A Landscape of Lignocellulosic Biopolymer Transformations into Valuable Molecules by Heterogeneous Catalysis in C’Durable Team at IRCELYON

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Abstract: This review article highlights part of the research activity of the C’Durable team at IRCELYON in the field of sustainable chemistry. This review presents a landscape of the work performed on the valorization of lignocellulosic biopolymers. These studies intend to transform cellulose, hemicellulose and lignin into valuable molecules. The methodology usually consists in evaluating the behavior of the biopolymers in the absence of catalyst under various conditions (solvent, temperature), and then to assess the influence of a catalyst, most often a heterogeneous catalyst, on the reactivity. The most significant results obtained on the upgrading of cellulose and lignin, which have been mainly investigated in the team, will be presented with an opening on studies involving raw lignocellulose.

Keywords: wood; cellulose; lignin; biomolecules; acid catalysis; metal supported catalysis

1. Introduction

The use of biosourced reactants for the production on molecules of interest is of definitive importance for the development of a green chemistry [1,2]. The C’Durable team at IRCELYON has been involved for many years in the valorization of various bioreresources. We particularly use heterogeneous catalysis, which is a primordial tool to implement sustainable procedures [3–6]. Initial works in the team focused on upgrading first generation biomass like vegetable oils [7–10], starch [11–14], glycerol [15–21], simple saccharides [22–31], and their furan derivatives [32–34]. Studies involving second generation biomass, especially lignocellulose (wood and grassy plants), have started more recently and are now predominant. All the past and current research in this area has given to our team a certain notoriety within the French catalysis and green chemistry communities [35]. We would like to present here an overview of the research performed in the C’Durable team on the transformation of lignocellulosic biopolymers. We will focus on the molecular aspects of the catalytic transformations through the main outputs coming from recent projects. Most of them involved the valorization of single biopolymers constituting lignocellulose which are cellulose and lignin. The use of raw lignocellulose has started more recently; general strategies and perspectives for its catalytic transformation will also be presented.

2. Transformation of Cellulose

Cellulose is the main component of lignocellulosic biomass [36]. It is a biopolymer composed of glucosyl units and the most straightforward molecular transformation of cellulose implies a depolymerization into glucose (Scheme 1). However, due to its structural arrangement made of a dense H-bonds network, cellulose is almost chemically inert and insoluble under usual conditions of solvent and temperature. If one excludes biotechnological methods, such transformation into light products requires demanding conditions,
like solvothermal treatments in the presence of a catalyst [37]. Catalysts for depolymerization are usually Brønsted acids, either homogeneous (HCl, H₂SO₄, H₃PO₄) or heterogeneous. Several types of heterogeneous catalysts have been used like zeolites, heteropoly acids [38], but best results were obtained with sulfonated materials. Here, research groups of Zhang [39], Hara [40] and Onda [41] pioneered studies with sulfonated carbons while Jacobs, Sels et al. [42] introduced sulfonated silica and carbon nanocomposites giving glucose with yields over 50 mol%. Water is the adequate reaction medium due to compatibility with formed products and its green solvent character [43,44]. However, due to the reaction conditions to get significant conversion (temperatures over 150 °C on day scale), glucose is rarely formed as sole product and always reacts to give 5-hydroxymethylfurfural (5-HMF) then levulinic acid together with uncontrolled formation of soluble/insoluble oligomers known as humins (Scheme 1) [45]. To try to overcome these issues, non-conventional reaction media like ionic liquids [46], deep eutectic solvents [47], or activation methods like ball-milling, non-thermal plasma and high frequency ultrasound have been successfully introduced, even in the absence of acid catalyst, as demonstrated by Jérôme et al. [48].

![Scheme 1. Cellulose depolymerization in the presence of a Brønsted acid.](image)

When we initiated our research on cellulose transformation with heterogeneous catalysts, we were first interested in evaluating the non-catalyzed reactivity of cellulose in water, that was rarely investigated at that time. We observed for the first time that a reaction temperature of 190 °C was sufficient to liquefy (i.e., dissolve and/or convert) cellulose up to 60 wt% after 120 h, even in the absence of a catalyst [53]. This was possible thanks to the water dissociation at this temperature providing protons in the reaction medium [54]. Nevertheless the latter presented less than 15 wt% of deep depolymerization products (glucose, 5-HMF . . . ). Therefore this non-catalyzed reaction formed preferentially what we name hydrosoluble oligomers and polymers (either as primary or as recondensation products) and hydrochar (Scheme 2) [55].

