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## Guidelines for performing lignin-first biorefining

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The valorisation of the plant biopolymer lignin is now recognised as essential to enabling the economic viability of the lignocellulosic biorefining industry. In this context, the “lignin-first” biorefining approach, in which lignin valorisation is considered in the design phase, has demonstrated the fullest utilisation of lignocellulose. We define lignin-first methods as active stabilisation approaches that solubilise lignin from native lignocellulosic biomass while avoiding condensation reactions that lead to more recalcitrant lignin polymers. This active stabilisation can be accomplished by solvolysis and catalytic conversion of reactive intermediates to stable products or by protection-group chemistry of lignin oligomers or reactive monomers. Across the growing body of literature in this field, there are disparate approaches to report and analyse the results from lignin-first approaches, thus making quantitative comparisons between studies challenging. To that end, we present herein a set of guidelines for analysing critical data from lignin-first approaches, including feedstock analysis and process parameters, with the ambition of uniting the lignin-first research community around a common set of reportable metrics. These guidelines comprise standards and best practices or minimum requirements for feedstock analysis, stressing reporting of the fractionation efficiency, product yields, solvent mass balances, catalyst efficiency, and the requirements for additional reagents such as reducing, oxidising, or capping agents. Our goal is to establish best practices for the research community at large primarily to enable direct comparisons between studies from different laboratories. The use of these guidelines will be helpful for the newcomers to this field and pivotal for further progress in this exciting research area.

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### Broader context

Conversion of polysaccharides from lignocellulose to biofuels and chemicals has long been the primary objective of the biomass conversion community. However, for the success of a global bio-based circular economy that employs lignocellulosic biomass as a feedstock, it is imperative to also derive value from the lignin component, beyond low-value heat and power. Despite a century of lignin research, this goal remains elusive, and yet is increasingly important. The international research community is now vigorously pursuing lignin valorisation approaches in response to the need for a more sustainable carbon economy. The use of active stabilisation methods for lignin fractionation and valorisation, generally dubbed “lignin-first” biorefining, is receiving significant attention. Here we provide our perspective on how to unify the lignin-first community around a common set of research practices to accelerate progress in this field.

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# 1. Introduction

It is now clear that humankind must rapidly transition from the use of fossil resources, given concerns over anthropogenic climate change. Accordingly, the development of alternative and sustainable feedstocks for energy, chemicals, and materials has become one of this century's most important societal challenges.<sup>1,2</sup> To that end, plant-based feedstocks are one of the most well-studied and promising green carbon sources available in sufficient quantities to contribute to a more sustainable global carbon economy without negatively affecting the climate.<sup>3</sup>

Large amounts of lignocellulose are processed today for pulp and paper production and, in some parts of the world, for heat and power. In recent decades, substantial governmental and industrial investments have created a strong driving force to commercialise the production of cellulosic biofuels and bioproducts, which has mobilised a large global research community to focus on this problem. Most of the leading lignocellulosic biofuel production paradigms that have employed selective approaches in the last two decades focused on the conversion of plant polysaccharides to biofuels, such as ethanol, gasoline, diesel, and jet fuel, and have slated the high energy density aromatic biopolymer, lignin, for on-site heat and power production. This approach is mirrored by pulp and paper mills that also combust lignin for power and to recover inorganic pulping chemicals.<sup>4,5</sup> Lignin combustion has long been thought to be the most viable approach for dealing with this biopolymer, given the challenges associated with isolating lignin from polysaccharides without making it more recalcitrant, and the difficulty in overcoming the inherent heterogeneity of lignin – the two main challenges facing lignin valorisation research efforts spanning the last century.

The canonical lignin building block is generally a phenylpropanoid group that usually exhibits one of three aromatic structures: syringyl (S), guaiacyl (G), and hydroxyphenyl (H) units. Each of these units derives from one of the monolignols: sinapyl alcohol, coniferyl alcohol, and *p*-coumaryl alcohol. Several types of C–O and C–C linkages are formed between these units during lignin biosynthesis to build the native polymer. In its native state in the plant, lignin is now regarded as being less branched than previously surmised,<sup>6–9</sup> and more tractable. Additional building blocks have been shown in the past decade to be incorporated into lignin, including flavonoids, hydroxystilbenes, and others.<sup>10</sup> Several lignin model structures are shown in Fig. 1 to illustrate some of the key features.

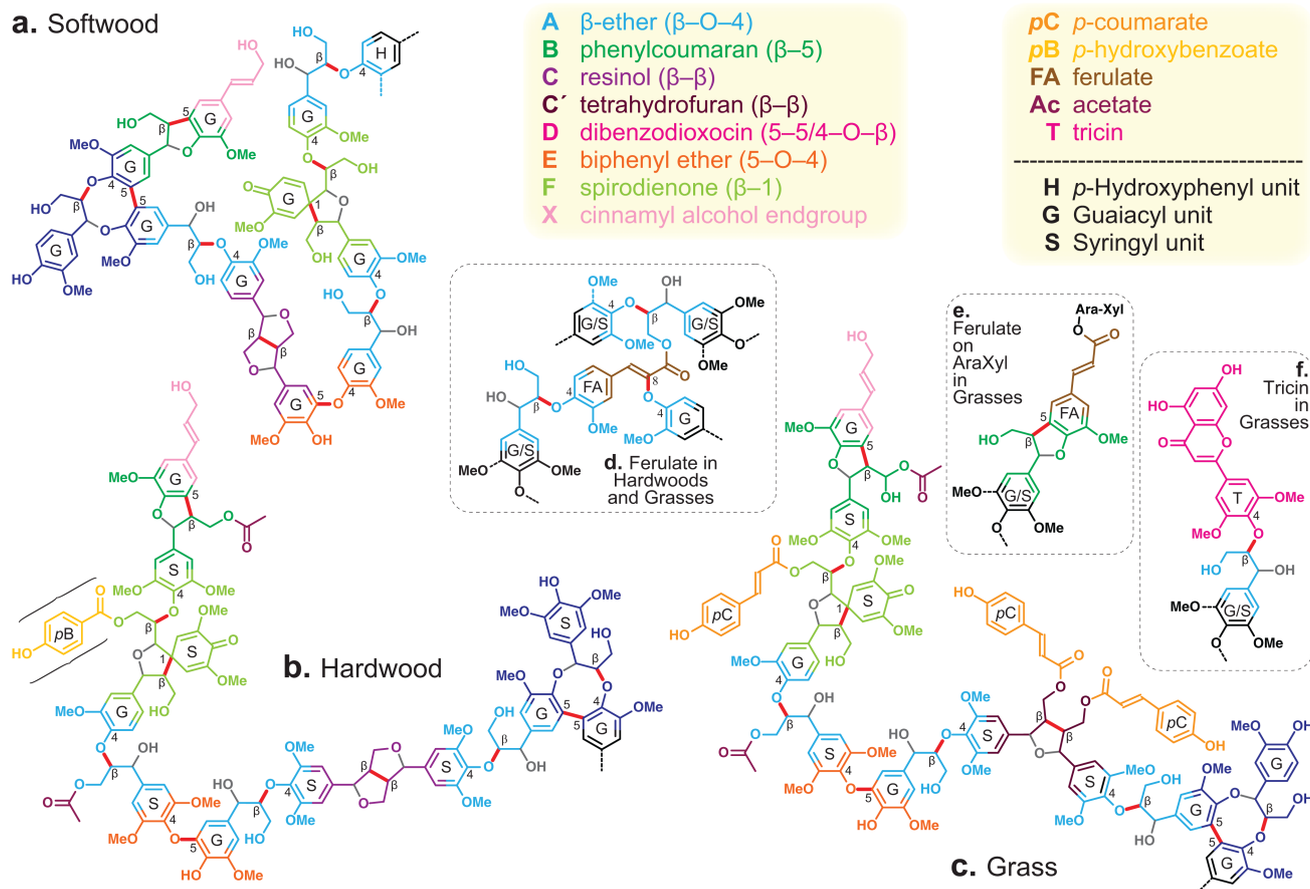
As lignin represents the largest source of sustainable aromatics on the planet, it is increasingly being recognised as foolhardy to ignore its potential value.<sup>11,12</sup> As a result, many researchers today are pursuing more holistic strategies for biomass utilisation that place substantial value on both lignin and polysaccharides, in many cases using selective fractionation technologies. Much of the motivation for this work is underpinned by technoeconomic analyses (TEA) and life-cycle assessments (LCA) that indicate that lignin valorisation can improve the overall biorefinery economics and sustainability footprint.<sup>13–16</sup> A fundamental challenge in

selectively fractionating biomass into its constituents is the propensity of carbohydrates, lignin, or both to degrade to undesirable products during processing. More efficient methodologies to fractionate and conserve cellulose, hemicelluloses, and lignin are therefore desirable.

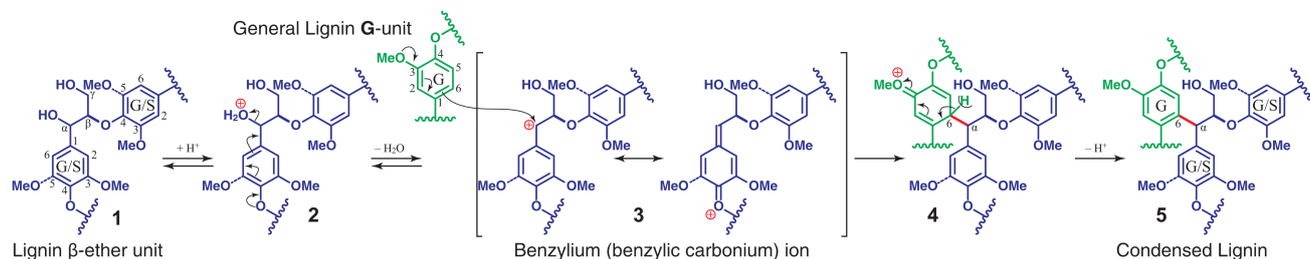
For selective lignin removal from intact lignocellulosic biomass, many fractionation strategies have been developed over decades of research.<sup>17–19</sup> By far the most common are organosolv processes, generally defined as processes that employ an organic solvent, most commonly with an acid co-catalyst and water.<sup>20,21</sup> These pretreatment strategies often liberate substantial amounts of lignin from biomass. Given the typical requirement for acidity, ether and ester bonds may be cleaved and, through reasonably well-understood mechanisms, condensation reactions occur to an extent dependent on the fractionation conditions (pH, residence time, solvent). As a result, non-native lignin-derived polymers (a.k.a. technical lignins) are formed.<sup>9,22–31</sup> The rationale for this condensation is that the solvolysis conditions, catalysed by acid, in addition to being able to cleave weak C–O linkages prevalent in the lignin, protonate the  $\alpha$ -OH in various lignin structures, leading to the ready formation of benzylic carbocations (benzyl cations, Scheme 1). These reactive intermediates readily participate in electrophilic aromatic substitution reactions on the electron-rich aryl groups of lignin to form recalcitrant C–C bonds that are not found in the native lignin; an equivalent chemical explanation is that the electron-rich aromatic rings readily participate in nucleophilic attack on the carbocation centres.<sup>24</sup> The insidious aspect of this reaction sequence is that, although phenolic benzyl cations are more stable than their etherified counterparts, the latter still easily form, so condensation can therefore occur within the chain, not just on endgroups. Similarly, although the nucleophilicity of a free-phenolic unit is higher, and will therefore more rapidly attack such carbonium ions, etherified units also readily undergo the condensation reaction. The propensity to spontaneously undergo these reactions in acid explains how condensed lignins result and why conditions that are sufficiently harsh to extract the lignin generally produce particularly intractable polymers. These types of organosolv processes, for which there is a huge number of variations, typically yield low-quality lignin streams that are, because of these condensed units, not suitable for subsequent depolymerisation processes. Typical thermochemical pretreatment methods, such as hot water, steam explosion, and acid pretreatment, that focus almost solely on maximising monomeric carbohydrates, lead to many of the same condensed lignin structures (in part because of the native acids released from esters in the cell wall during such pretreatments), rendering lignin into a more recalcitrant polymer than its starting native form in the plant cell wall. High-severity alkaline conditions will similarly lead to undesirable lignin condensation.<sup>9,32–36</sup>

Lignin-first processing is the broadly accepted umbrella term for solvent-based methods in which lignin preservation, together with that of the polysaccharides, is considered upfront, moving away from the current practice of having to deal with an intractable lignin product at the end of a





**Fig. 1** Model lignin structures containing 11 units for: (a) a softwood, (b) a hardwood, and (c) a grass. These structures in no way represent the proportion of the various functionalities that are found analytically, but rather show the major and more interesting lignin units with 'legal' inter-unit bonding. Notable features include: (a) a dibenzodioxocin unit **D**, free-phenolic, from 5-5-coupling of oligomer units followed by 4-O- $\beta$ -coupling with a single coniferyl alcohol monomer (thus not really creating a "Y-type" branchpoint), a spirodienone **F** from  $\beta$ -1 coupling, and a 4-O-5 (biphenyl ether) unit **E** also from the coupling of oligomer units but, again, not producing a real branchpoint because it is found only in its free-phenolic form shown; (b) the same range of units as for the softwood, but involving S-units where appropriate, and also showing, acylating a  $\gamma$ -OH, one acetate and one *p*-hydroxybenzoate, the latter being a feature in primarily poplar/aspens, willow, and palms; (c) a similar range of units again, except with the  $\beta$ - $\beta$ -coupled unit arising from initial dimerisation of sinapyl *p*-coumarate to produce a tetrahydrofuran **C'** rather than the resinol **C** that results from dimerisation of (un-acylated) monolignols, showing the natural lignin acylation by acetate and *p*-coumarate, features in all grasses; and inserts: (d) one of many structures involving ferulate units **FA** derived from lignification with monolignol ferulate conjugates that is a feature of all grasses and some hardwoods; (e) one of many units derived from the cross-linking of lignin and arabinoxylan *via* ferulates acylating the latter, a feature of all grasses (and commelinid monocots); (f) the triclin **T**, a flavone, that acts as a chain-starter in 'all' grasses. The figure was modified from Ralph *et al.*<sup>8</sup>



**Scheme 1** Mechanism for the presumed major condensation reaction occurring under acidic aqueous conditions. A lignin  $\beta$ -ether unit **1** readily forms resonance-stabilized benzylium (benzyl carbocation, or benzyl carbonium) ions **3** following protonation of the  $\alpha$ -OH to produce intermediate **2**. Nucleophilic attack of a general lignin G-unit on this carbonium ion or, as alternatively viewed, electrophilic aromatic substitution by the carbonium ion on the G-unit, produces an intermediate **4** that rearomatizes by losing a proton to produce **5**, a new 'condensed unit' with a particularly recalcitrant non-native 6- $\alpha$ -bond in so-called condensed lignins. G-units are favoured over S-units for attack by carbonium ions because of their more accessible (less sterically encumbered) 6-positions. Lignins that have undergone such condensation are difficult to degrade to monomers. Note: the numbering here is specific to this figure.



