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Lignin-based porous carbon adsorbents for CO₂ capture

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A major driver of global climate change is the rising concentration of atmospheric CO_2 , the mitigation of which requires the development of efficient and sustainable carbon capture technologies. Solid porous adsorbents have emerged as promising alternatives to liquid amine counterparts due to their potential to reduce regeneration costs. Among them, porous carbons stand out for their high surface area, tailorable pore structure, and exceptional thermal and mechanical properties, making them highly robust and efficient in cycling operations. Moreover, porous carbons can be synthesized from readily available organic (waste) streams, reducing costs and promoting circularity. Lignin, a renewable and abundant by-product of the forest products industry and emerging biorefineries, is a complex organic polymer with a high carbon content, making it a suitable precursor for carbon-based adsorbents. This review explores lignin's sources, structure, and thermal properties, as well as traditional and emerging methods for producing lignin-based porous adsorbents. We examine the physicochemical properties, CO_2 adsorption mechanisms, and performance of lignin-derived materials. Additionally, the review highlights recent advances in lignin valorization and provides critical insights into optimizing the design of lignin-based adsorbents to enhance CO_2 capture efficiency. Finally, it addresses the prospects and challenges in the field, emphasizing the significant role that lignin-derived materials could play in advancing sustainable carbon capture technologies and mitigating climate change.

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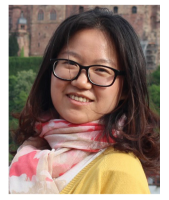
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1 Introduction

The mitigation of climate change is a pressing contemporary issue that motivates research across diverse scientific disciplines and influences evolving government policies worldwide. The consequences of climate change are realized in their impacts on marine and freshwater ecosystems, water and food security, economy and human migration, health and wellbeing.1 Climate change is driven by a combination of natural variations and other contributing factors, including anthropogenic activities, such as atmospheric pollution. Among the main greenhouse gases (GHG)—carbon dioxide (CO₂), methane (CH₄), nitrous oxide (N₂O), and water vapor (H₂O_{vap})—the atmospheric concentration of CO2 shows a particularly strong correlation with rising surface temperatures.²⁻⁵

Since pre-industrial times, atmospheric CO₂ has risen from 280 ppm to over 400 ppm.6 This trend is expected to continue unless drastic changes take place following increased public awareness, and the adoption of new regulations and policies along with remediation via sustainable and cost-effective technologies. A dominant source of anthropogenic GHG emissions has been CO2 emissions from the combustion of fossil fuels, accounting for 88% of the total emissions over the last decade.⁷ In the long term, a significant amount of fossil energy is anticipated to be replaced with 'green' sources (e.g., solar, wind, nuclear, and hydro). However, it will take time to phase out fossil fuels and products due to their ingrained relationship with modern civilization. Beyond petroleum, many other industrial activities also produce CO2, for instance, as reaction by-products (e.g., cement and steel). Therefore, in the short



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Ingo Burgert

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building materials, focusing on enhancing the functionality and life cycle efficiency of wood-based composites.

term, emission reduction and mitigation via CO2 capture are necessary.8

Generally, CO₂ capture works by separating and concentrating it from gas streams, whereupon the CO2 is compressed and transported for storage or utilization. In the broad context, carbon capture approaches can be categorized as carbon capture and storage (CCS), carbon capture and utilization (CCU), and reactive carbon capture (RCC). In CCS, the captured carbon is typically stored in reservoirs such as geological formations for permanent storage. Alternatively, CCU approaches seek to utilize CO2 after capture and storage. In contrast to CCS and CCU, RCC immediately converts CO2 to another product, potentially significantly improving energy efficiency by eliminating the thermal energy requirement for CO2 release.

Current carbon capture technologies include postcombustion (CO2 and N2 from flue gas), pre-combustion (CO2 and H₂ from syngas), or oxy-fuel combustion (O₂ from air) as classified by the composition of the stream (i.e., the compounds to be separated).¹⁰ The implications of these distinctions are realized in the different processes and conditions required for separation. Today, with currently available carbon capture technologies, power plants could reduce their carbon emissions to close-to-zero, however, at the cost of considerable energy and financial penalties. 11 For example, within the framework of post-combustion carbon capture, amine-based liquids are among the most mature technologies applied to scrub CO2 from flue gas based on the formation of watersoluble carbamates or bicarbonate species. Yet, major challenges exist in the form of the regeneration of aqueous solvents, which requires CO₂ stripping at high temperatures. ¹² Such demands, associated with major capital and operation costs, energy, and solvent intensity, limit or prevent the adoption of technologies that have already been demonstrated for their technical feasibility.

To fulfill the ever-increasing pursuit of high-performance carbon capture materials, porous solid adsorbents have garnered attention due to their high surface area and CO2 selectivity. Compared to liquid adsorbents that rely on chemisorption, porous solids mainly work by physisorption, leading to efficient gas separation, thereby reducing energy requirements during regeneration, especially under the conditions of a low adsorption enthalpy. Porous solids can be sourced at low costs and can be tailored for morphology and stability under harsh operating conditions (high temperatures, pressures, and humidity). 10,13 Of critical importance, solid adsorbents can be regenerated more efficiently than liquid adsorbents, for example, by pressure or temperature swings. Notably, these regeneration approaches imply reduced space requirements and system complexity, although, in practice, their implementation is highly dependent on the type of solid adsorbent used and the possibility of use at scale. 14 Regeneration of solid adsorbents often drives the techno-economic feasibility of the technology.