2.1. Transformation of Cellulose in Water

When we initiated our research on cellulose transformation with heterogeneous catalysts, we were first interested in evaluating the non-catalyzed reactivity of cellulose in water, that was rarely investigated at that time. We observed for the first time that a reaction temperature of 190 °C was sufficient to liquefy (i.e., dissolve and/or convert) cellulose up to 60 wt% after 120 h, even in the absence of a catalyst [53]. This was possible thanks to the water dissociation at this temperature providing protons in the reaction medium [54]. Nevertheless the latter presented less than 15 wt% of deep depolymerization products (glucose, 5-HMF . . . ). Therefore this non-catalyzed reaction formed preferentially what we name hydrosoluble oligomers and polymers (either as primary or as recondensation products) and hydrochar (Scheme 2) [55].
Moreover it appeared that the intrinsic characteristics of the cellulose are a parameter influencing its reactivity and the morphology plays a role more important than polymerization degree or particle size. Celluloses with a fiber structure, despite a much higher polymerization degree (DP), react faster than those in particle forms (Figure 1) [56].

Then we assessed for the first time the behaviour of solid Lewis acids as catalysts for the depolymerization of cellulose [57,58]. We studied the reaction in the presence of tungstated zirconia $\text{WO}_x\text{ZrO}_2$ and alumina $\text{WO}_x\text{Al}_2\text{O}_3$. Yields are determined based on the amount of moles of initial glucosyl units $C_6H_{10}O_5$, $(n_{\text{glucosyl units}} = m_{\text{dry cellulose}}/162)$, corrected by the number of carbons in the product. We observed a different reactivity than that occurring with Brønsted acids (see Scheme 1). While the latter form preferentially

\[ In the absence of catalyst: k_2 \gg k_1 = k_3 \gg k_4 \]
levulinic acid, Lewis acids formed lactic acid as main product up to 30 mol% yield [59,60], a possible monomer for biosourced plastics. This corresponds to the first report on the formation of lactic acid from cellulose, even more with a water-tolerant heterogeneous catalyst [61]. Indeed, studies reported at the same time by Taarning et al. described the formation of lactic acid with zeolites from monomeric sugars only [62]. The reaction mechanism supposedly involves the presence of vacant Lewis sites at the surface of the catalyst able to promote dihydroxylation and carbon-carbon bond cleavages of the glucose chains in addition to hydride transfers giving pyruvaldehyde as intermediate [60]. Note that when lignocellulose was used as reactant in place of cellulose in same conditions, very close results were obtained, showing that the presence of other wood components may have no impact on the transformation of cellulose into lactic acid (see Section 4 for more details) [63]. When WO\textsubscript{x}-ZrO\textsubscript{2} or WO\textsubscript{x}-Al\textsubscript{2}O\textsubscript{3} served as support for Pt, and the transformation performed under H\textsubscript{2}, we observed the reduction of the intermediate to acetol and propylene glycol through successive hydrogenation steps (Scheme 3) [64,65].

![Scheme 3](image)

**Scheme 3.** Reaction pathways of cellulose transformation in the presence of solid Lewis acids (water 65 mL, Avicel cellulose 1.6 g, catalyst 0.68 g, 190 °C, 5 MPa 24 h). Yield = 100 × (n_{product}/n_{glucosyl units}) × (3/6). Adapted from [60,64].

Interestingly, in the absence of Pt but still under H\textsubscript{2} atmosphere, another different route occurred forming 2,5-hexanediol as main product (Scheme 4) with WO\textsubscript{x}-ZrO\textsubscript{2} [66]. This was possible from hydroxyl groups abstraction promoted by this catalyst under the hydrothermal conditions as proven by quantitative Infra-Red spectroscopy analysis and kinetic investigations [67]. This highlights the different mode of action of WO\textsubscript{x}-ZrO\textsubscript{2} vs. WO\textsubscript{x}-Al\textsubscript{2}O\textsubscript{3} in the depolymerization of cellulose. That was the first report on direct 2,5-hexanediol formation from cellulose, a molecule of high interest for green fuels aviation synthesis by aldol condensation [68].