biorefining process. Here we define 'lignin-first' as an active stabilisation approach that liberates lignin from the plant cell wall and prevents condensation reactions through either catalysis or protection-group chemistry. Importantly, lignin-first biorefining is not a synonym for lignin valorisation, but rather an integral approach that derives value from both lignin and polysaccharides, towards an atom-efficient and more sustainable utilisation of lignocellulosic biomass. Most commonly, lignin-first processes involve three steps: (i) the lignin is removed from whole biomass using an organic solvent through solvolysis or acid-catalysed reactions (similarly to organosolv pretreatment); (ii) the resulting intermediates are stabilised, with the intention of preventing condensation of reactive species generated by lignin depolymerisation, and (iii) further depolymerisation occurs if not fully depolymerised at the stabilisation stage.<sup>9,37–39</sup>

To date, there have been several approaches reported for lignin-first refining, the chemical steps of which are illustrated in Fig. 2. The most common methodology comprises solvent-based lignin extraction from biomass in the presence of a transition metal under hydrogen atmosphere or with the aid of a hydrogen-donor solvent or another reducing agent.<sup>40–44</sup> This methodology, which has been termed 'Catalytic Upstream Biorefinery' (CUB) or 'Early-stage Catalytic Conversion of Lignin' (ECCL) for the process using 2-propanol as an H-donor,<sup>9,40,45–50</sup> is generally now termed Reductive Catalytic Fractionation regardless of the H-source (RCF, Fig. 2a). It should be noted that RCF carried out under H<sub>2</sub> pressure was first practiced/developed in the late 1930s and 1940s as a methodology to study lignin,<sup>51–57</sup> and as a means for pulping and high-yield production of lignin-based chemicals, but this was not commercialized at the time.<sup>58–61</sup> Variants of the RCF approach include systems in which sugars in the biomass operate as reducing agents,<sup>62</sup> and recently flow-through operations or catalyst baskets have been applied such that biomass solvolysis and hydrogenation/hydrogenolysis reactions on lignin intermediates have been separated in time and space.<sup>63–67</sup> The primary roles of the metal catalyst are reductive stabilisation of reactive intermediates from lignin that result from the solvolysis process and depolymerisation of the solubilised lignin oligomers. Solid(biomass)/solid(catalyst) contact is therefore not essential, as demonstrated in reactor set-ups that physically separate the solid biomass and catalyst.<sup>63,65,66</sup> The solvent mixture and the potential presence of an additional acid or base catalyst can influence the yield of monophenolic compounds and (hemi)cellulose retention (Fig. 2).<sup>58,68</sup> The solvent determines not only the delignification degree (*i.e.*, solvolysis and extraction of lignin) but also the retention of (hemi-)cellulose as a pulp, as well as the selectivity and distribution of monophenolics.<sup>38</sup> Usage of water, or protic solvents with a high proportion of water, may produce high delignification but will also hydrolyse and solubilise carbohydrates, which can be hydrogenated by the catalyst or react with the solvent.<sup>40,46,47,69</sup>

Other active stabilisation approaches during acidic fractionation in the lignin-first sphere comprise the rational use of protection-group chemistries.<sup>70,71</sup> Acid-catalysed depolymerization of lignin *via* acidolysis of the  $\beta$ -O-4 linkage proceeds through two pathways,<sup>72,73</sup> one of which results in the

formation of the so called Hibbert's ketones,<sup>74,75</sup> and the other delivers a C2 aldehyde (as mixture of H, G or S, depending on wood), which is unstable under the reaction conditions. Trapping this C2 aldehyde by using diols, typically ethylene glycol, to form more stable cyclic acetals results in suppressing recondensation phenomena and very high selectivity to the corresponding C2-acetals.<sup>76</sup> This has been shown on a variety of organosolv lignins,<sup>77,78</sup> but also in a 'metal-free' lignin first process, dubbed here diol-assisted fractionation (DAF), where C2-acetals are directly obtained from lignocellulose under carefully selected reaction conditions.<sup>79</sup> Newly generated aldehydes produced in a process may be protected as acetals by using ethylene glycol (Fig. 2b).<sup>76,79</sup>

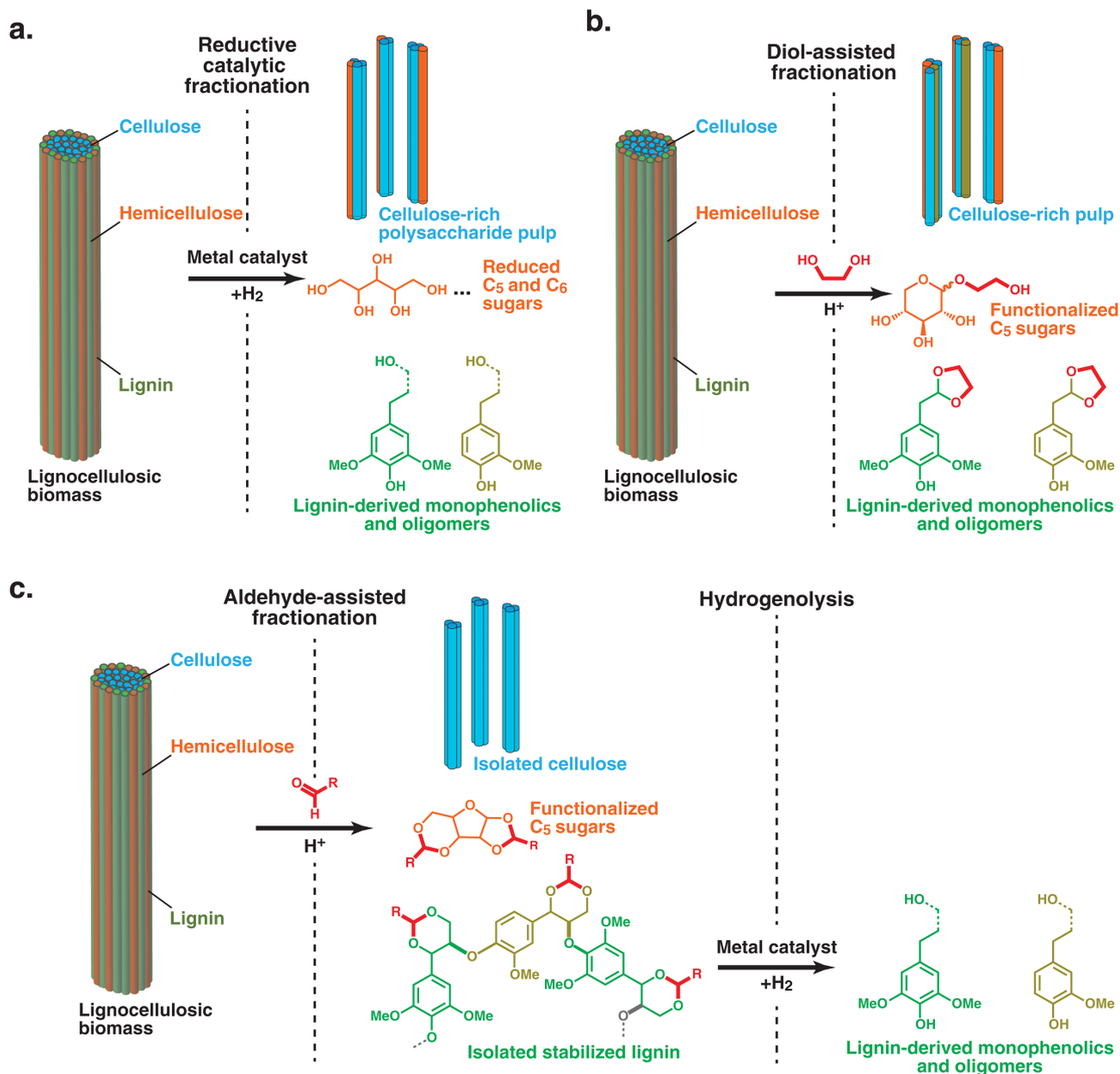
Alternatively capping the benzylic alcohol, for example by exploiting the natural 1,3-diol in lignin sidechains to produce an acetal with formaldehyde or other simple aldehydes, stops benzylic cation production, dubbed aldehyde-assisted fractionation (AAF) (Fig. 2c).<sup>24,80</sup> The latter strategy, which involves capping with stoichiometric reagents, enables further downstream depolymerisation and transformations of chemically stabilised lignin. Hydrogenolysis reactions for example, can be used to yield narrow product distributions.<sup>24,59,80,81</sup>

Efforts towards developing the lignin-first concept for fractionation of lignocellulosic biomass have significantly accelerated in the past several years and new studies are continuously being published in this area by the global biomass conversion community. A challenge in this emerging field is that there are no commonly accepted standards for the choice of feedstocks, product analysis, or evaluation of process performance.<sup>82</sup> This is a severe limitation as: (1) quantitative comparison of methodologies and results between laboratories is challenging, (2) reproduction of other research groups' procedures becomes difficult or even unfeasible, and (3) standardised lignocellulosic materials are not available. We therefore posit that to advance this research field from fundamental studies to a de-risked technology that industry could harness to fractionate and valorise lignocellulosic biomass, best practices should be established and implemented across the lignin-first research field. In this perspective, we thus present a set of recommended guidelines to establish common practices in this promising research direction for lignin valorisation. This perspective is organised into sections describing feedstock preparation and analysis, reactor configurations for performing lignin-first processing, measuring the efficiency of the catalyst performance, and determining product yields and mass balances. We conclude with future perspectives and propose several next steps to further advance the growing lignin-first biorefining field.

## 2. Feedstock preparation and characterisation

In simple chemical reactions, mass or mole balances are straightforward to establish both for products and reactants, which makes the calculation of typical reaction parameters, such as conversion, yield, and selectivity, straightforward to accomplish.





**Fig. 2** Three lignin-first strategies reported to date using solvolysis and catalytic stabilization of reactive intermediates to stable products or protection-group chemistry and subsequent depolymerisation. (a) Reductive catalytic fractionation (RCF) relies on the use of a metal catalyst and  $H_2$  (or a hydrogen donor) in a polar, protic solvent to selectively extract lignin from the cell wall, which is depolymerised and further stabilised *via* reduction chemistry. This results in the direct production of monophenolics as well as the reduction of any carbohydrates that are solubilized to hydrogenated sugars and sometimes polyols. (b) Diol-assisted fractionation (DAF) relies on the use of an appropriate solvent (typically non-protic), acid catalyst (triflic acid, metal triflates, or sulphuric acid) and diol (typically ethylene glycol).<sup>79</sup> During this process lignin is depolymerized by acidolysis of the  $\beta$ -O-4 moiety (C2 and C3 pathways) followed by the trapping of the unstable C2-aldehyde species in the form of their more stable (typically cyclic) acetals.<sup>76</sup> (c) Aldehyde-assisted fractionation (AAF) similarly involves fractionation using a non-protic solvent and an acid (typically HCl or  $H_2SO_4$ ) but in the presence of an aldehyde, which reacts with the diol on the  $\beta$ -O-4 structure to form an acetal. The acetal prevents condensation reactions during fractionation to yield stabilized lignin oligomers that can then be depolymerized to lignin oligomers at yields comparable to RCF.<sup>83</sup>

With biomass conversion, however, these parameters are more difficult to establish due to the solid, heterogeneous nature of the substrate. Nevertheless, to accurately quantify yields, it is crucial to characterise the biomass thoroughly before conducting further experimental work. The following subsections propose guidelines to report the feedstock characteristics accurately.

We note that the US Department of Energy's National Renewable Energy Laboratory (NREL) has published Laboratory Analytical Procedures (LAPs) for the relevant techniques described in Sections 2.1 and 2.2.<sup>84-91</sup> Overall, we suggest that these methods be followed as a consistent means to prepare and characterize biomass feedstocks. The NREL website also



contains Microsoft Excel-based spreadsheets to use these LAPs. While ASTM International has published similar procedures, NREL's LAPs are periodically updated to reflect new advances in the field of biomass analytics.

### 2.1 Sourcing and preparing the feedstock

The raw material understandably has a profound effect on the outcome of any transformation. In the case of lignocellulosic biomass, this is even more important as the biomass will differ depending on factors that, for woody biomass, include: species, age of the wood, sapwood or heartwood, bark content, regional factors, and time of harvesting. For agricultural residues, it is necessary to again know if the biomass consists of stems and/or leaves, contains seeds or cobs, and the physiological state of the harvested plants. For all biomass feedstocks, the source must be specified. When possible, referencing the plantation establishment and maintenance should be included: namely, watering practices, fertiliser, and herbicide use, as well as growth stimulation. Harvesting, initial processing, handling, and storage conditions of biomass should be described including the dimensions of the sample and what anatomical fractions of the plant have been used, and whether it is whole (above-ground) biomass.

Before storing biomass in the laboratory for extended periods of time, the feedstock is typically dried, according to the NREL/TP-510-42620.<sup>86</sup> This method employs air drying, a convection oven, or lyophilisation. Fortunately, cell wall material is reasonably stable, but care should be taken to dry the material, to < 10% moisture content, before fungal infection can occur. Biomass is subsequently knife-milled (Wiley-mill or other) to pass through a 2 mm screen. This biomass is then milled further to pass through a 20 mesh (1 mm) screen affording ~300–600 µm particles. This sizing usually obeys the rules of the minimal suspension criteria during its processing in stirred tank reactors typically used in the laboratory (> 50 mL).

The next step is to determine moisture content on a mass basis, *via* NREL/TP-510-42621 at 105 °C.<sup>88</sup> Oven-drying to a constant weight, in triplicate, is the recommended method and can be combined with NREL/TP-510-42622<sup>87</sup> to determine ash content and limit sample consumption. Each sample requires ~0.50 g (accurately weighed) of dried biomass. Even after the initial moisture content of the samples is determined, this measurement must be taken before each of the following procedures to accurately correct to a dry-weight basis and account for any changes to the moisture content due to humidity, storage, or location changes.

Extractives may be removed in ethanol/water by simple sonication treatment, Soxhlet extraction, or by using Dionex™ accelerated solvent extraction (ASE) *via* NREL/TP-510-42619.<sup>84,92–94</sup> The use of flow-through systems in which the solvent composition can be varied systematically is advantageous for the selective removal of extractives.<sup>95</sup> The typical procedure involves Soxhlet extraction with water and then ethanol.<sup>84,92,93</sup> Also popular (but not in the aforementioned NREL LAP) is a method in which the ground biomass is suspended in 80% v/v EtOH/H<sub>2</sub>O and sonicated.<sup>94</sup> The amount of water and ethanol extractives is determined by mass loss

in the recovered biomass sample. In poplar wood, for example, these extractives account for 7–8% of the biomass by weight, ranging from roughly 2–10% for various woods.<sup>96</sup> It is not necessary to analyse extractives in detail, although metabolite profiling may be useful, especially for transgenic materials or if extractives themselves are targets for further valorisation.<sup>97–100</sup>

For seed or other fatty-acid-rich samples, a hexane or chloroform extraction is also necessary. Protein-rich samples benefit from initial water extraction as proteins may aggregate, denature, and precipitate, so becoming difficult to subsequently remove if initially treated with organic solvents.<sup>101</sup> We note that protein content can be estimated as well *via* NREL/TP-510-42625.<sup>85</sup>

After determination of all other measurable components, lignin, cellulose, and hemicellulose content in the biomass feedstock sample (*vide infra*), the remaining mass is assumed to be ash inorganics, which can be estimated *via* NREL/TP-510-42622.<sup>87,89</sup> Elemental determination of ash content is not necessary unless there is indication that it might affect the reaction chemistry; above 10% ash content in the extracted sample may cause problems with hydrolysis. For example, palm wood and wastes may contain significant levels of iron<sup>102</sup> that may interfere with catalysis and certainly makes NMR analysis challenging.<sup>103</sup> In such an instance, atomic emission or absorption spectroscopy can be used for determining the element of interest.<sup>104,105</sup>

#### Minimum reporting requirements for feedstock sourcing and preparation:

- Origin and species of the feedstock.
- Biomass particle size used for reaction studies.
- Moisture, extractives, and ash content, and the methodology used to determine them.