Typical solid adsorbents include zeolites, metal-organic frameworks (MOFs), covalent organic frameworks (COFs) and activated carbons, all of which display high porosity, available specific surface area and physisorption capability. However, zeolites, MOFs, and COFs are often moisture-sensitive, expensive, and difficult to regenerate.¹⁴ Various alternative technologies have also been developed, including renewable feedstocks such as amine-functionalized fibrillated cellulose, or nanochitin aerogels. 15-18 Among these materials, porous carbons are promising solid adsorbents for CO2. Porous carbons can be produced from renewable, carbon-rich agricultural or industrial by-products and residual streams, offering an opportunity to add value to these resources. Current efforts highlight material conversion from a variety of bioresources such as wood sawdust,19 fruit seeds,19 oil palm,20 gelatin and starch,²¹ date seeds,²² nutshells,²³ and food wastes,^{24,25} to name just a few. While these materials present significant opportunities for conversion into activated carbons, a key challenge persists: their lower CO2 adsorption capacity and selectivity



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Orlando Rojas is a Canada Excellence Research Chair and Director of the Bioproducts Institute, where he synergizes a distinguished group of researchers conducting multi-disciplinary research to create fundamental knowledge and applications of biobased products. He is a highly cited researcher (top 1% by citations, Clarivate) and recognized as a worldwide leader in the area of renewable nanomaterials. This is a result of the efforts of the talented members of his Bio-

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compared to zeolites or MOFs. This has prompted new strategies for controlling the material pore structure and surface interactions of porous carbon, critical to adsorption performance.

In this context, lignin has emerged as a promising precursor of solid porous adsorbents. Lignin is the most abundant aromatic biopolymer found in nature, accounting for up to 10–35% by weight and 40% by energy of plant biomass. ²⁶ Lignin-containing by-products from industry represent a viable source of this biopolymer, which can be isolated downstream through fractionation processes. In such conditions, industrial lignin is produced in large quantities (~ 50 –70 million tons annually), ²⁷ and used primarily as a low-value biofuel through combustion that enables chemical recovery after wood pulping.

Lignin's aromatic structure endows it with a high carbon yield (~40%, the highest of all biopolymers), while its functionalities offer various pathways for physical and chemical modification. In fact, many of the alternative resources selected for the production of activated carbon (e.g., coconut shells, wood) are actually lignin-rich, endowing them with higher carbon yields. Indeed, using lignin alone may be a more efficient approach. Further, there are pathways for exploiting the polymeric structure of lignin for strategic upgrading strategies that are not possible for other feedstocks that are complex mixtures of several biopolymers and biomolecules. Therefore, lignin has great potential as a major feedstock for producing renewable activated carbons. The use of lignin in material applications, particularly in carbon capture technologies, sequesters biogenic carbon, providing industries with a viable path toward achieving net-zero—and potentially even net-negative—CO2 emissions.

Despite decades of efforts from various stakeholders, the full potential of lignin has yet to be fully realized on the industrial scale, primarily due to the inherent composition variability and chemical complexity. However, owing to its inherent advantages, there are major incentives to find cost-effective and scalable methods that adhere to green chemistry principles for the commercialization of lignin-based carbon capture technologies. In this review, we offer a perspective on lignin-based solid gas adsorbents for CO2 capture. We discuss lignin sources and availability, its structure and chemistry, as well as the strategies employed for lignin deployment in the field. A review of the key metrics for the properties and performance of ligninderived carbon materials is included, which is critical for developing structure-property-performance relationships. We also describe the traditional and emerging methods for processing lignin into porous materials designed for CO₂ capture. We provide critical viewpoints and identify key gaps in innovation. Finally, we discuss the prospects of lignin as a technological alternative for CO2 capture and outline future directions for research and development.

2. Lignin chemistry and properties

The term "lignin" encompasses a broad class of biomacromolecules exhibiting significant variability in their chemistry, structure, bonding patterns, and properties.²⁸ Special consideration of lignin's origins and structure is essential for processing it into value-added materials; lignin-based carbon adsorbents are no exception. In this section, we review the origins of lignin, its chemical structure, and relevant definitions. The discussion then covers lignin's thermal properties and current processing strategies. Finally, we address the sustainability aspects associated with the utilization of technical lignins in the context of the main processing routes.