2.2. Transformation of Cellulose in Liquid and Supercritical Organic Solvents

The reactivity of cellulose in alcohols was studied with the aim of forming levulinic esters as high-interest biosourced molecules for energy or chemical applications [69]. The alcohol serves as reaction medium and esterifying agent. Levulinic esters are usually obtained by esterification of levulinic acid in the presence of a variety of catalytic systems; however, the possibility of forming these compounds from cellulose has been previously demonstrated by Garves, however, without real further investigation since then [70].
Scheme 4. Formation of 2,5-hexanedione from cellulose in the presence of solid Lewis acid (water 65 mL, Avicel cellulose 1.6 g, catalyst 0.68 g, 190 °C, 5 MPa 24 h). Yield = 100 × (n_{product}/n_{glucosyl units}) × (6/6). Adapted from [66].

We showed that reacting cellulose in methanol under supercritical conditions led to the deep liquefaction within few minutes, even in the absence of catalyst. The supercritical conditions were necessary to get this very fast reactivity (Figure 2). The presence of the solid Brønsted acid Cs_{52.5}H_{0.5}PW_{12}O_{40} formed as expected methyl levulinate in 20 mol% yield [71]. For such reactions the presence of 10 wt% of water in methanol enhanced the overall reactivity.

The reaction was extended to butanol to form butyl levulinates, and in this case we did not observe the influence of these supercritical conditions on the global reactivity. However, we demonstrated the importance of the butanol isomer on the formation of the corresponding butyl levulinate. Here the primary butanols gave the best yields up to 50 mol%, both in the presence of H_{2}SO_{4} or Cs_{52}HPW_{12}O_{40} (Figure 3) [72]. Note that tert-butyl levulinate was formed as traces due to fast dehydration of tert-butanol preventing its role of esterifying agent.

Figure 2. Cellulose transformation into methyl levulinate in supercritical MeOH (methanol 90 wt%, 25 g, cellulose 0.35 g, catalyst if present 0.15 g). Yield = 100 × (n_{product}/n_{glucosyl units}) × (5/6). Adapted from [71].
With the aim of avoiding water formation during the esterification step with a more efficient atom utilisation, we assessed butene s as esterifying agent. Previous study has been reported by Dumesic and co. forming butyl levulinates in two-step procedures through reactive extraction in the presence of sulfuric acid [73]. In our case, the challenge was to find conditions able to accommodate all reagents for a one-step transformation. With the use of H$_2$SO$_4$ in an organic solvent, iso-octane, we were able to form a levulinic ester from the reaction of cellulose with an olefin (Scheme 5). Sec-butyl levulinate was formed in a 19 mol% yield, higher than using the corresponding alcohol. Attempts to change iso-octane to a biosourced solvent, γ-butyrolactone, led to the formation of sec-butyl levulinate, with a 5 mol% yield. This confirms the difficulty to find suitable conditions for a highly sustainable procedure [74].

Finally, as a chemical application of butyl levulinates, with showed for the first time their use as biosourced solvents for synthetic chemistry like C-C couplings. Vaccaro et al.
previously demonstrated the ability of γ-valerolactone (GVL) as biosourced solvent for this chemistry with therefore the potential to replace (toxic) organic solvents generally used for such transformations [75]. However, since alkyl levulinates are the precursors of GVL, it would be more interesting to use them directly as solvents. Here, n-butyl levulinate showed to be as suitable as GVL or organic solvents for Heck C-C coupling, even in the presence of a heterogeneous catalyst. The system solvent/catalyst has shown to be completely recoverable and reusable compiling many requirements for a green synthetic route (Scheme 6) [76].

Finally, as a chemical application of butyl levulinates, with showed for the first time

\[
\begin{align*}
\text{Pd@NaY} & \quad \text{solvent: n-butyl levulinate} \\
\text{R = o-, m-, p-MeO, NO}_2 & \quad \text{reusable system} \\
& \quad 85-99 \text{ mol\% yield}
\end{align*}
\]

Scheme 6. Heck coupling in n-butyl levulinate as solvent (solvent 6 mL, aryl halide 6 mmol, n-butyl acrylate 7 mmol, NET3 6.6 mmol, Pd 6 µmol, 150 °C, 6 h). Yield_{product} = 100 × (n_{product}/n_{Ar-I}). Adapted from [76].

3. Transformation of Lignin

Lignin is another biopolymer constituting lignocellulose. Different from cellulose and hemicellulose, which are polysaccharides, lignin is constituted of aromatic units linked through a variety of bonds, each having a special reactivity. Indeed, the lignin structure (Figure 4) is obtained from the polymerization of three aromatic alcohols (coniferyl, sinapyl, and coumaryl), leading to a variety of compositions depending on the lignocellulose type (soft or hard wood, grass plants, vegetal wastes . . . ).