#### Preferred reporting recommendations for feedstock sourcing and preparation:

- Part of the feedstock used, growth location, and harvest parameters.
- Procedures for drying, sizing, and storing (and duration).

### 2.2 Compositional analysis

Reliable standard methods are available for identifying and quantifying the carbohydrate constituents of biomass, described in detail in NREL/TP-510-42618.<sup>89</sup> Even when the focus is placed on lignin deconstruction, quantifying the carbohydrate fractions is essential for the calculation of overall mass balances and yields to specific components. Furthermore, preserving and/or valorising the carbohydrate fraction is vital to make any lignin-first biorefinery economical. In these techniques, biomass is hydrolysed in the presence of a mineral acid in two stages. The first step involves the dissolution of polysaccharides in 72% w/w sulfuric acid at 30 °C, which is analogous to the first step of a Klason lignin determination.<sup>106</sup> Therefore, it is convenient to perform both analyses simultaneously. The dissolved polysaccharides are diluted to form a dilute acid mixture (4%) and hydrolysed to monosaccharides at a higher temperature. At this stage, standards are treated under the same conditions to



account for sugar degradation. The final yields are then corrected for degradation by comparison with the standards. Once the acid-insoluble fraction is filtered out, quantification follows HPLC separation of the monomeric sugars in the resulting liquor. Depending on the source of biomass, different monomers should be used as standards besides D-glucose. For example, D-xylose, D-galactose, D-mannose, and L-arabinose are used as standards to quantify hemicellulosic sugars in the case of most hardwoods and grasses. Alternatively, GC can be used. In this case, the sugars are first reduced and then acetylated to their alditol acetates.<sup>107</sup> Here, the monosaccharide standards also need to undergo this pretreatment. For both HPLC and GC data processing, quantification can be performed by analysing the resulting sugar peaks using a calibration curve built using a dilution series with external standards. NREL/TP-510-42618 contains detailed quality control and error analysis methods.<sup>89</sup>

The quantified and corrected values can then be used to calculate the various polysaccharide fractions. The measured glucose is largely from biomass' cellulose fraction, but a fraction also derives from hemicelluloses; xylose, galactose, arabinose, and mannose are assumed to have been produced from hemicelluloses. Researchers should also consider free sugars and starch, if applicable. When calculating the original mass of these simple sugars in the native biomass, the water molecule added during hydrolysis has to be taken into account such that, rather than the monosaccharides themselves (glucose and xylose, for example) their forms in the polymer (glucan, *i.e.* (C<sub>6</sub>H<sub>10</sub>O<sub>5</sub>)<sub>n</sub>; and xylan, (C<sub>5</sub>H<sub>8</sub>O<sub>4</sub>)<sub>n</sub>) are used.

The percent by mass content of cellulose and hemicelluloses in the original biomass sample is calculated as follows:

$$\% \text{Cellulose} = \frac{m_{\text{glucan}}}{m_{\text{substrate,dry}}} \times 100\% \quad (1)$$

$$\% \text{Hemicelluloses} = \frac{m_{\text{xylan}} + m_{\text{galactan}} + m_{\text{arabanan}} + m_{\text{mannan}}}{m_{\text{substrate,dry}}} \times 100\% \quad (2)$$

The total polysaccharide content is taken as the sum of cellulose and hemicelluloses (in which the latter may also include acetate). The quantification procedure should always be reported and referenced even for the general procedure described. All deviations from established protocols must be reported.

#### Minimum reporting requirements for structural carbohydrate quantification:

- Fraction of major structural carbohydrates (cellulose and hemicelluloses) of the feedstock biomass based on at least three replicates with the resulting standard deviation.

### 2.3 Lignin characterisation

**Lignin content.** The Klason lignin (acid-insoluble lignin, AIL, and acid-soluble lignin, ASL) determination method,

described in NREL/TP-510-42618,<sup>89</sup> provides reliable lignin quantification for most traditional biomass samples that have been pre-extracted with water and solvents (*vide supra*)<sup>94,108</sup> to be free of extractives. This extractive-free material has been termed 'cell wall residue' (CWR), and in more recent alcohol-based extraction methods, may be referred to as an 'alcohol-insoluble residue', that essentially represents the plant cell wall fraction.<sup>108-110</sup> The basic procedure involves dissolving polysaccharides and acid-soluble lignin using 72% w/w sulphuric acid. After dilution, hydrolysis, and filtration, the remaining solids are considered to be acid-insoluble lignin and ash. Ash is defined as the remaining solids after a subsequent calcination of all remaining solids.<sup>87</sup> Although the lignin structure undergoes significant transformation during this process, the mass that is determined is thought to accurately represent lignin in wood.

It is noteworthy that the Klason lignin method is considerably more problematic for materials with bark and leaves, and for grasses and legumes that may contain proteins, suberins, and other complex components not found in extracted wood.<sup>101</sup> The source of the non-lignin components in 'Klason lignin' has only occasionally been delineated; such is the case for cereal grains and other plant-based foods in which the limitations of "non-specific lignin determination methods"<sup>111</sup> was recognised.<sup>111,112</sup> A quote from an abstract notes the extent of the problem: "Estimation of the contribution of non-lignin compounds to the Klason lignin contents reduced the non-corrected Klason lignin contents of the insoluble fibres from 28.7% (kale), 22.8% (pear), 14.8% (wheat), and 9.9% (corn) to maximum lignin contents of 6.5% (kale), 16.4% (pear), 4.9% (wheat), and 2.3% (corn)."<sup>112</sup> A seedcoat material containing an intriguing C-lignin (derived from the novel monomer, caffeyl alcohol) was originally reported to be 90% lignin from Klason lignin determination,<sup>113</sup> but was later revised down to only 10% lignin in a subsequent study,<sup>28,114</sup> which used methods that are unfortunately not viable for widespread corrections of Klason values. For now, Bunzel's methods appear to be the best for elucidating interfering components derived from fats, waxes, cutin, and suberin.<sup>112</sup> Other methods exist for lignin determination, but all have their own limitations.<sup>101</sup> This long-term unsatisfactory situation makes it difficult to make strong recommendations here except to note that Klason lignin remains the gold standard for woody biomass,<sup>89</sup> and may also be reasonable for forage grasses and legumes.<sup>101,115</sup>

**S : G : H ratio.** Measurement of the ratio of S, G, and H units within lignin is important because of the ramifications for depolymerisation and the value of the products. Having two, one, or zero methoxyls *ortho* to the phenolic group (that may be etherified in the polymer) affects more than just the electronic characteristics of the S, G, and H aromatics. Guaiacyl units are typically more condensed (*i.e.*, have more C-C links with other lignin units) than S units because the availability of the aromatic C5 position for radical coupling leads to their being involved in 5-β-, 5-O-4-, and 5-5-coupled units that have no S counterparts;<sup>116</sup> obviously S units may still be involved in



4-O-5-coupled units (with the G-unit linked at C5). S units are therefore more heavily involved in the  $\beta$ -ether units that are the weaker bonds that are cleaved in essentially all of the depolymerisation methods, including those used for analytics. S units are, however, also more heavily involved in (condensed)  $\beta$ - $\beta$  units, because of the long lifetime and favourable dimerisation of the sinapyl alcohol radical during lignification.<sup>116</sup> Incidentally, 4-O-5 and 5-5 units (Fig. 1) are difficult to quantify, and may not contribute to S:G:H measurements, depending on the method. They are important because they represent connections formed during lignification from two growing polymer chains.<sup>8,116</sup> However, these were always thought to produce real 'Y-type' branchpoints in the polymer chain but, to date, evidence only for free-phenolic units can be obtained, meaning that the lignin polymer chains may be less branched than previously thought.<sup>6-8,117</sup>

Other factors can influence bond speciation in lignin, which is an active area of research.<sup>118,119</sup> We comment only briefly on H units because they are minor and may often be neglected, except in targeted transgenics, and in softwood compression-wood zones in which the level may reach some 30%.<sup>120</sup> In principle, H-units have even more options for radical coupling than G and S units because of the additional open C3 position on the ring but, although H units may be involved in more extensive 5/3 bonding, the longstanding assumption that H-lignins are substantially more condensed is not borne out experimentally –  $\beta$ -ether units still predominate even in H-rich lignins or H-only synthetic lignins.<sup>121</sup> Unfortunately, due to conflation from other units that are not polymer units derived from *p*-coumaryl alcohol,<sup>122-124</sup> H units are often reported at a far higher level than they actually are in lignin (*vide infra*). Care in analyses must therefore be taken to exclude compounds arising from components in lignin that are not derived from the prototypical monolignols. Examples include the vinylphenol and vinylguaiaicol that efficiently arise from abundant *p*-coumarate and ferulate esters in grasses as analysed by analytical pyrolysis followed by mass spectrometric detection such as GC/MS or Molecular Beam Mass Spectrometry (MBMS).<sup>125-127</sup> The common refrain that softwoods are G-lignins (with low H levels), hardwoods are S/G-lignins (with very low H levels), but grasses are H/G/S lignins largely results from this mischaracterisation; grass lignins rarely contain more than 5% H units.

We illustrate below how to obtain (more) reliable values but acknowledge that none of the methods listed below is capable of determining the actual distribution in the polymer. Although the NMR methods can in principle, limitations that are discussed below persist in practice. All of the degradative methods release only a fraction of the polymeric units for quantification. We will not discuss secondary spectroscopic methods such as FT-IR, Raman, or NIR because they are all compromised by their reliance on the other methods here for their calibration or have insufficient 'peak purity' to allow single- or multiple-peak direct quantification at this point.

**Solution-state NMR methods for S:G:H determination.** As (solution-state) NMR can, in principle, measure the signals

from the entire lignin, NMR might be seen as the only method that can determine the relative levels in the entire lignin polymer. There is a long history of 1D proton (<sup>1</sup>H) and carbon (<sup>13</sup>C) methods used for (attempted) lignin quantification, as reviewed.<sup>128</sup> The major problem is one of insufficient dispersion of assignable resonances that can be reliably used. Two-dimensional (2D) NMR methods, particularly the <sup>1</sup>H-<sup>13</sup>C correlation experiments such as HSQC (heteronuclear single-quantum coherence), largely solve the dispersion problem; correlation contours (peaks) for S, G, and H units are strikingly well dispersed.<sup>128</sup> A major breakthrough was the ability to run solution-state NMR spectra on whole-cell-wall material (and, essentially, whole biomass) by dissolution or swelling of the finely divided (ball-milled) material in a good lignin solvent.<sup>128-131</sup> Even in HSQC spectra, a few overlaps can still occur; for example, the C2/H2 and C6/H6 peaks from minor 4-O-5-linked G units may coincide with (redundant) C2/H2 and C6/H6 peaks from normal S units.<sup>7,117</sup> H unit determination is made particularly difficult by the coincidence with a phenylalanine peak from proteins that are often associated with biomass samples; protease treatment of the sample before NMR can help alleviate this issue and provide more realistic estimates of H levels.<sup>123</sup>

The peak contours can be volume-integrated to provide comparative S:G:H estimates based on peak ratios. As far as anyone can discern (as there is no method to provide independent and accurate data), even when HSQC spectra are acquired under qualitative conditions, the ratios are reliable. Recent advances such as the more energy-efficient adiabatic-pulse variants of HSQC-type experiments are recommended as standard experiments on newer high-field instruments, particularly those equipped with cryogenically cooled probes. They offer the advantages of a wide inversion bandwidth meaning minimized <sup>13</sup>C-pulse offset effects, lower power, and a wider decoupling range using various decoupling schemes, as reviewed in the context of lignin spectra.<sup>128</sup> They are also less sensitive to spin-spin coupling effects, for which there are other solutions.<sup>132-134</sup> To obtain S : G : H ratios, it is strongly recommended to volume-integrate only the well-dispersed S2/6, G2, and H2/6 (which is unfortunately compromised by coincidence with phenylalanine) because these all have similar coupling environments – all are coupled only to the 6-proton with a small  $J_{H-H}$  of 1.6–2.0 Hz, all have a similar  $J_{C-H}$  coupling constant, and all are in the same small region of the spectrum.<sup>128</sup>

Extrapolating the reliability of qualitative HSQC for S:G:H quantitation to other units or structures of interest is not, however, sound. Regular HSQC data are not quantitative. For example, it is well known that mobile end-units, with their longer relaxation times, become over-represented *vs.* units buried within the lignin backbone because the fast relaxation of the latter units during the actual NMR pulse sequence (and therefore after excitation but before the actual signal detection) is significant.<sup>128,131</sup> Mobile, pendent units such as *p*-coumarates (in grasses) and *p*-hydroxybenzoates (in poplar, aspen, palms) adorning some lignins (but that should not be quantified as actual lignin units) are therefore seriously overestimated by such



HSQC methods. As this longer relaxation giving higher signals is contrary to expectations for researchers familiar with 1D NMR, it is worth a simple illustration of why this is so. In such sophisticated NMR experiments, the length of the pulse sequence applied following the initial excitation of the proton signals, prior to the signal acquisition, is a non-trivial fraction of the relaxation time. If a long-relaxing end group's resonance has decayed to 90% of its original level before the start of the acquisition, but an internal unit's rapidly-relaxing nuclei have decayed to 10%, the former will be over-represented relative to the latter by a factor of 9.

Although current HSQC NMR methods must be used cautiously to quantitatively analyse whole cell wall samples, isolated lignins are more soluble and have sufficiently improved relaxation properties to allow utilisation of a quantitative method such as HSQC<sub>0</sub>.<sup>135</sup> In this experiment, three 2D spectra with different starting points are acquired, the peak volumes are measured, and these are projected back to a theoretical time-zero (before relaxation has occurred). It works well in some cases, as illustrated recently,<sup>136</sup> but we stress that quantification remains non-trivial. The recommendation for original biomass, therefore, is to use the described qualitative HSQC methods<sup>130,131</sup> on whole-cell-wall (whole biomass) or lignin samples for reasonable S:G:H estimates. If accurate values are needed, the lignin must be isolated for the use of HSQC<sub>0</sub> methods.<sup>135,136</sup> The most straightforward way is to produce a so-called enzyme lignin (EL) by digesting away most of the polysaccharides while retaining essentially the entire lignin component.<sup>123,137</sup> If keeping all structures in their native form is not considered crucial, a more rapid and convenient method that provides a large fraction of the lignin in rather clean form [but with spirodienones ( $\beta$ -1-linked units) hydrolysed to their open form, for example] is a mild acidolysis method.<sup>138,139</sup> Solution-state NMR is indispensable for validating and profiling the incorporation of non-canonical monomers, including those from beyond the monolignol biosynthetic pathway (such as tricin and hydroxystilbenes) into the lignin polymer.<sup>10,126,128,140–142</sup> Modern solid-state NMR (being currently used more for understanding polysaccharide interactions),<sup>143,144</sup> and Dynamic Nuclear Polarisation-enhanced methods,<sup>145</sup> are producing exciting new insights and are worth monitoring in the quantification space.

