *in situ* flow-through solvolysis indicates a transient lignin stream. As also noted by Lan *et al.*<sup>15</sup> and Wang *et al.*,<sup>16</sup> this feature will facilitate extended catalyst activity and stability studies.

While an in-depth exploration of solvolysis residence times is outside the scope of this work, it is worth considering the present results with respect to previous batch solvolysis work. Previous batch solvolysis residence times have been 2-3 h.<sup>8, 12, 13</sup> The flow-through solvolysis reported here employed a 5 g biomass bed centered inside a tube of approximately 35 cm total length and 1.6 cm inner diameter. The 5 g bed occupied approximately 5 cm in the axial dimension, thus the lignin traveled through a heated zone ranging between 15 and 20 cm in length, depending on where in the biomass bed it originated. At a 2 mL min<sup>-1</sup> flow rate of methanol, the mean residence time for the lignin fragments to be quenched was thus  $\sim 17$  min.



Figure 5. Time-on-stream measure of cumulative monomer yields for in situ and ex situ hydrogenolysis. Ex situ flow-through RCF provides a consistent feed to the catalyst bed, resulting in a linear increase in monomer production, compared to the nonlinear, transient monomer production during *in situ* flowthrough RCF. These data are also provided in **Table S4**. All experiments were conducted in duplicate, and the error bars show the range. Reaction conditions: 2 mL min<sup>-1</sup> methanol or ex situ solvolysis liquor, 5 g poplar (*in situ*) or no biomass (ex situ), 0.9 g 15 wt% Ni/C, 1,600 psig, 200 sccm H<sub>2</sub>, 225°C, 3 h (not including 1 h temperature ramp for *in situ* runs). Ex situ runs included a 1.5 h methanol flush at the end of the reaction. 4week and 7-week ex situ runs profiles are offset by 1 h and 2 h time-on-stream, respectively, for visual clarity.

Additional studies will ascertain if this *ex situ* solvolysis approach to produce native-like lignin is sufficiently general. A system like that used here will be challenging to directly implement at scale, given that the solvent-tobiomass ratio is exceptionally high relative to the predicted range needed for industrially relevant operation.<sup>17</sup> However, these results suggest that native-like lignin extraction may be possible with appropriate reaction engineering measures to rapidly quench solvolysis liquor, which will be pursued in future studies. Also, the ability to isolate native-like lignin demonstrated here enables steady-state catalyst evaluation, potentially including with neat lignin oil and in other solvents after methanol removal. Moreover, the current setup is a convenient approach to isolate native-like lignin from different feedstocks and with different solvents.

#### Conclusions

Overall, this study demonstrates that flow-through solvolysis can produce native-like lignin using poplar and methanol at 225°C. Specifically, reductive catalytic treatment of the isolated lignin from a flow-through system produces monomer yields equivalent to a two-stage *in situ* flow-through setup, indicating that the aryl-ether bonds are conserved in the flow-based methanol extraction. From a lignin-first biorefining research perspective, these results demonstrate that immediate catalyst action is not necessary for passivation of reactive components of lignin, and that the lignin extract can be dried to an oil for storage or stored in methanol and processed later without losing substantial reactivity. Similarly, it may also be feasible to use the extracted lignin in materials. Taken together, these results suggest that lignin can be successfully isolated without significant chemical modification, which can enable both improved understanding of lignin structure and continuous catalytic processing of this important biopolymer.

#### Author Contributions

DGB, JSK, NET, GGF, JKK, RJD, ARCM, TR, NCC, RMH, RK, TBV, and DGW executed the study. The paper was written by DGB, JSK, and GTB with contributions from all authors.

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