Lignin can be a renewable source of aromatic chemicals, either for energy like toluene and for chemistry like vanillin, and their derivatives. Therefore, the number of studies has expanded rapidly during the past few years to valorise this biopolymer [77,78]. Its selective transformation represents a great challenge and in addition to the fact that lignin structure depends on its botanical origin, the extracted lignin is different than the native one because the extraction method can create new bonds between aromatic moieties. In consequence, and despite the fact that β-O-4 linkages correspond to half of the total interunit bonds, it appears initially difficult to consider general rules for catalytic lignin reactivity. Various methods of transformation into value-added molecules have been assessed in the literature, from enzymatic to chemo and organometallic catalysis. As we will see below, heterogeneous catalysis has also a role to play because it allows transformations using relatively demanding conditions leading to a specific reactivity of lignin.

3.1. Transformation of Lignin under Neutral Atmosphere

Preliminary studies in the team consisted in comparing, under neutral atmosphere, the solvolysis of lignins from different origins and extraction methods: a spruce-lignin and a pine-lignin obtained from Kraft processes, a wheat straw-lignin obtained from Soda process, and a wheat straw lignin obtained from Organosolv (CIMV®) process [79]. After full sample characterization by a variety of techniques, solvolysis was studied in EtOH-H$_2$O under continuous flow conditions at 225 °C under 8 MPa of N$_2$, preventing char and foam formation. Generally, after treating the reaction mixture three phases were obtained; the Klasson phase corresponding to non-liquefied lignin, the aqueous phase containing mainly aliphatic water-soluble compounds, and the organic phase containing the target aromatic products. We showed the importance of the amount of EtOH (up to a 50 wt% mixture) on the content of monomeric organic products formed during the treatment. EtOH favours lignin conversion and stabilises organic products in which guaiacol was predominant due to β-O-4 bond cleavages (Figure 5). In these conditions, Kraft-spruce-lignin presented a higher reactivity (up to 76% of conversion) due to its natural solubility in EtOH-H$_2$O while
the other lignins necessitated the addition of K$_2$CO$_3$. Globally, it appears that product formation depends more on the origin of the lignins, rather than on their extraction process. When the reaction was performed on Kraft-spruce-lignin but in a closed reactor, the nature of the alcohol co-solvent had an impact on the amount of monomeric products. It increased from 0.8 to 4.8 wt% in the presence of iso-PrOH due to the formation of alkylated guaiacol species. After these non-catalyzed transformations, the influence of a supported metal catalyst under similar conditions using H$_2$O-iso-PrOH solvents in an autoclave reactor was assessed. Pd/ZrO$_2$ showed almost no influence on organic product formation, giving 4.4 wt% of aromatic products but increased significantly the total yield of organic products mainly oligomers from 45 to 54 wt%. Note that when using Pd(OAC)$_2$ as soluble catalyst, close results were obtained indicating in both cases the homogeneous nature of the reaction from soluble Pd species leached from Pd/ZrO$_2$. In the last case we supposed that leached species performed primary depolymerization of lignin into monomers that further reacted on supported Pd species [80].

![Figure 4](image_url)

Figure 4. Example of lignin structure highlighting the diversity of constituting units and interunit bonds.