**Degradative methods for S:G:H determination.** The primary degradative methods for S:G:H determination are: nitrobenzene oxidation, thioacidolysis, derivatisation followed by reductive cleavage (DFRC), RCF, and analytical pyrolysis. All of these methods deliver the ratio derived only from the releasable (and quantifiable) monomers and therefore do not represent the entire lignin; arguments can be made that the values are or are not representative. Dimers may also be analysed but these are usually used to identify resistant structural units and are typically not included in the S:G:H determination. Dimer fractions are particularly quantitatively distorted, with  $\beta$ -1 units that are known to comprise only a percent or two of the polymer, accounting for 30% or more of the dimers.<sup>146,147</sup> The information obtained from these methods is subtly or

overtly different. Two (thioacidolysis and DFRC) are considered most diagnostic for lignin because they operate on the principle of cleaving  $\beta$ -ethers to generate monomers, and, importantly, leave a signature to verify that an ether has been cleaved.<sup>148–150</sup> These are the methods that must be used to distinguish real lignin units from other structures associated with the cell wall that may produce the same monomers, a particular problem with nitrobenzene oxidation. Some of the features of each method, with the exception of analytical pyrolysis which has been recently reviewed elsewhere,<sup>127</sup> are noted here:

- Nitrobenzene oxidation produces the highest monomer yields as certain C–C-bonded structures may cleave to produce a monomer.<sup>151</sup> Micro-methods have been developed to improve both throughput and safety,<sup>152</sup> as nitrobenzene explosions are well-known. The product hydroxybenzaldehydes and hydroxybenzoic acids are all available as standard compounds so they are easily quantified. Researchers need to be aware that certain lignins (poplar, aspen, willow, and palms) contain *p*-hydroxybenzoate units on their lignins, and grasses (and all comelinid monocots) similarly contain *p*-coumarates that will produce the same monomers. Conflating all these molecular species into a single H-level number elevates the apparent H-levels in a way that does not reflect the true lignin composition; methods (*vide infra*) that do not produce the same products as lignin should be used in such cases. Grass cell walls also contain ferulates, mostly on arabinoxylan polysaccharides, and their oxidation to the same monomers may also artificially inflate G-levels.

- RCF conditions have been used to explore lignin structure since the 1930s,<sup>51,153</sup> and may yet become a preferred method;<sup>24,42,57,60,62,118,154–159</sup> comparisons have already been made with thioacidolysis.<sup>118,160</sup> The reactions are simple, need little optimising, and produce a modest array of simple products largely retaining their H, G, and S signatures, products for which standards may be available (for authentication and as quantification standards). Catalyst choice (*e.g.*, Ru *vs.* Pd) allows selectivity for primarily arylpropanes *vs.* arylpropanols.<sup>24,28,59,61,80,81</sup> The only complication is that 4-O-5-coupled units may partially cleave to contribute (in only a small way) to the monomers, but also produce novel 'rearrangement monomers' and dimers diagnostic of their origin.<sup>161</sup> Care should be taken when using RCF for analytical approaches to avoid demethoxylation reactions, which would skew the resulting S:G:H ratio. We stress that RCF is not yet a fully established analytical method and note that opportunities exist to develop this approach further.

- Thioacidolysis is perhaps today's premier diagnostic method for characterising lignin, releasing *p*-hydroxyphenyl-trithioethyl-propyl monomers solely by cleaving  $\beta$ -ethers.<sup>150,162,163</sup> Again, small-scale methods are available.<sup>164,165</sup> Unfortunately, the standards are not available and need to be synthesised.<sup>166</sup> The desulphurisation method that produces the arylpropanes is not generally used for monomers analysis, but is used for dimers characterisation.<sup>147,150,162,167</sup> In addition to the foul smell of the reagents, thioacidolysis has one shortcoming – many units



in lignins are acylated (primarily by acetate, *p*-hydroxybenzoate, *p*-coumarate, or ferulate) but these are neither fully retained nor fully cleaved during the procedure, resulting in some distortion of values.<sup>168,169</sup>

• DFRC was invented to possibly circumvent the noxious odour from thioacidolysis by using a different mechanism, reductive cleavage, to specifically cleave  $\beta$ -ethers.<sup>149,170</sup> In fact, thioacidolysis generally produces higher monomer yields, but DFRC is particularly useful for certain determinations. Like thioacidolysis, it leaves a diagnostic signature that a  $\beta$ -ether has been cleaved, specifically the double bond in the product monolignol acetates. It is somewhat attractive that the products are the monomers (although as their peracetates) from which the lignin was originally biosynthesised in the plant. Its huge advantage or disadvantage, depending on the level of information required, is that it absolutely does not cleave esters in the acylated monolignols noted above.<sup>169,171</sup> That means that any unit that is acylated, except by acetate, will not yield its H, G, or S monomer, but will instead be released as a monolignol conjugate. Quantification therefore needs to either be measured with the normal and acylated monomers summed, or the esters should be first cleaved in a separate step (below). The DFRC procedure for cleaving ethers but leaving esters intact has become a powerful tool for studying such conjugates in lignin.<sup>149</sup>

In principle, it is straightforward to assure that acylated lignins do not pose a problem for nitrobenzene oxidation, thioacidolysis, DFRC, or RCF. Adding a saponification (and extraction) step prior to the procedure, removes these esters. It is not completely straightforward, however, and reliable protocols do not appear to have been developed to date.

#### Minimum reporting requirements for lignin characterisation:

- Klason lignin content based on at least three replicates along with the resulting standard deviation.
- Based on carbohydrate and lignin analysis, the overall mass balance of the substrate (lignin, cellulose hemicelluloses, ash, extractives, protein, etc.).

#### Preferred reporting recommendations for lignin characterisation:

- S : G : H ratios determined by thioacidolysis (or *via* DFRC, RCF, nitrobenzene oxidation, or analytical pyrolysis).
- 2D NMR volume-integrals for S:G:H from whole-cell-wall or isolated lignin samples.

## 3. Reactor design for lignin-first biorefining

For any chemical transformation, selecting the appropriate type of reactor is crucial for the overall process design. Of all process equipment, reactor design requires consideration of rate constants, reaction enthalpies, heat and mass transfer coefficients, and phase equilibria data. In addition, other factors, such as process economics, scale-up, and safety of operation, influence this choice.<sup>172</sup>

The two primary reactor configurations reported to date in lignin-first biorefining are batch and flow-through (Fig. 3). Although both reactor designs can be used to extract experimental data, it is imperative to understand the benefits and limitations of each configuration when delineating experimental goals. In this section, we describe innate advantages and common pitfalls for each reactor type and present guidelines for reactor selection, operation, and reporting. We provide recommendations and best practices for data collection, including heuristics for identifying heat and mass transfer limitations. We note that lignin-first processes often require high temperature and pressure; thus, operators must receive proper training and adhere to strict safety protocols and standards in the construction and utilization of chemical reactors.

### 3.1 Batch reactors

A stirred autoclave vessel, commonly known as a batch reactor, has been the typical reactor used to investigate lignin-first fractionation processes in the condensed phase (Fig. 3a). In this type of reactor, the reagents, solvents, and catalysts are sealed and heated for a predetermined amount of time and can feature intermittent sampling to track reaction progress. Batch reactors are simple to operate and are used commercially to produce low-volume, high-value chemicals. To date, most biomass processing in the pulping and bioethanol industries is performed in batch or semi-batch mode. Understanding both the inherent practical advantages and the limitations of batch reactors is important to avoid data inconsistencies. The following suggested guidelines can be used to improve reporting and reproducibility across laboratories.

Reactions carried out in batch systems are transient in nature, complicating the collection of reaction rate data, as reactant and product concentrations change as a function of time. Instantaneous rates that are not only specific to a particular set of reaction conditions (*e.g.*, partial pressure, temperature, and initial concentrations), but also to the extent of reaction, are not readily available. Initial rate data can be obtained, for example, by calculating the slope of regressed datapoints collected in the near-linear, low-conversion regime (typically <10%) in a conversion as a function of time plot. Data fitting may be another way to obtain rate data, but this requires a reliable kinetic and/or mass transfer model that might not be trivial to develop.<sup>119</sup> Reporting individual yields might generate misleading data given that product yields, particularly at high conversion extents, can be almost identical at two different timepoints as reactions rates slow down with reactant depletion or as reactions approach equilibrium. These drawbacks are exacerbated when trying to collect rate data for more complex reaction networks (*e.g.*,  $A \rightarrow B \rightarrow C$ ), which is always the case for lignin-first processes.

Ideally, the reaction should start being timed ( $t = 0$ ) when the reactants are put into contact with the catalyst at the reaction temperature. However, in practice, this is often not possible as reagents and catalysts are mixed together and heated to the reaction temperature with the initial time chosen



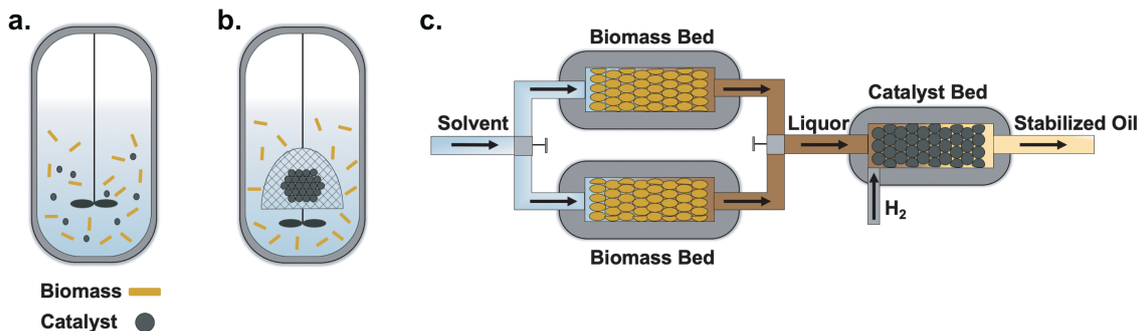


Fig. 3 Reactor configurations reported to date for lignin-first biorefining. (a) Batch reactors wherein whole biomass, catalyst, solvents, and other species (e.g., H<sub>2</sub>) are mixed and reacted. (b) The use of a catalyst basket in a batch reactor system can physically separate the catalyst and biomass particles, thus simplifying post-reaction biomass and catalyst analyses.<sup>66</sup> (c) Flow-through systems are also useful to physically separate the biomass and catalyst during lignin-first processing and to study solvolysis-limiting or hydrogenolysis-limiting reaction conditions. Therein, a heated solvent flows over a switchable biomass bed, and the liberated, lignin-rich liquor passes over a catalyst for depolymerization and stabilisation.<sup>63–65,118</sup> Image credit: Gregory Facas, NREL.

as the point at which heating starts; this heating period can take minutes to tens of minutes, depending on the laboratory setup. Using the starting time when the reactor is at reaction temperature is similarly problematic as reactions already occur earlier in the process during the heat-up period. Adding the biomass later after proper heating seems like a solution, but this is not commonly practical at laboratory scales. Construction materials, vessel volume, use of liners, heating and cooling method, stirring regime, and total mass loaded into the reactor all drastically alter the heating and cooling profiles, thereby introducing large errors particularly at the crucial early reaction times needed to calculate initial rate data. Similarly, although adding the catalyst at reaction temperature may be an option in specialised reactor configurations, depolymerisation and condensation can occur even in the absence of catalyst as a result of solvolysis. Regardless of the batch reactor setup, the heating and cooling profiles for the vessel should be reported.

In typical batch lignin-first processing, the solid biomass is mixed with a solid heterogeneous catalyst, which makes post-reaction biomass and catalyst characterisation a substantial challenge. One solution to overcome this problem includes the use of catalyst baskets in an autoclave (Fig. 3b).<sup>66</sup> Using catalyst baskets and pre-sized catalyst pellets in these reactor types enables practical separation of catalysts and pulp after reaction. Such systems can also be used to investigate the stability of the metal catalysts by renewing the solvent and biomass loaded into the reactor with the same basket. Sizing the catalyst particles is important to prevent catalyst escape into the bulk liquid with the biomass. However, the size cannot be too large or diffusion limitations will occur, hampering catalytic depolymerisation and the required rapid intermediate stabilisation.<sup>66</sup>

Non-uniform stirring caused by vessel geometry or the use of baffles, and/or the type of stirring mechanism (magnetic, mechanical), can cause the formation of dead spots with poor mixing that potentially introduce heat and mass transfer artefacts (*vide infra*). Although in some cases, liquid-phase

experiments can be carried out under static conditions, in most biomass fractionation processes (particularly when dealing with three-phase systems), mixing is an important factor for obtaining reproducible data. Depending on the size of the biomass particles, slow stirring could cause particles to sink to the bottom of the reactor, thus changing reactivity profiles. The same occurs when using baffle systems that are beneficial to increase gas/liquid mass transfer. Similarly, using high biomass-to-solvent ratios, that are required for the economics of lignin-first processing, may result in highly viscous slurries that are difficult to mix without powerful overhead stirrers. For example, although Sels and co-workers were able to use very high biomass to solvent ratios (e.g., 60 g<sub>biomass</sub> in 240 mL of solvent) in batch RCF experiments using an autoclave with overhead motorised stirring,<sup>43</sup> similar conditions would have resulted in less effective mixing in vessels relying on stir bars powered by magnetic stir-plates. For this reason, small reactors, below 50 mL, with similar loadings generally give irreproducible results. For experiments involving three-phase systems (e.g., those involving solid biomass, solid catalyst, solvent, and reactant gas), the use of gas-entrained impellers is ideal to maximise gas-liquid interfacial area. However, caution must be exercised when operating at supercritical conditions, at which solid particles can enter and clog the impeller due to the loss of a significant density difference between the gas and liquid phases needed for gas entrainment.

Many autoclaves are equipped with a sampling port in which a small volume of the slurry (containing the solvent, liquid products, solid biomass, and the catalyst) is isolated from the rest of the reactor for collection. However, if the reaction volume loaded into the reactor is low compared to the volume of the sampling tube, then frequent sampling will change the reaction profile producing non-uniformities in the slurry and changes to the gas-liquid proportions in the vessel; preferably not more than 5–10% volume should be removed from the reactor (in total). As an alternative, several identical reactors can be run in parallel, stopping each one at a different time interval to construct a reaction time profile.



**Minimum reporting requirements for batch reactor use in lignin-first processes:**

- Reagents, catalysts, and solvents and their nominal quantities.
- Vessel geometry, material of construction, reactor liner, stirring method (if used), and stirring rate.
- Heating, cooling, and pressure profiles.
- Quantity and frequency of sampling.
- Definition of the reaction time.

**Preferred reporting recommendations for batch reactor use in lignin-first processes:**

- For kinetic studies, the initial rate data or full reaction time profiles rather than single/final yields.

**Best practices:**

- Total sampling volumes should not exceed 5–10% of the reaction volume.
- Reaction vessels above 50 mL are recommended, especially for high solid loadings (above 10 wt%).
- The use of overhead motorised stirring is recommended especially for high solid loadings (above 10 wt%).
- Use Hastelloy reactors or an inert reactor liner to avoid leaching of reactive metal species from the reactor walls.

### 3.2 Flow-through reactors

Disadvantages with batch reactors include: (1) slow heating, (2) inability to optimise and also study solvolysis and hydrogen-transfer reactions separately, (3) tedious separations of catalyst from the pulp, and (4) mechanical disruption of the pulp. A strategy to separate the solvolytic extraction of the lignin from the metal-catalysed reactions is to separate these two processes in space and time by employing a flow-through reactor (Fig. 3c).<sup>63–65,118</sup> In such a system, the biomass is loaded into a percolation chamber and the metal catalyst is loaded into a downstream reactor. Solvent, which can be pre-heated to the desired reaction temperature, is then pumped through the percolation chamber to extract lignin (and some polysaccharides) and transfer them to the reactor containing the transition metal catalyst bed. In between the percolation chamber and the reactor, a T-coupling with an outlet may be fitted to allow real-time analysis of both solvolysis and hydrogen transfer reactions. A back-pressure regulator stabilises the pressure in the system.

In contrast to batch reactors, flow-through reactors with switchable beds convert substrates in a semi-continuous manner (Fig. 3c) Using this configuration, steady-state can be approximated by switching biomass beds such that the concentration of extracted lignin reaching the catalyst bed remains relatively constant within a range that does not significantly alter reaction rates. In its simplest form, a packed-bed flow reactor consists of a tube packed with a catalyst bed held in place by a frit, mesh screen, or plugs of inert material (*e.g.*, glass beads, quartz wool). Under ideal conditions, all substrate elements flow at the same velocity, parallel to the reactor axis, without back-mixing; plug flow conditions can be measured and reported as the dimensionless Reynolds number, *Re*. This scenario allows the assumption that all material present at any given reactor cross-section has had an identical residence time. Depending on the application, these reactors can be operated either in isothermal (*i.e.*, constant temperature throughout the reactor) or adiabatic (*i.e.*, varying temperature across the

reactor length) modes to process feedstock over a wide range of throughput volumes. The advantages of flow systems were recently demonstrated for RCF of poplar using flow-through setups with separated biomass and catalyst beds to obtain intrinsic kinetic data and identify mass-transfer limiting conditions for the entire process.<sup>64,118</sup> Decoupling the conditions for solvolysis from those of reductive stabilization in flow enabled the interrogation and optimization of crucial aspects of the RCF process that are difficult to access with batch reactors. For instance, operating at the limiting conditions for solvolysis (*i.e.*, when lignin fragment detachment is slow relative to the time scale of the total number of turnovers for reductive bond cleavage at the catalyst surface) allowed the influence of the solvent composition on lignin solubilization to be isolated, while operating at limiting conditions for reductive stabilization (*i.e.*, when lignin solvolysis is fast relative to the time scale of reductive bond cleavage of lignin fragments) allowed catalyst activity and stability to be studied. The following suggested guidelines for flow-through lignin-first reactors can be used to improve reporting and reproducibility across laboratories.

Although turbulent flow regimes are preferred to laminar flows for improved mixing and heat transfer normal to the flow direction, achieving high Reynolds numbers (*Re*) could require excessively high flowrates. Under laminar flow regimes, the flowrate is proportional to the pressure drop, which is a function of bed height, linear flowrate, and dynamic viscosity of the fluid. The pressure drop can increase with bed deformation caused by high flowrates, pellet dissolution, or structural changes in the bed, eventually leading to over-pressurisation and even complete loss of flow through the reactor.

Catalyst beds act as deep-bed filters, capturing and precipitating colloidal material that can cause bed fouling and clogging. Researchers must pay close attention to flowrate/pressure drop effects and always install the appropriate safety measures (guard beds, rupture disks, pressure monitors, as well as emergency over-pressurisation alarms and shutdown valves) to prevent accidental releases in the case of partial or full blockage.

Non-uniformity in the bed packing is a major culprit for flow non-idealities and should be evaluated when working with solid biomass substrates. For example, channelling (in which fluid (solvent or gas) passes through one part of the reactor bed more rapidly than other parts) or hold-up (in which a fraction of the substrate resides in stagnant areas with reduced flow) can drastically change reaction profiles. For catalytic beds in microreactors, it is recommended to pelletise the powdered catalyst and sieve it to a size large enough to prevent large pressure drops or add in inert particles of a larger diameter, in both cases to avoid external mass-transfer limitations (*vide infra*). A common heuristic is to allow space for  $\geq 100$  particle diameters in the axial direction and maintaining at least a 4 : 1 height-to-width aspect ratio of the bed to ensure uniform contact of all substrates with the solids.<sup>173</sup> This is generally accomplished using inert fillers (*e.g.*, quartz chips or silicon carbide particles) having a similar particle size to the sieved



catalyst to reach this ratio for cases in which a lower amount of catalyst is required. Similarly, for the biomass beds, the use of inert filler particles may be necessary to avoid bed collapse, especially in cases where high extraction extents occur.<sup>174</sup>

Deviations from ideal plug flow due to back-mixing will cause product streams to have a distribution of residence times. For extremes of back-mixing profiles, the plug flow reactor approaches the behaviour of a continuously stirred reactor (CSTR), potentially limiting expected conversion values – the mathematical description of this phenomenon is beyond the scope of this work and can be found in standard reaction engineering textbooks.<sup>173,175,176</sup> When using fixed biomass beds in flow-through reactors, as in the flow-through RCF studies described above, it is crucial to check the integrity of the bed during solubilisation. Indeed, although some solvents (*e.g.*, methanol, ethanol, and dioxane) might selectively extract lignin from the cell wall, the use of water, acid, or base additives may also remove the carbohydrate fraction and cause severe structural changes to the biomass bed, which can cause bed collapse and plugging. Lastly, we note that changes in residence times in a flow-through reactor may result in different degrees of undesirable reaction products. For example, when performing flow-through RCF in reactors with separated biomass and catalyst beds, the flow profiles and residence time distribution between the beds may cause higher condensation rates compared to batch reactors unless a solvent system is used that temporarily stabilises the intermediates.<sup>177</sup>

In a sealed autoclave, the pressure is established by the vapor-liquid equilibrium at the reaction temperature of the liquid and gas substrates loaded into the reactor, whereas in a flow-through system the pressure is regulated by the user (*e.g.*, by using a back-pressure regulator). If the appropriate pressure is not selected when desiring to work exclusively in the liquid phase at the reaction temperature, the feed may exist in a mixed liquid-vapor or entirely in the vapor phase when flowing through the catalyst bed, generating irreproducible data. For this reason, it is recommended to carry out the appropriate vapor-liquid equilibrium calculations prior to the design of the experimental stage using, for example, tabulated data or a software package such as ASPEN Plus.

### 3.3 Reactor selection for lignin-first studies

Reactor selection and experimental design must be performed in concert to obtain the desired information accurately, with the minimum number of measurements and the least expense. This is particularly important for lignin-first processes, such as RCF, that comprise two independent consecutive steps, each featuring intrinsic parameters that can dominate the apparent behaviour of the overall system. In this section, RCF will be used to illustrate the importance of reactor selection to meet experimental goals.

Depending on reaction conditions, the overall performance of typical lignin-first processes can be limited either by the solvolysis or the stabilisation step. In RCF, solvolysis-limited conditions occur when lignin detachment from the cell wall is slow relative to the timescale of catalytic turnover, whereas the

stabilisation-limited conditions are observed when lignin solvolysis is fast compared to the timescale of reduction.<sup>64</sup> A solvolysis-limiting condition is required when investigating the dynamics of lignin detachment from the cell wall, including, for example, comparisons of product distributions across biomass types, extraction solvents, and additives (*e.g.*, inorganic acids or co-solvents). AAF is almost always operated in solvolysis-limiting conditions, ensuring rapid stabilisation and condensation prevention, whereas for RCF, the limiting conditions depend on the goals of the study. Batch reactors are ideally suited for solvolysis-limited studies because different variables, including substrates, solvents, and temperatures can be screened with ease across a wide range of lignin extraction extents. Using low biomass-to-catalyst ratios is common practice to ensure operation in the solvolysis-limiting regime. For this reason, however, it is also common to observe invariant monomer yields and fully saturated monomer sidechains regardless of catalyst type used. Note that lignin solvolysis from the cell wall can be limited by mass transfer depending on, among other factors, biomass type and particle size.<sup>119</sup> Mass transfer effects must be identified and reported, as it may cause data irreproducibility across experiments and laboratories.

Operation in a reduction-limited regime is necessary to study catalyst activity and stability, as well as to perform comparisons across catalyst types. This is always the case for AAF because stabilisation is temporally and spatially decoupled from downstream catalytic depolymerisation. Typically, intrinsic kinetic data (*e.g.*, activation barriers) are used to assess catalyst performance, including reaction rates and product selectivity, whereas catalyst stability is evaluated from deactivation rate profiles coupled with catalyst characterisation. Extracting these data with batch reactors is cumbersome due to difficulties in measuring initial rates accurately and separating the catalyst from the biomass particles. Flow reactors, on the other hand, are ideally suited to this task. Having precise control over residence time allows collection of steady-state data under differential conditions, which is necessary to calculate initial rates and to identify primary *vs.* secondary products in complex reaction networks. Similarly, catalyst stability studies are readily accessible by measuring deactivation profiles at steady-state intermediate conversion levels and performing characterization studies post-reaction (or even under *operando* conditions) on the catalyst bed that is physically separated from the biomass bed. Section 3.4 outlines several strategies to check for the presence of transport artefacts in experimental data.

### 3.4 Transport effects

The rate of diffusion of reactive substrates across phases can alter the observed rate of chemical reactions. Similarly, the rate of heat transport into (for endothermic reactions) or away from (for exothermic reactions) a catalyst can influence the temperature-dependent rate constants for the system. To properly evaluate or compare lignin-first methods operated either in solvolysis- or reduction-limited regimes (*vide supra*), intrinsic kinetic data that are free from heat and mass transport artefacts must be measured.<sup>119</sup> If the observed rate is modified by



these artefacts, then new conditions must be selected to either lower the reaction rates or increase the rate of heat/mass transfer. Rigorous calculations of reaction-diffusion profiles to extract intrinsic parameters, (*e.g.*, Thiele moduli and effectiveness factors) require numerical techniques and high-fidelity models, as recently shown by Thornburg and co-workers for lignin-first RCF.<sup>119</sup> In this section, we show simplified methods to assess the influence of mass transfer limitations in an approximate manner.

Gas-liquid diffusion is important when consuming gaseous reagents in a liquid solvent. For example, the transport of hydrogen gas across the interface between the headspace and the agitated solvent in a batch reactor performing RCF could limit the apparent rate if the catalyst is starved of hydrogen. It is recommended to always perform a “maximum rate of gas transfer” analysis on the reactor that will be used for the lignin fractionation studies under identical conditions of liquid volume, gas pressure, temperature, and agitation to those used in the actual fractionation experiment. This measurement is particularly important when working with systems featuring very fast apparent rates (typically those exceeding 1 turnover per site per second). Several methods exist to evaluate the maximum rate of gas transfer. For hydrogenation reactions using molecular H<sub>2</sub>, Meille *et al.*<sup>178</sup> and Chaudhari *et al.*<sup>179</sup> outlined experiments using either physical adsorption or reaction of a substrate with well-established kinetics (*e.g.*, styrene or  $\alpha$ -methylstyrene) to evaluate the gas-to-liquid mass transfer coefficients. A comparison of the experimentally determined maximum rate of transfer coefficients to those observed under catalytic conditions will indicate if the catalyst will be gas-starved. Note that if the system is gas-liquid mass transfer limited, increasing the stirring rate alone may not be able to overcome this limitation.

The transport of liquid substrates and dissolved gases to the external surface of the solid catalyst pellet can also disrupt intrinsic reaction rates. As the substrates approach the catalyst pellet, they must cross a stagnant boundary layer in which the flux across this layer will be a function of the concentration difference between the bulk phase and the external catalyst surface and the mass transfer coefficient. This phenomenon is often modelled by assuming a linear concentration profile across these two regions by assuming that no reactions take place at the boundary layer. A simple mathematical analysis combined with engineering correlations (not shown here) generates the following relationship between mass transfer coefficient and important process variables when flowing a fluid around a spherical catalyst particle:<sup>173</sup>

$$k_c \propto \frac{D_{AB}^{2/3} \rho^{1/6} u^{1/2}}{R_p^{1/2} \mu^{1/6}} \quad (3)$$

where  $k_c$  is the mass transfer coefficient,  $D_{AB}$  is the diffusivity of species A in the bulk fluid B,  $\rho$  is the fluid density,  $u$  is the linear fluid velocity,  $R_p$  is the radius of the catalyst particle, and  $\mu$  is the fluid viscosity. This equation shows why either increasing fluid velocity (*e.g.*, by increasing stirring speed) or decreasing particle size are commonly used practices to check

for mass transfer effects from the surrounding fluid to the particle surface (*i.e.*, external mass transfer), as both can be readily changed to influence the value of  $k_c$ . Thus, if the reaction rate profile does not change by increasing fluid velocity or decreasing particle size, it is generally assumed that the system is not externally mass-transfer limited. Care must be taken when increasing fluid speed in a flow reactor, as it may result in excessively high pressure-drop values.

If  $k_c$  is known, then a simple criterion to evaluate if external mass transfer does not affect the rate is as follows:

$$\frac{|r_{\text{obs}}| R_p}{k_c C_{AB}} < \frac{0.15}{n} \quad (4)$$

where  $|r_{\text{obs}}|$  is the experimentally observed rate on a volume basis (*e.g.*, moles of monomer production per bed volume per time),  $C_{AB}$  is the concentration of substrate A in the bulk fluid B (*e.g.*, the Klason lignin mass in the loaded content divided by the total solvent used), and  $n$  is the reaction order.

#### Minimum reporting requirements for flow-through reactor use in lignin-first processes:

- Reagents, catalysts, and solvents and their quantities.
- Detailed diagram of the system (including bed aspect ratio, volume of reaction system, thermocouple location, and flow configurations).
- Flowrates of liquids and gases.
- System pressure and temperature.
- Catalyst pellet size.
- Type of solid diluent (*e.g.*, silicon carbide).

#### Preferred reporting recommendations for flow-through reactor use in lignin-first processes:

- Description of reactor packing profile.
- Description of product collection and quenching system.
- Description of start-up phase, biomass and catalyst pretreatment.

#### Best practices for flow-through reactor use in lignin-first processes:

- Pressure in the flow system should be monitored actively during operation.
- Pressure drop measurements across the catalyst bed reactor must be performed to identify appropriate fluid flow and mixing regimes.
- Installation of pressure relief valves to avoid over-pressure from plugging (especially during bed compaction phase) is strongly recommended.
- The use of check-valves to mitigate back-flow issues during plugging is recommended.
- Back-pressure regulators to ensure that pressure is controlled and maintained are recommended.
- Appropriate catalyst bed packing procedures should be followed to avoid data discrepancies (*e.g.*, at least 10 particles in the radial and 50 particles in the axial direction to prevent stochastic effects).
- The use of a flow profile that approaches an ideal plug flow, which requires turbulent flow.