3.2. Transformation of Lignin under Oxidative Atmosphere

The reactivity of lignin in neutral conditions appears to give a very low amount of value-added molecules. Therefore, main catalytic approaches for the up-grading of lignin involve a reacting atmosphere, either reductive (H$_2$) to produce deoxygenated aromatics for energy use, or oxidative (O$_2$, air) to produce oxygenated aromatics for fine chemistry [81]. As far as heterogeneous catalysis is concerned, oxygenated aromatics can be obtained particularly with perovskites [82], mixed-metal oxides such as FeCoO as demonstrated by Barakat et al. [83], all giving global yields into aromatic aldehydes (vanillin, syringaldehyde . . . ) in the 15–20 wt% range. Systems with supported metals have also been investigated with Co, Au, and Pd. Sales et al. have extensively studied Pd catalysts like Pd/γ-Al$_2$O$_3$, leading also around 10 wt% of aromatic aldehydes [84].
Due to the molecular nature of the products obtained from the oxidative route, our team has been involved in the oxidative depolymerization of lignin for many years, particularly in water solvent. Pursuing with Pd systems, we prepared new Pd catalysts based on soluble PVP-stabilized or oxide supported nanoparticles (Pd/TiO₂) (Figure 6). The homogeneous system was very efficient for the selective oxidation of benzyl alcohol as lignin models such as vanillic and veratryl alcohols into the corresponding aldehydes into mild conditions (80 °C, O₂ flow) [85]. However, when applied to the oxidative depolymerization of a Kraft-pine-lignin [86], despite enhancing the lignin conversion compared to the absence of catalyst, the two Pd-systems did not improve the formation of the target products like vanillin and acetovanillone for which ones the yield remain low compared to catalyst free oxidative lignin depolymerization (i.e., for softwood Kraft lignins: from 2.4 wt% for non-catalyzed reaction to ca 2.2–2.5 wt% for Pd-catalyzed reactions) [87]. However, this reflects only an apparent lack of Pd-catalyst activity. Indeed, we demonstrated that these catalysts act more probably for oxidizing the Cα-OH (or Cγ-OH) junctions in β-O-4 bonds according to our studies on benzyl alcohol, before base catalyzed reactions take place (i.e., retro-aldol, tautomerization . . . ) for breaking these bonds and delivering the expected aromatics [86]. Additionally, Pd-catalysts seem to favor recondensation reactions, thus enhancing the amount of Kason phase and, despite good reactivity in catalyzed reactions, did not deliver further aromatics.

Catalysts based on other metals, also known for their oxidation features have been employed (Figure 6). Here, we showed that Au/TiO₂ had the tendency to further oxidize formed aromatics into volatile molecules. Thus very low yields into vanillin (0.2 wt%) were achieved. This result contrasts with that obtained when using Au nanoparticles in suspension that deliver results close to what was observed with Pd [86]. Supporting Au nanoparticles onto TiO₂ seems to affect their activity. Pt/TiO₂, unlike Pd catalysts, gave slightly enhanced yields of vanillin (from 2.4 wt% to 3.4 wt% for softwood Kraft lignin) and “syringaldehyde and vanillin” (from 4.6 wt% to 7.7 wt%, mainly on syringaldehyde). A kinetic study with this last catalyst indicated that it preserved the lignin substructures leading to a better yield of aromatics [88]. Later on, catalysts based on Cu have been prepared and used for the oxidative depolymerization of Kraft-pine-lignin. Compared to the non-catalyzed transformation, the presence of CuO/TiO₂ did not affect the global lignin liquefaction (remaining around 70%) but increased the amount of valuable oxygenated aromatics especially vanillin formed up to 5 wt% in an autoclave reactor. In a continuous
flow reactor this catalyst allows full lignin liquefaction with a higher aromatic molecule yield but without enhancing vanillin yield [89,90].

Figure 6. Catalyst development in C'Durable team for oxidative lignin depolymerization to aromatics, mainly vanillin.

4. Transformation of Lignocellulose

As presented above, the selective transformation to molecules of interest of isolated lignocellulosic biopolymers is in itself difficult. Our team reported interesting and promising results during the past decade but progress to enhance our initial results was far from what we could expect. There are possible reasons for that. Firstly, it is now well accepted that the lignocellulose fractionation processes into isolated cellulose and lignin modify significantly the structure of the native biopolymer and make it less reactive. This is clearly the case for all commercial samples used in the studies described above. For example, commercial celluloses are more crystalline than the native cellulose and are available with different characteristics (morphology, polymerization degree, crystallinity). These parameters influence the depolymerization and valorization efficiency. This explains why various pre-treatments are often applied to cellulose to enhance reactivity. The drawback is an increase of the heterogeneity of the solid reactant giving uncontrolled variabilities of the physico-chemical features. This also true for lignin, for which the origin and extraction method impact very much subsequent transformations. As a consequence, for both biopolymers data comparison has become very difficult for the scientific community. Secondly, a limitation relies on the difficulty to quantify the substrate conversion. Most of the studies on cellulose or lignin reactivity monitor the reaction progress based on the mass of solid residue. This can be mistaken in case of high molecular weight solid products (oligomers) formation and/or the use of a solid catalysts present in the residue (char, Klason lignin). Also, soluble oligomers present in product solution are usually not analyzed, hiding part of the products
and distort data analysis. In consequence, it is of crucial importance to develop efficient and rapid analytical tools to accurately assess the biopolymers reactivity from a conversion and product yield point of view. In the case of cellulose, we attempted to improve this point using Infra-Red spectroscopy to quantify the unconverted solid to get real kinetics of conversion or applying liquid chromatography to analyse soluble oligosaccharide [67]. For instance, these reliable tools demonstrated the impact of solid catalysts WO$_x$/ZrO$_2$ on the initial rate of non-pre-treated cellulose conversion (see below).