Once substrates reach the surface of the catalyst, they need to diffuse through the internal pore structure of the catalyst to reach the active sites. Depending on the size of the pores and the intrinsic rate of reaction, the observed rate might be internal-transport-limited if the substrates are consumed before reaching all the active sites within the particle. The non-linear analysis coupling the simultaneous diffusion and reaction events required to evaluate the extent of catalyst pellet utilisation (*i.e.*, the effectiveness factor) is complex. Fortunately, Weisz and Prater developed a simple criterion based on



measurable quantities to determine whether the observed rate is not influenced by internal mass transfer artefacts.<sup>180</sup>

$$\frac{|r_{\text{obs}}| R_{\text{P}}^2}{D^{\text{eff}} C_{\text{AS}}} < \frac{1}{n} \quad (5)$$

where  $|r_{\text{obs}}|$  is the experimentally observed rate on a volume basis (e.g., moles of monomer production per bed volume per time),  $R_{\text{P}}$  is the radius of the catalyst particle,  $D^{\text{eff}}$  is the effective diffusivity (defined as the diffusivity adjusted for both porosity and tortuosity of the catalyst, which together decrease diffusivity by approximately a factor of 10) and  $C_{\text{AS}}$  is the concentration of substrate A at the external surface of the catalyst S (which can be set equal to  $C_{\text{AB}}$  if the criterion outlined in eqn (6) is satisfied).

For highly exothermic or endothermic reactions, the temperature within a catalyst pellet might be very different from that measured by the thermocouple in the bulk phase if heat cannot be transported quickly enough within the boundary layer around the pellet (external limitation) or within the pellet (internal limitation). Fortunately, an external criterion developed by Mears (eqn (6)) and an internal criterion developed by Anderson (eqn (7)) can be used to evaluate whether heat transfer effects influence the observed reaction rate as follows:<sup>181,182</sup>

$$\frac{|\Delta H_{\text{R}} r_{\text{obs}}| R_{\text{P}}}{h_{\text{t}} T_{\text{B}}} < 0.15 \frac{R_{\text{g}} T_{\text{B}}}{E_{\text{a}}} \quad (6)$$

$$\frac{|\Delta H_{\text{R}} r_{\text{obs}}| (R_{\text{P}})^2}{\lambda T_{\text{S}}} < 0.75 \frac{R_{\text{g}} T_{\text{S}}}{E_{\text{a}}} \quad (7)$$

where  $\Delta H_{\text{R}}$  is the heat of reaction,  $h_{\text{t}}$  is the heat transfer coefficient (estimated from correlations using the Nusselt number, thermal conductivity, and the particle radius),  $T_{\text{B}}$  is the bulk fluid temperature,  $T_{\text{S}}$  is the fluid temperature at the surface of the particle (which can be set equal to  $T_{\text{B}}$  if the criterion outlined in eqn (6) is satisfied),  $R_{\text{g}}$  is the ideal gas constant,  $\lambda$  is the heat conductivity within the particle (which can usually be approximated by the solvent's conductivity), and  $E_{\text{a}}$  is the true activation energy.

Diagnosing transport effects with eqn (3)–(7) above requires prior knowledge of several critical parameters, such as effective diffusivity and thermal conductivity, that may not be straightforward to obtain for a given system. For this reason, a simple, yet powerful, experiment to verify the presence of both heat and mass transfer limitations is known as the Madon-Boudart test<sup>183</sup> (also known as the Koros-Nowak criterion). This test involves measuring rates on catalysts in which the concentration of active material is changed deliberately, comparing both concentrations at two different temperatures. In the absence of heat and mass transfer effects, the reaction rate is proportional to the concentration of active material (*i.e.*, the TOF should be invariant) and this proportionality should hold at different temperatures. A common way of varying the concentration of the active material is simply by diluting the bed with an inert material while keeping the total bed volume constant, making sure that the inert packing has the same diffusional characteristics and comparable particle size to that of the active catalyst. This way, in the absence of transport effects, reducing the active material concentration by half, should result in a 50% reduction in rates. Importantly,

this test does not require measuring the number of active sites in the catalyst. Examples applying the Madon-Boudart test for liquid-phase and three-phase systems can be found in the literature.<sup>184,185</sup>

#### Minimum reporting requirements for assessing transport effects for lignin-first processes:

- Assessment of mass and heat transfer limitations *via* either of the following criteria:
  - The Weisz-Prater, Mears, and Anderson criteria which should be applied for both external and internal heat and mass transport.
  - The Madon-Boudart test at two different temperatures.
- Verification that the gas loading or gas flows are not limiting unless so intended.

#### Best practices for assessing transport effects for lignin-first processes:

- Rule out gas-liquid mass transfer limitations by evaluating the maximum rate of gas transfer to the liquid.

## 4. Catalyst performance in lignin-first fractionation of biomass

The lignin-first process comprises two catalytic operations: (1) the first is solvolysis, which may be catalysed (usually by an acid), leading to fractionation of the biomass *via* liberation of lignin and varying amounts of polysaccharides. This initial step can occur in the presence or absence of stabilising reagents. However, in their absence, the resulting intermediates will eventually undergo condensation. Subsequently, (2) catalytic reactions occur that stabilise reactive intermediates and/or cleavage of ether bonds. These processes can occur simultaneously as is the case for RCF, or subsequently when stabilisation/capping agents are used. Each of these processes needs to be adequately characterised.

### 4.1 Catalysed solvolysis

The solvolysis step is crucial in lignin-first methods and can be promoted by a catalyst. Therefore, reporting solvent mixture, reaction concentration, temperature, and amount of (generally acid) catalyst or stabilising reagent is necessary. In addition, both solvents and stabilisation reagents can degrade and or be further transformed during subsequent or simultaneous hydrogenolysis. Therefore, although not commonly reported, we recommend that the field moves toward reporting recovery of solvent mixtures, including the catalyst and stabilisation reagents.

### 4.2 Catalytic depolymerisation and intermediate product stabilization

Lignin-first processes are usually conducted on feedstocks with inorganics and other impurities that can be detrimental to transition metal catalysts due to poisoning reactions. These impurities are usually present in significant quantities when the catalyst is in the presence of whole biomass, as is the case with RCF, and are somewhat reduced when in the presence of isolated lignin such as those extracted in processes such



as AAF. The high temperature solvent at high pressure, present during hydrogenation/hydrogenolysis can also facilitate catalyst deactivation including *via* leaching, sintering, and strong adsorption of heavy degradation products.<sup>65</sup> Stirring is required for batch lignin-first processes, whether solid biomass or soluble lignin is used, to limit external mass transfer limitations. Under these conditions, the mechanical force might cause catalyst attrition and result in a loss in performance. Therefore, the thermal, catalytic, and mechanical stability of the catalyst needs to be considered. Ultimately, catalyst regeneration will become an important consideration as the lignin-first field moves toward integration and scale-up, as these processes lead to both reversible and irreversible deactivation.

When no stabilisation agents are used, the solubilised lignin is unstable at high temperatures and must be rapidly brought into contact with the catalyst. In batch RCF processes, this implies having the catalyst in close proximity with solid biomass, after which separation and recycling of the solid catalyst from the solid residue is a challenge. Although using granular catalyst in a basket<sup>66</sup> can solve the downstream separation challenge, the mechanical stirring can lead to attrition and mass loss in the form of fine particles. These fine powders are mixed with the solid residue or lost in the filtration. Alternatively, some catalysts (such as RANEY<sup>®</sup> Ni) can be magnetically separated from solid polysaccharide-rich residues.<sup>40,46</sup> The chemical and thermal stability of a catalyst determines its lifetime in industrial reactors and thus its recyclability is an important factor in determining process feasibility. The stability and recyclability of the heterogeneous catalyst should therefore be considered for such batch reactions. However, when recyclability studies are undertaken, care should be taken to demonstrate recyclability with subsequent batch runs below full conversion, otherwise a catalyst might appear artificially to be stable due to its deactivation not being sufficient to drop the reaction below maximum conversion/yield. For reactions in flow-through conditions, runs at less than maximal conversion/yield should be compared to ensure that catalyst deactivation can be measured. To ensure that recyclability is assessed in a statistically significant manner, we recommend at least three runs, specifying any regeneration operation, to assess the stability of catalyst both in batch and flow-through operation. Additionally, pre- and post-reaction catalyst characterisations (*e.g.*, with microscopy, X-ray photoelectron spectroscopy, inductively-coupled plasma/mass spectrometry, chemisorption, and physisorption) should be conducted. Notably, microscopy and chemisorption together can be used to understand changes in metal particle size, whereas physisorption can be used to assess lignin deposition effects on catalyst porosity.

When evaluating catalyst systems for lignin-first processes, proper control experiments should be performed. For example, when using heterogeneous catalysts, and especially when developing new catalyst systems, reactions that employ only the catalyst support should be reported. In all cases, reactions with no catalyst should also be performed as a negative control.

In typical kinetic reactions involving a single known reaction with a heterogeneous catalyst, TOF (*i.e.*, the moles of converted substrate/number of active centres/time) is used to describe the catalyst efficiency and can be measured repeatedly to determine its stability. However, determining heterogeneous catalyst active centres for real lignin conversion is quite difficult due to the complex substrates and different reactions that are involved. Different active centres may be involved and affect the overall process simultaneously. For real lignin, a simple measurement of the catalyst productivity (*i.e.*, moles of desired products/weight of whole catalyst/time) and its evolution over time can be used to compare catalysts. Including the mass of the support in this calculation can be justified given that it may play a role in the catalytic cycle. TOF can in turn be used to describe the catalytic efficiency especially for well-defined catalysts (*e.g.*, homogeneous or single-site catalysts) and model compounds of lignins (*e.g.*, model lignin dimers), for which the substrate and active centre are easier to identify.

#### Minimum reporting requirements for assessing catalyst performance in lignin-first processes:

- Catalyst loading, biomass:catalyst ratio, and use and loading of any capping agents in the solvolysis step.
- Depolymerisation catalyst properties including composition and its preparation method.
- When using a new catalyst, control experiments performed with no catalyst and with only the support (for a heterogeneous catalyst).

#### Preferred reporting recommendations for assessing catalyst performance in lignin-first processes:

- Stability and recovery of the catalysts, any stabilising agents, and solvents involved in the solvolysis reactions.
- Depolymerisation catalyst productivity = moles of monophenolics/(total mass of catalyst × time).
- With well-defined catalysts and reactants (*e.g.*, lignin models), TOF can be reported in addition to productivity.
- The structure (*i.e.*, appearance under microscopy, and pore size distribution) and composition (*i.e.*, the metal content) of any heterogeneous catalysts used in the process must be characterised before and after reaction.

## 5. Mass balances and product yields

One of the most acute challenges in lignin-first research is the comparison of data across different research articles. However, such comparisons are key because they facilitate the direct evaluation of the research field's progress by providing a common basis for expressing mass and carbon balances, and product yields. Calculating these yields also facilitates an assessment of the extent to which a lignin-first biorefinery uses the renewable carbon in lignocellulosic biomass to its fullest.

In this section, we distinguish yields that describe the recovery of different component fractions and the yields of individual products. In some processes, the production of component fractions occurs simultaneously with the deconstruction of these fractions to individual products (*i.e.*, specific molecular products). In these cases, yields and balances can be calculated for fractions and individual products for the same



product mixture. One such example includes lignin-derived products (e.g., specific arylpropanols or arylpropanes) and some polysaccharide-derived products (e.g., xylose, glucose) resulting from RCF of whole biomass, in which mass balance, component fraction yields, and individual molecular yields can be estimated from analysis of the liquid-phase RCF products. In other processes (Fig. 2c), fractions will first be isolated but not fully deconstructed. After this initial fractionation step, only initial mass balance and fraction yields can be calculated rather than individual product yields. This is the case for AAF, in which stabilised lignin oligomers are first separated from other biomass fractions. Following this step, an initial yield of isolated lignin can be calculated. The stabilised lignin is subsequently depolymerised in a second process, the analysis of which allows the determination of individual product yields. If subsequent upgrading/refining processes occur, molecular product yields can also be established for these transformations.

In the two subsequent sections, we detail overall and fraction yields and balances and discuss individual product yields that result from depolymerisation and any further transformation.

### 5.1 Overall component balances and fractionation yields

It is important to determine the mass and carbon fluxes throughout process stages. Beginning with lignocellulosic materials, RCF and the initial fractionation stage of other lignin-first processes usually generate two to three main streams: polysaccharide-rich pulp, a possible carbohydrate-derived soluble stream, and a lignin-rich stream. The pulp typically contains cellulose, with variable quantities of hemicelluloses and some residual lignin. In several RCF procedures, this pulp will be physically mixed with the catalyst and difficult to separate. If separation is not possible, an analysis of this entire mixed solid fraction (residual biomass solids and catalyst) can be performed using a standard biomass analysis procedure (see Section 2.2) to measure the carbohydrates. The resulting glucan can be assumed to correspond to the cellulose, and other sugars can be assumed to correspond to the remaining hemicellulose fraction in this pulp. The total solids to which is subtracted the catalyst loading and this cellulose and hemicellulose fraction can give an estimate of the residual lignin. However, this residual lignin content estimate should be considered imprecise because it can be influenced by ash content and any mass loss from the catalyst.

The liquor typically contains solubilised lignin possibly together with carbohydrate derivatives.<sup>47,69</sup> This stream can generally be further purified to isolate the lignin and carbohydrate-derived streams. The lignin stream generally takes the form of a lignin oil in the case of RCF, whereas it can be precipitated as a solid in the case of AAF.<sup>40,83</sup> We recommend at minimum calculating an overall mass balance, and recommend performing an overall carbon balance following any initial conversion stage from native biomass using eqn (8) and (9), respectively:

$$\text{Mass balance(\%)} = \frac{m_{\text{pulp,dry}} + m_{\text{lignin}} + m_{\text{sugars}}}{m_{\text{substrate,dry}}} \times 100 \quad (8)$$

Carbon balance(%)

$$= \frac{\%C_{\text{pulp,dry}}m_{\text{pulp,dry}} + \%C_{\text{lignin}}m_{\text{lignin}} + \%C_{\text{sugars}}m_{\text{sugars}}}{\%C_{\text{substrate,dry}}m_{\text{substrate,dry}}} \times 100 \quad (9)$$

where  $m_{\text{pulp,dry}}$  corresponds to the mass of isolated pulp (on a dry biomass basis),  $m_{\text{lignin}}$  is the mass of isolated lignin or lignin oil from the solvent (if a stabilising functionality has been added, the weight of the stabilising agent should be subtracted from the lignin, *vide infra*),  $m_{\text{sugars}}$  is the mass of sugars or their derivatives solubilised in the liquor (note that the mass will be increased during hydrolysis and/or *via solvent/reagent incorporation* that should be taken into account),  $m_{\text{substrate,dry}}$  is the mass of the substrate;  $\%C_{\text{pulp,dry}}$  is the carbon content of the pulp,  $\%C_{\text{lignin}}$  is the carbon content of the isolated lignin stream,  $\%C_{\text{sugars}}$  is the carbon content of the solubilised carbohydrates or their derivatives from the liquor, and  $\%C_{\text{substrate,dry}}$  is the carbon content of the substrate.

These equations can be adapted to calculate the mass yields of component fractions compared to their respective fractions in the original feedstock. As the carbohydrate-rich pulp and sugars or their derivatives solubilised in the liquor can originate from either cellulose or hemicelluloses, we only recommend estimating the yield of isolated lignin.

$$\text{Lignin yield(\%)} = \frac{m_{\text{lignin}}}{m_{\text{substrate,dry}} \text{ OR } m_{\text{Klason}}} \times 100 \quad (10)$$

where  $m_{\text{Klason}}$  is the total mass of Klason lignin (ASL + AIL) in the original quantity of starting material.