The above comments give a clear idea on what can be the challenge with these biopolymers within a native lignocellulosic matrix. Indeed, if the main advantage is to have the biopolymers presumably more reactive, here come the potential issues of their interference during their respective catalytic transformation to a specific product, for instance impact on the accessibility to a heterogeneous catalyst, on poisoning of active sites by uncontrolled conversion of the other biopolymers. Nevertheless several groups are investigating the reactivity of raw biomass with heterogeneous catalysts. One can cite Fukuoka et al. [91] and Zhang et al. [92,93], who applied catalytic systems to wood reactivity to transform the carbohydrate biopolymers (cellulose and hemicellulose), into sugar-derived products giving close yields without apparent conversion of the lignin fraction. Wang et al. reported the valorization in a one-pot procedure with a single catalyst of all components of a lignocellulosic substrate into polyols from carbohydrate fractions, and cyclic alkanes from lignin [94]. While most of such studies try to valorize the carbohydrate fraction, releasing degraded lignin for challenging subsequent reactivity, Sels et al. developed the lignin-first concept of biomass transformation. This concept aims, during lignocellulose deconstruction, at lignin stabilization when depolymerization for example through hydrogenolysis for a more efficient catalytic valorization [95].

Another issue when the objective is the selective formation of a single product (or more modestly on a family of products), is to be able to convert one biopolymer keeping the others unchanged the most as possible, for parallel valorization. In this objective, again, analytical methods are very important when using lignocellulosic substrates of different nature depending on their origin and obtaining. Pertinent procedures for the preparation of the sample (drying, milling, sieving), the determination of its exact composition (cellulose, hemicellulose, lignin) before reaction and in solid reaction products are necessary in order to monitor the transformation progress. Among others, techniques for sample preparation, compositional analysis (acid hydrolysis combined to advanced Infra-Red spectroscopic methods), structural analysis (DRX, TGA, NMR) are particularly mastered in our research team [67,96]. The results we obtained on cellulose and lignin have participated to pave the way to study the reactivity of lignocellulose as a whole.

We have presented above the peculiar case of lactic acid formation from pine wood sawdust hydrothermal treatment in the presence of WO$_x$/ZrO$_2$ Lewis acid catalyst. Compared to commercial cellulose, results showed similar kinetic profiles indicated that the presence of other wood components like lignin did not affect the accessibility of the catalyst to the carbohydrate fractions (intrinsic hindrance), nor poison the active sites responsible for lactic acid formation. Moreover, a higher initial rate of product formation was observed from wood (Figure 7) certainly due to the different feature of cellulose (fiber morphology, lower crystallinity) and/or the contribution of hemicellulose in wood making its carbohydrate fraction more reactive [63]. Interestingly, it was shown by Infra-Red spectroscopy of the solid residue that the lignin component remained unchanged, certainly contributing to this result.

While the above result suggests that a fractionating pre-treatment is not always necessary to get specific molecules from lignocellulose [97,98], our team is investigating an original concept consisting in lignocellulose fragmentation in supercritical organic solvents in the presence of a catalyst. The supercritical conditions of an organic solvent may allow a partial selective conversion/liquefaction of lignocellulose components. Indeed, it is known that supercritical fluids possess solvation properties different than those under usual state. The solvation abilities of supercritical solvents can be tuned near the critical point by small
change in the temperature, the fluid density. The composition and nature of the supercritical fluid (pure solvent or mixture) is also of prime importance regarding the chemical nature of the solid substrate to be solubilized. Thus, this methodology relies on the assumption to selectively solubilize one of the wood components, (hemi)cellulose vs. lignin by choosing the nature of the supercritical fluid and conditions. As an example, we investigated the behaviour of pine wood sawdust in ethanol and heptane in supercritical state (critical points 241 °C and 6.1 MPa, 267 °C and 27 MPa, respectively) at 280 °C in autoclave reactor. Figure 8a shows that depending on the supercritical fluid, the biopolymers conversions are different. For example, lignin was more converted than hemicellulose in ethanol at 280 °C (10 MPa), and this was the contrary in heptane. In heptane, under the same conditions, hemicellulose was completely converted vs. 40 wt% for cellulose and lignin. The product families were also different, giving mainly bio-oils in ethanol (35 wt%) and bio-char in heptane (40 wt%) (Figure 8b) [99].