When using a fractionation with stoichiometric capping agents such as AAF, the formation of an additional functionality can increase the mass of the recovered lignin beyond its original mass.<sup>83,186</sup> For AAF, the mass of isolated lignin can be corrected by estimating the fraction of acetal-functionalised groups using the HSQC NMR spectra of the isolated lignin. Specifically, this acetal-functionalised fraction is estimated by taking volume-integral ratios of the peaks corresponding to the appropriate lignin units in the HSQC spectra and then subtracting the calculated mass of the additional functionality from that of the isolated lignin.<sup>186</sup>

Care should also be taken when using biomass with high extractives content because it can interfere with lignin yield estimation. Inadvertently including some extractives in the isolated lignin mass is common and will overestimate the delignification yield. Using pre-extracted biomass (*vide supra*) is a useful control experiment to check for any yield over-estimation.

Given that lignin and carbohydrate substrates may undergo catalytic deoxygenation – often resulting in the production of water – a carbon balance should accompany a mass balance. Specifically, even if no mass losses occur during lignin isolation and solids workup, water release may represent a significant fraction of the missing mass in the lignin-first mass balance. Additional reactions can also occur, including hydrodeoxygenation, cracking of the propyl sidechain of the lignin C<sub>9</sub>-units, demethoxylation, or reactions with C<sub>5</sub> and C<sub>6</sub> sugars, further



increasing the complexity of the product distribution. Moreover, these catalytic processes can also lead to the formation of gaseous products, such as CO<sub>2</sub> and CH<sub>4</sub>, that are difficult to distinguish from those formed by the catalytic decomposition of alcohol solvents often employed in the RCF processes or AAF hydrogenolysis. We note more generally that solvent conversion to less-volatile products and solvent-substrate cross reactivity can also occur, generating species that are difficult to distinguish from those of substrate conversion, but should be considered in overall mass and carbon yields.

Following the catalytic process, work-up losses can also constitute a substantial portion of carbon and mass losses for lignin upgrading processes. For example, successful isolation of the lignin from the liquor will invariably remove water and other volatile components. Rotary evaporation of solvents (under reduced pressure) can also remove some light components of the crude lignin stream.

#### Minimum reporting requirements for mass balances and fractionation yields for lignin-first processes:

- Overall mass balance for biomass components.
- Mass yield of the isolated lignin stream.

#### Preferred reporting requirements for overall mass and carbon balances for lignin-first processes:

- Total mass balance, including all solvents, catalysts, and capping agents.
- Carbon balance for biomass components.

#### Best practices for estimating overall mass balances and fractionation yields:

- Account for addition of any functionality by a stoichiometric capping agent when calculating the mass of isolated lignin or oil.
- Conduct control experiments with and without extraction to account for extractives in mass balances, especially in lignin.

## 5.2 Lignin depolymerisation and transformation

In this section, we discuss the reporting of yields to individual, identifiable products derived from lignin after its depolymerisation. Lignin depolymerisation to monophenolic compounds in RCF occurs simultaneously with fractionation, whereas other methods isolate a lignin fraction to be subsequently depolymerised. For depolymerisation and subsequent catalytic transformations of the resulting monophenolics, mole- and carbon-balances are more straightforward to calculate, as the elemental composition of reactants and products can be determined either *via* elemental analysis or, for simple mixtures, using chromatographic techniques to quantify individual components.<sup>101,187–189</sup> Here, we strongly recommend that yields of individual products be reported whenever possible, rather than reporting more loosely identified fractions (*e.g.*, ethanol-soluble fractions, gel-permeation chromatography (GPC) isolated fractions, or functional groups measured using chemical reagents, *etc.*).

As previously discussed, a pragmatic basis for calculating the yield of lignin-derived monophenolics is the total content of original lignin estimated using the Klason method.<sup>89,187</sup> However, as this method is based on the degradation of the lignocellulosic materials under severely acidic conditions

that substantially alter the structure of the native lignin, estimating the initial share of carbon in native lignin is difficult. Although this can limit the understanding of the molecular mechanisms underlying the deconstruction of native lignin, the expression of yields (*vide infra*) of lignin products relative to the total Klason lignin content of the lignocellulosic substrate is useful. In this context, another way to express product yields is relative to 100 kg of lignocellulosic substrate (on a dry basis).

In both cases, it is essential to base the yields on the lignin content or total weight of the original biomass. This is especially important in cases in which fractionation and depolymerisation are run separately. When lignin is progressively separated from biomass, the first fractions of lignin that are removed from biomass typically give higher monophenolic yields than those removed subsequently.<sup>24,186</sup> If only a small fraction of the lignin was extracted from the original biomass sample, this phenomenon leads to artificially high yields when expressed on an isolated lignin basis.

The next step in calculating lignin yields is to analyse lignin-derived products. The vast majority of lignin monophenolics (or monomer derivatives) that require identification and quantification have structures for which standards can be purchased or synthesised. Furthermore, monophenolics are usually volatile or can be rendered volatile by derivatisation, such as by trimethylsilylation.<sup>24,43,190</sup> Therefore, we recommend initial identification of lignin-derived monophenolics by comparison of their retention times in gas chromatography (GC) or liquid chromatography (LC) with those of authentic standards and by comparison of the mass spectra to those of authentic standards (Fig. 4). To authenticate new compounds, high-resolution MS data and NMR spectra are required. Mass spectra for a large number of lignin monomers have also been published in the literature, which can also be used as a basis for identification.<sup>191</sup> When possible, the similarity index of the EI-MS identification and the calculated *vs.* observed high-resolution mass (of the parent ion) should be reported.

Quantification is often straightforward with GC using a flame-ionisation detector (FID). In this technique, peak areas of identified monomers are converted to concentrations using either calibrations built with external or internal standards (*e.g.*, 2-isopropylphenol or decane). Where standards are available, we recommend using authentic standards with their respective calibration curves. However, as shown several times for lignin monomers,<sup>24,43,186</sup> the ratio of the peak area of the monomer to that of the internal standard can be used to adequately estimate their molar ratio times a correction factor based on the ratio of their effective carbon number (ECN).<sup>192</sup> (**Caution** should be exercised when using the ECN method if the response factors (RFs) of the internal standard to the target compound are > 2). Nevertheless, RFs are significantly affected by the concentration of the internal standard and the target compounds. RFs therefore need to be determined/optimised based on the real concentration of the products. The ECN can also vary slightly depending on the solvent used,<sup>5</sup> so checking the accuracy with standards that are close in structure to the



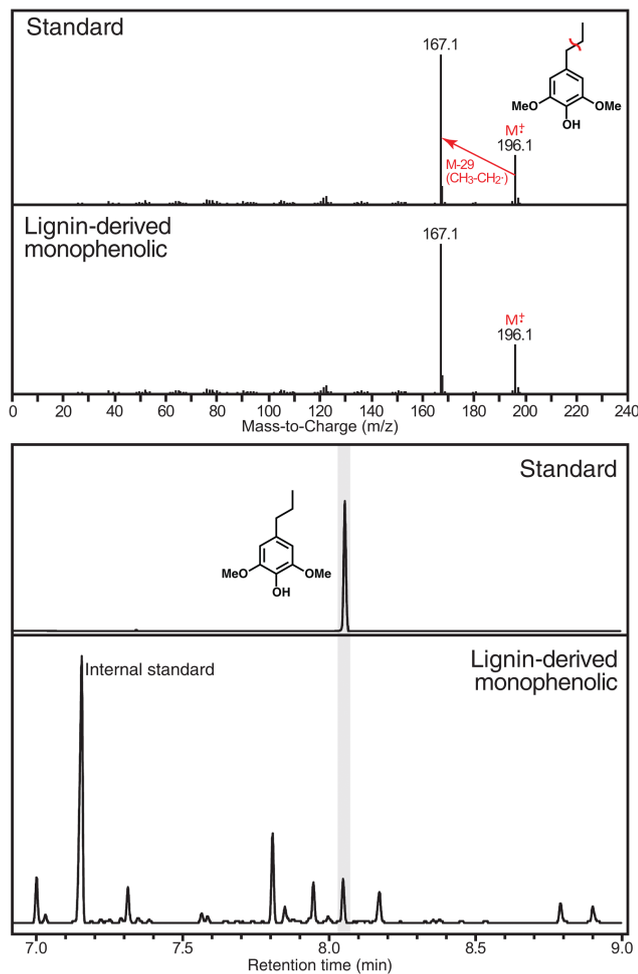


Fig. 4 Representative monophenolic identification by MS comparison combined with retention time similarity in GC—for propyl syringol found in lignin-derived liquor. The top two panels represent the comparison of MS spectra obtained by GC-MS of an authenticated standard and that of the lignin monophenolic within the liquor resulting from hydrogenolysis of AAF lignin. The bottom two panels compare the retention time in GC-FID of the authenticated standard and that of the monophenolic within that same liquor resulting from AAF lignin hydrogenolysis. Other peaks in the bottom GC-FID trace include lignin monophenolics and the internal standard. Adapted from Shuai *et al.*<sup>24</sup>

molecules that require quantification is advised. RFs should be measured for the principal monophenolics and reported. For LC, similar calibrations can be used with UV, diode-array detection (DAD), or MS; external standard calibration curves are generated using pure authentic compounds (which can further aid confirmation of structural assignments), and ideally an internal standard is used. The calibration curves need to be updated frequently as the UV lamp and MS conditions change over time. Calibration standards therefore need to be carefully maintained. Quantification by MS for both LC and GC is fraught with difficulties and, unlike GC-FID RFs, MS response factors are neither universal (across machines) nor stable over time. Use of an internal standard added to the reaction mixture is viable, as long as the internal standard has

similar ionization efficiency in MS (and consequently has a relative RF of < 2 again). For MS-based quantification, the gold standard approach is therefore to use stable-isotope-labelled versions of the exact same compounds injected (in known amounts) into the sample.<sup>171,193,194</sup> As these labelled compounds must often be synthesised in-house, it is hard to make a generic recommendation.

Once the monophenolics are quantified, mass yields can be calculated directly using the mass of the original lignin or biomass (eqn (11)). Typically, RCF or AAF with subsequent hydrogenolysis can produce monophenolics at total individual yields exceeding 5–10 kg per 100 kg of lignocellulosic substrate.

$$\text{Mass yield}_{\text{MP},i}(\%) = \frac{N_i \times \text{MW}_i}{m_{\text{substrate,dry}} \text{ OR } m_{\text{Klason}}} \times 100 \quad (11)$$

where  $\text{Mass yield}_{\text{MP},i}$  represents the mass yield of an individual monophenolic  $i$  produced from lignin,  $N_i$  is the total number of moles of the monophenolic recovered for a given mass of starting material ( $m_{\text{substrate,dry}}$ ),  $\text{MW}_i$  is the molecular weight of the monophenolic from lignin, and  $m_{\text{Klason}}$  is the total mass of Klason lignin in that same mass of starting material.

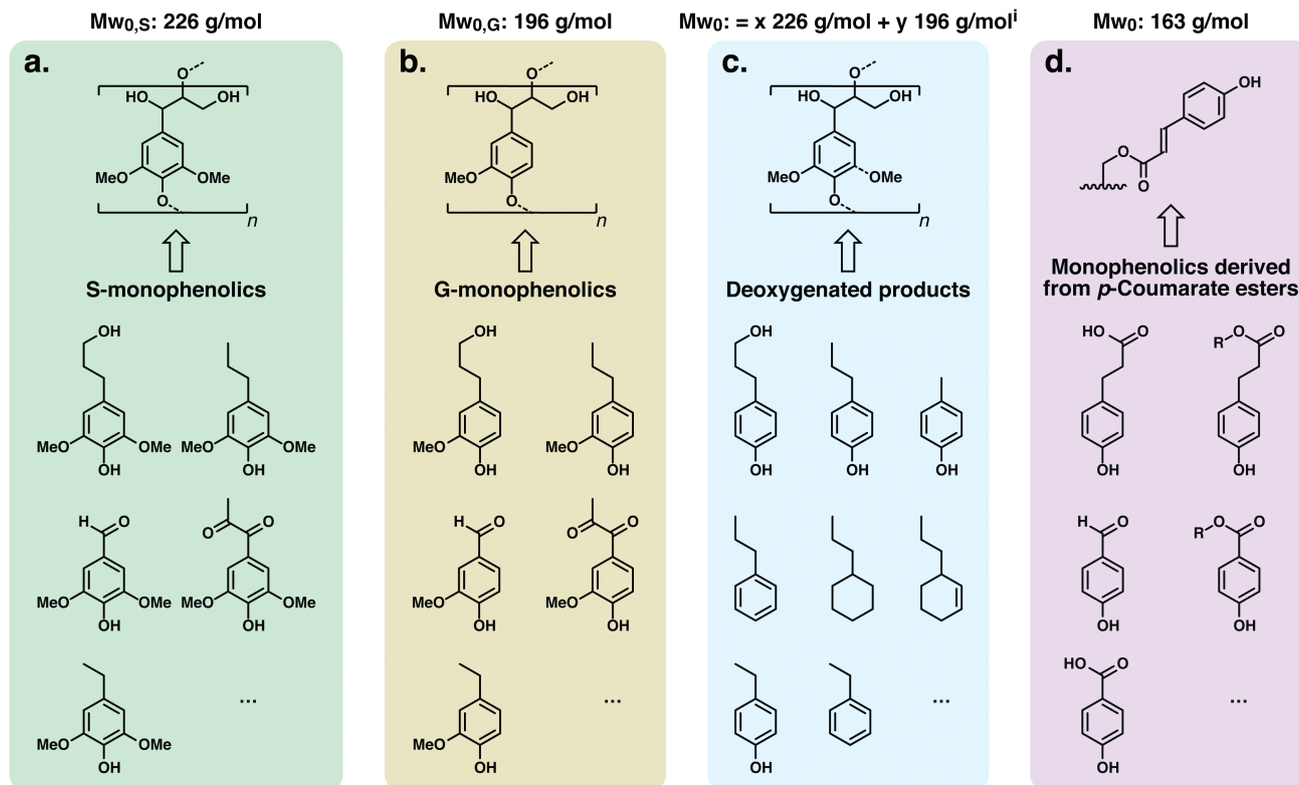
It may also be useful to calculate yields that consider the chemical modifications during depolymerisation (*i.e.*, that account for the transformation and its associated change in mass without affecting the yield). Lignin-first depolymerisation processes will yield monophenolics that differ from their original form in the native lignin structure. Thus, yield calculations can be adapted to account for this transformation and avoid artificially decreasing or increasing the yield. Specifically, this is done by relating the targeted monophenolic to its original structure, which can be assumed to be within a  $\beta$ -ether unit or an ester and using its original molecular weight within native lignin ( $\text{MW}_0$ ). The original molecular weights for various common monophenolic products of lignin depolymerisation are shown in Fig. 5. Eqn (12) can then be used to calculate this yield ( $\text{Yield}_{\text{MP},i}$ ).

$$\text{Yield}_{\text{MP},i}(\%) = \frac{N_i \times \text{MW}_{0,i}}{m_{\text{substrate,dry}} \text{ OR } m_{\text{Klason}}} \times 100 \quad (12)$$

where  $\text{Yield}_{\text{MP},i}$  is the adjusted yield and  $\text{MW}_{0,i}$  is the presumed original molecular weight of the resulting monophenolic in native lignin. The lignin oligomer yield will presumably be the difference between total extracted lignin yield (eqn (10)) and total monomer yield.