![Figure 7](image1.png)

**Figure 7.** Comparison of cellulose and lignocellulose transformation in the presence of WOx-ZrO2 (water 65 g, wood 0.68 g, catalyst 0.3 g, 190 °C). Adapted from [63].

![Figure 8](image2.png)

**Figure 8.** Pine wood (a) conversion and (b) products yields, in supercritical ethanol or supercritical heptane (solvent 27 g, wood 2.5 g, no catalyst, autogenous pressure, 1 h), in the absence of catalyst.

For a specific solvent, fine condition tuning such as the fluid density influences wood conversion and product distribution. Treating pine wood with supercritical ethanol of higher density led to more bio-oil and less bio-char, most likely explained by clustering effect which may favour reaction between ethanol and the biopolymer fragments at the expense of their recondensation (Figure 9) [100]. These results confirm that (partial) selective fractionation of lignocellulose is feasible by using a solvent of adequate polarity and
functionality and playing around the reaction conditions, especially near the critical point of the solvent.

![Graph showing yield and conversion (%)](image)

**Figure 9.** Pine wood conversion in supercritical ethanol at 250 °C. Influence of the fluid density on the yields of bio-char, bio-oils, light products, and gases, in the absence of catalyst. Adapted from [100].

In the absence of catalyst, the selective formation of liquid light products was clearly not favoured, and the conversion/liquefaction mainly produced bio-oil up to 30 wt% of initial substrate, using supercritical ethanol for example.

This is why the transformation strategy can also include the presence of a catalyst, at the early stage of the fractionation, in order to drive a liquefaction into the formation of a specific product. Some possibilities are presented in Figure 10. Studies are currently in progress in the team to determine the influence of the characteristics of various solvents and catalyst on the sequential lignocellulose component liquefaction and the nature and composition of the products [101].

![Diagram showing lignocellulosic biomass, supercritical organic solvent, catalyst, and products](image)

**Figure 10.** Strategy for lignocellulose fractionation in supercritical solvents in the presence of a catalyst.

An additional possibility to perform more selective transformations is the use of a semi-continuous flow reactor in which the solvent circulates through a fixed-bed formed of reactant and catalyst. Advantages are the possibility to finely and independently tune the temperature and pressure and to control more precisely the residence time under certain conditions. This may impact favourably the selectivity of the transformation, for example, by avoiding modification of products as observed in autoclave reactors [99], or limiting successive reactions. We initiated this concept by reacting a fixed-bed of cellulose or spruce wood maintained in an open tubular reactor (beforehand wet-impregnated with 0.1N solution of mineral acid) through the percolation of supercritical 2-butene (critical
points 146 °C and 4.0 MPa) under various conditions. A yield up to 13 mol% of sec-butyl levulinate (based on cellulose content) was achieved through a protolysis and esterification sequence (Figure 11) [102,103]. Note the importance of the presence and nature of adsorbed mineral acid on the efficiency of the reaction under these conditions.

Figure 11. Biomass liquefaction and sec-butyl levulinate formation after treatment under super-critical 2-butene (flow 0.1 mL·min⁻¹, reactant 2 g, 10 MPa He, 1.5 h).

5. Conclusions

This review describes the main advances obtained in the C’Durable team at IRCELYON in the area of molecular transformation of lignocellulose biopolymers for the last decade. Our goal is to form value-added molecules, using sustainable processes as much as possible. In this view the use of heterogeneous catalysis is a tool with definitive advantages to drive the reactivity towards robust and selective formation of products of interest. To date we have been mainly working on cellulose and lignin transformation applying an approach consisting in studying the non-catalyzed transformation, then the impact of heterogeneous catalysts on the conversion and product formation. We had the chance to face with many problematics that can be encountered in this field of research, giving us a deep scientific and technical know-how to develop catalytic processes for the valorization of this bioresource. Henceforth we try at present to apply all this knowledge to efficient transformation of raw lignocellulose, which we believe is going to be one of the future challenges in the field of green chemistry for a sustainable development.

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