These total adjusted yields of monophenolic compounds have a strong correlation to  $\beta$ -O-4 bond content. Several models have attempted to calculate a theoretical monophenolic yield based on the cleavage of ether linkages in lignin. Models that assume a random distribution of C–C and ether linkages within a linear lignin polymer and neglect end-group effects, predict theoretical yields that are equal to the square of the linkage fraction within native lignin that are ethers.<sup>37,44,68,195</sup> These estimates are often surprisingly close to the yields that result from RCF or stabilised lignin on a Klason lignin basis, which are in the low 20% for softwoods, up to ~50% of Klason lignin





**Fig. 5** Approaches to estimating the presumed original molecular weight of units within the native lignin ( $MW_0$ ) for (a) syringyl, (b) guaiacyl, or (c) ill-defined monophenolics.  $i$ , where  $x$  and  $y$  represent the fractions of syringyl and guaiacyl functionalities measured in the original lignin. Fractions of syringyl or guaiacyl functionalities can be measured by various methods including NMR, thioacidolysis, DFRC, nitrobenzene oxidation, or RCF (Section 2.2). Various phenolic esters are associated with lignin (or, in fact, the cell wall polysaccharides) including *p*-hydroxybenzoate in poplar/aspen, willow, and palms, and *p*-coumarate and ferulate in grasses (or commelinid monocots in general) and will produce monophenolics or derivatives from degradative lignin-first methods. (d) *p*-Coumarate as an example ( $MW_0 = 163$ ); the products all derive from *p*-coumarate esters, the level of which is best determined by saponification (and quantified as the *p*-coumaric acid ( $MW = 164$ ) released, by GC-FID, GC-MS, or LC-MS), or perhaps, as the dihydro analog ( $MW = 166$ ) by RCF; estimates from 2D NMR, at least without careful use of quantitative methods, will always be high because *p*-coumarates are (free-phenolic) end-groups that relax much more slowly than internal polymer units. A similar approach can be taken for monophenolics derived from ferulate ( $MW_0 = 193$ ) and *p*-hydroxybenzoate ( $MW_0 = 137$ ) units, if products can be accurately traced back to their corresponding esters on the native lignin. Note: not all possible products from lignin-first methods are shown, as indicated by the ellipsis in each box.

for hardwoods, and as high as almost 80 or even 90% for specially engineered trees containing linear lignins high in ether linkages, such as  $\sim 100\%$  S-lignin poplar,<sup>24,80</sup> or 100% C-lignin seedcoats.<sup>23,28</sup> Thus, it is also recommended to report yields of monophenolic compounds with respect to the  $\beta$ -O-4 ether content determined by NMR, thioacidolysis, or nitrobenzene oxidation (see Section 2.2) and, where applicable, yields of esters from saponification or RCF.<sup>196–200</sup>

To expand the analysis to heavy species, GPC and more elaborate methods can be used. Dimer and oligomer identification and quantification is challenging without model studies to identify their derivation and authentic standards for quantification. GPC can provide an apparent distribution of dimers and oligomers, which is useful as a guide for comparative analyses. However, the data should be analysed with caution. When applied to product mixtures obtained from lignin, direct information regarding the content of species cannot be retrieved from a UV-vis detector as the detector response is not universal. The use of polystyrene standards for GPC

calibration is useful for comparisons but may not be accurately translated to lignin, which can be directly addressed by use of MALS.<sup>201</sup> Moreover, in samples obtained from reductive treatment of lignin streams that contain aliphatic hydrocarbons, compounds may be invisible to the UV-vis detector. Despite these limitations, GPC coupled with UV-Vis spectroscopy provides useful information on the apparent distribution of  $M_w$  and chemical structural uniformity of the eluting species, even if it provides little insight into accurate yields of dimers and higher fractions, or their structures.

Finally, we also recommend performing 1D and 2D NMR, including HSQC, of the product mixture after depolymerization to determine what inter-unit linkage motifs remain in the lignin-derived stream. Model compound experiments are often required to determine what units (with their characteristic inter-unit linkages) have reacted during the depolymerization process, and to provide a reference for assigning resonances in NMR. Examples of NMR-identified and assigned products from RCF are becoming more common.<sup>24,28,67,118,136</sup>



**Minimum reporting requirements for yields of lignin-derived molecules:**

- Monophenolic yields based on the quantification of individually identified products.
- Precise description of the method for formally identifying said molecules (*e.g.*, NMR spectrum, comparison of GC or HPLC retention time to an authenticated standard, comparison of MS spectra with an authenticated spectrum).

**Preferred reporting requirements for yields of lignin-derived molecules:**

- Comparison of yields on a Klason lignin basis with the  $\beta$ -ether content (and ester content, where applicable) of the original biomass, and the associated theoretical maximum yield.
- If targeting individual molecule yields, reporting yields based on the original molecular weight ( $MW_0$ ), based on eqn (12).
- GPC spectra to characterise lignin  $M_w$  distribution, but not for quantification purposes or to calculate yields.
- 2D NMR of product mixtures to gain insight into the composition of product mixtures; caution should be used when deriving quantitative information.
- Response factor calculations for quantified compounds.

**Best practices for estimating yields of lignin-derived molecules:**

- Yields based on the weight of the original biomass or original lignin fraction within that biomass, rather than on the basis of an isolated fraction.

### 5.3 Pulp yields and composition

The lignin-first processes also generate a polysaccharide-rich fraction for further valorisation as a pulp or *via* catalytic, biological, or thermal conversion processes. When performing lignin-first biorefining in batch mode without catalyst baskets, the catalyst and the polysaccharide-rich pulp will be mixed (Fig. 3). Purifying cellulose from the catalyst requires extensive washing, but this separation is needed to determine the degree of delignification and the quality of the polysaccharides. Alternatively, a catalyst basket, magnetic catalyst, or flow-through setup can be used to avoid tedious separations of catalyst and pulp after the reaction (Section 3).

We recommend reporting the weight of the pulp and its composition, which can be determined with the same methods as for whole biomass (Section 2.2). Additionally, we also recommend reporting the digestibility by commercial cellulase cocktails (an NREL LAP is available for this protocol as well at low solids loadings).<sup>91</sup> Determining the digestibility of the pulp with enzymes is useful for several reasons. First, it can be used as a secondary confirmation of the pulp composition. Second, it can demonstrate the potential of the pulp for use in a cellulosic biorefinery, which can be useful for performing initial techno-economic analyses.<sup>16,202</sup> Third, it can indirectly verify the occurrence of reactions between fractionation solvents or capping agents and cellulose, to which enzymes are sometimes sensitive.<sup>83,203</sup>

### 5.4 Product yields from liberated carbohydrates

Lignin-first experiments are usually performed in a solvent mixture containing water and sometimes acid, which typically hydrolyses some or most of the polysaccharide fraction, resulting in simple sugars, their dehydration and hydrogenation products,<sup>69</sup> or carbohydrate oligomers in the liquor. Sugars can also be present in modified forms due to reaction with the solvent (*e.g.*, methyl xyloside when performing RCF in

MeOH),<sup>43</sup> or with a capping agent (*e.g.*, acetal-functionalised sugars).<sup>24,204</sup> If simple sugars need to be isolated, the lignin-derived oil resulting from RCF can be decanted and/or extracted, whereas stabilised lignin can generally be precipitated from an aqueous phase containing sugar-derived products. Sugars can be quantified as described in Section 2.2. However, for modified sugars, including acetal-functionalised sugars, the corresponding external standards need to be used to build a calibration curve. Such standards will not necessarily be available commercially and so may have to be synthesised and purified. Once these sugars or their derivatives are quantified, yields can be calculated using the following equation:

$$\text{Yield}_j(\%) = \frac{N_j \times MW_{(\text{xylan or glucan})}}{m_{(\text{xylan or glucan})}} \times 100 \quad (13)$$

where  $\text{Yield}_j$  is the yield of any carbohydrate derivative ( $j$ ) resulting from one sugar molecule, including a monosaccharide, dehydration product, or modified sugar;  $N_j$  is the total number of moles of  $j$  (products recovered from a single xylan or glucan unit);  $m_{(\text{xylan or glucan})}$  is the total mass of xylan or glucan in the original biomass; and  $MW_{(\text{xylan or glucan})}$  is the molecular weight of this product in the original polysaccharide. For instance, xylan could be converted into xylose, methyl xyloside, or furfural, but all originate from a single xylan unit.

**Minimum reporting requirements for yields of pulp, sugar, or sugar-derived molecules**

- The mass of recovered purified pulp.
- The carbohydrate and Klason lignin composition of the pulp.
- Yields of major monosaccharides or their derivatives.

**Preferred reporting requirements for yields of pulp, sugar or sugar-derived molecules**

- Enzymatic digestibility of the pulp and the yield of the resulting sugars.
- Enzymes (or enzyme cocktail) used, and enzyme loading per mass of polysaccharides.

## 6. Outlook on the analysis and development of lignin-first biorefining

The lignin-first biorefining field has expanded dramatically in the past decade,<sup>9,23,38,47,50,70,79,159,205–209</sup> spurred by the realisation that reductive catalytic processes, although long practiced on whole biomass, are active stabilisation methods that enable production of narrow product slates from lignin to a degree simply unmatched in other lignin deconstruction processes. The advent of protection-group chemistries as alternative lignin-first approaches, such as AAF and DAF, to hydrogenolysis has provided new options to achieve similar process outcomes. Alternative stabilisation chemistries and approaches beyond these three have begun to emerge, including oxidative and photocatalytic routes,<sup>207,210</sup> and undoubtedly many more approaches will be reported in the coming years. The applicability of the guidelines presented here, although considered here in the context of the now well-accepted methods of RCF,



AAF, and DAF, should be broadly applicable to other lignin-first biorefining approaches. In this last section, we briefly discuss the outlook for unifying the lignin-first field around these guidelines.

The chemical composition of whole biomass feedstocks, as described in detail in Section 2, is typically measured *via* summative chemical methods that are now considered standard in the biomass conversion field. Many of these wet chemistry approaches have a long history of development over many decades (and indeed, Klason lignin analysis is over a century old), and are documented in open-access LAPs from NREL.<sup>84–91</sup> These LAPs contain information that researchers should be able to carefully follow to quantify polysaccharide, lignin, protein, ash, extractives, and moisture content in biomass. All equipment and materials in the standard NREL compositional analysis LAP<sup>89</sup> are readily available in most modern chemistry laboratories. This in turn should ideally present a low barrier to entry for laboratories new to the broader biomass conversion field, and certainly lignin-first biorefining, to accomplish these analyses. We stress that for all substrates treated in a lignin-first context, it is imperative to use these standard compositional analysis methodologies to ensure direct comparison of studies over time, across feedstocks, and, as stressed throughout this perspective, across different laboratories.

Eventual accessibility to standardised feedstocks for lignin-first biorefining would also be of significant benefit for the research community, as it would provide a benchmark substrate for all lignin-first researchers to compare new innovations against existing processes. The availability of a benchmark substrate would find utility, whether a researcher is focused on solvolysis, stabilisation chemistry, the catalytic process or catalyst materials development, downstream processing of either the lignin oil or pulp, or both, or overall process development. NIST supplies Biomass Reference Materials that were originally created in the 90s. Moreover, some large government-funded research organisations have established well-characterised feedstock repositories, such as in the US Department of Energy Bioenergy Technologies Office (*via* Idaho National Laboratory) or in the US Department of Energy's Bioenergy Research Centers. However, these substrates are typically not readily available to researchers around the world, which inherently limits the ability to have a truly “universal” feedstock for benchmarking. Going forward, this will become an important need for the growing lignin-first biorefining community – we anticipate pursuing efforts to this end.

Beyond analysing overall biomass composition, quantitative analytics for lignin chemistry have and continue to be a long-standing challenge.<sup>101,211</sup> The RCF version of lignin-first biorefining approach was actually first proposed as an analytical tool to essentially study lignin structure, and recent results<sup>42,62,118,157–160</sup> have shown that thioacidolysis and RCF provide similar C–O bond content measurements from the same feedstocks. Given the potential for an easier analytical approach, we propose that the lignin community might shift away from thioacidolysis and DFRC as routine approaches, and use simpler, less labour-intensive RCF-based chemical methods

for the same measurements. Additional work remains to be done to definitively show that RCF-based chemical methods can substitute for well-accepted approaches, and this work is currently ongoing.

The lignin oil resulting from lignin-first chemistry typically contains monophenolics associated with the cleavage of ether and, where relevant, ester bonds. Beyond monomers, there are dimers and oligomers associated with lignin-first biorefining that derive from C–C-bonded lignin units that are not cleaved in essentially any RCF approaches tested to date, and that may not participate in the protection-group chemistries, as well as unreacted  $\beta$ -ether units. Several reports have used GC-MS to identify dimers,<sup>43,57,212–219</sup> but a re-evaluation is required to thoroughly identify and quantify dimers and oligomers. As an example, products from 4-O-5-linked units using proper lignin models have recently revealed some interesting pathways and a unique dimer.<sup>161</sup> Full analysis of dimers and oligomers represents a major analytical undertaking, as has been accomplished for thioacidolysis recently,<sup>147</sup> but will be crucial to both understanding the impact of lignin-first processes on lignin more comprehensively and for designing downstream separations and valorisation strategies more holistically for the resulting lignin-rich streams.

Similarly, in most lignin-first contexts, some polysaccharides and extractives are liberated along with the lignin. The fate of the carbohydrate-derived compounds has been little explored to date, but here again, more inclusive analyses remain to be done on these compounds as well. For both lignin and off-target products, detailed experimental approaches combined with emerging computational tools,<sup>220–222</sup> will be a key component of a more holistic characterisation of the resulting products from lignin-first biorefining.

Increased standardisation will improve comparisons and process performance across laboratories conducting research around the world. In tandem, improved analytics will allow us to gain better insights into aspects that are not yet understood in lignin-first processing including the fate of more minor products such as certain carbohydrates or extractives. The fate of minor products, small losses of solvents, purification and separation steps, recycling of catalysts, solvents and capping agents, and energy demand are usually ignored at the laboratory scale. However, these aspects are likely to become increasingly important as lignin-first processes move towards industrial implementation. Improved analytics and experimental standardisation, as described here, will allow more effective and quantitative comparisons of studies across the literature, which will in turn enable generation of rigorous process inputs for economic and sustainability modelling, towards robust and realistic scale-up.<sup>172</sup> To this end, industrial implementation of lignin-first processes would represent the ultimate success for this field.

## Conflicts of interest

The authors declare the following competing financial interest(s): MMAO is founder and part owner of Spero Renewables, LLC,



a technology company making plant-based alternatives to petrochemicals. JSL is part owner of Bloom Biorenewables Ltd, which is exploring commercial opportunities for AAF lignin. JSMS is part owner of RenFuel AB and RenFuel Materials AB, that are exploring opportunities for products obtained by lignin-first approaches.

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