2.1 Lignin origins and resources

Originating roughly 400 million years ago as land plants adapted to terrestrial ecosystems - lignin is a class of polyaromatic compounds mainly derived from three canonical monolignols, p-coumaryl alcohol, coniferyl alcohol, and sinapyl alcohol, which differentiate by the number of ortho methoxy groups on the phenolic ring.²⁹ These fundamental building blocks are synthesized by plant cells in the cytoplasm from phenylalanine and transported to the cell walls, where they undergo chemically controlled free radical coupling reactions to form polymeric constructs. 30 The monolignols are therefore monomeric units of the lignin polymer and are referred to as p-hydroxyphenyl (H), guaiacyl (G), and syringyl (S), respectively. Due to the absence of biochemical control dictating polymer formation and the high functionality of its monomer precursors, lignin is a highly complex, three-dimensional random polymer with a variety of linkages and structures (Fig. 1) including covalent linkages to the polysaccharide framework of the cell wall. Further, lignin variation according to the plant source and the plasticity of the lignin biosynthetic pathways result in high compositional and macromolecular variability.³¹ Hence, there is no strict definition of lignin, and there are still ongoing critical discussions on key elements of its structure and definition.

A distinction should be made between native lignins, and lignins that have been extracted from biomass, termed technical lignins or those that remain attached to other polymeric structures, namely, residual lignins. Technical lignins are of practical relevance and have a significantly altered structure compared to their native counterparts. The former distinguish from the latter as far as its (i) reduced primary and secondary aliphatic hydroxy groups, (ii) reduced oxygenated aliphatic moieties, such as the β-O-4, and (iii) higher phenolic hydroxy and ester groups, saturated aliphatic moieties, and degree of condensation.33 The extent of these differences can vary significantly depending on processing history. Technical lignins can be bound to carbohydrate units, the so-called lignin-carbohydrate complexes (LCCs), especially lignosulfonates and hydrolysis lignins. These complexes include both covalent and non-covalent bonding between lignin and carbohydrates, and therefore, their separation and/or characterization introduce further challenges. Depending on the dissolution and fractionation processes used, technical lignins can be modified to include heteroatoms. For example, kraft lignin contains small amounts of organically bound sulfur (thiol compounds).³⁴ Further, a significant number of native linkages are cleaved and the propyl side chain on the lignin

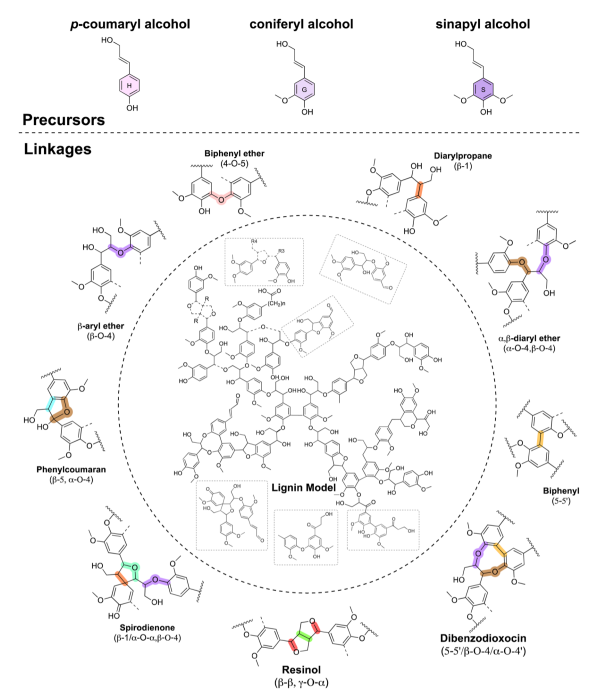


Fig. 1 Main lignin precursors (top), typical inter-unit linkages and proposed structural model macromolecule corresponding to spruce milled-wood lignin. Some of the elements shown in this illustration are inspired by ref. 32. Adapted under terms of the CC-BY 4.0 license.³² Copyright 2020, Royal Society of Chemistry.

subunits is modified, resulting in a change of chemical functionality, new linkages, and a new polymer chain architecture.

According to the plant source, three main categories of lignin exist: softwood lignins, hardwood lignins, and herbaceous lignins. Softwood lignins are primarily composed of G-type subunits and possess less compositional variability. In comparison, hardwood lignins have varying proportions of S and G units (*i.e.*, a diverse S/G ratio), whereas herbaceous lignins are considered to be a complex mixture of H-, S-, and G-types.

In terms of processing differences, technical lignins are primarily produced by the pulp and paper industry during the chemical digestion of biomass, where lignin is dissolved to liberate the cellulosic fibers. Kraft lignins and lignosulfonates, which originate from the kraft and sulfite pulping processes, respectively, are commercially available. Lignosulfonates are a special case of technical lignins with different behavior relative to other technical lignins. The chemistry of the sulfite pulping process results in the production of highly sulfonated and

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amphiphilic lignin, which is soluble in water at neutral pH and possesses enhanced surface activity. Due to the significant differences of lignosulfonates with other technical lignins, in this review, it will be specified clearly when discussing them. Despite sulfite pulping being much less common than kraft, lignosulfonates have been the main source of lignin-derived commercial products.35 However, the kraft pulping process can afford partial lignin removal without offsetting the chemical and energy recovery cycle. This can be done by implementing separation processes from the so-called black liquor by the LignoBoost and the LignoForce process.36-38 As a result, the Kraft lignin market has the potential to surpass that of lignosulfonates in the near future.³⁹

Beyond traditional pulping processes, there have been significant developments in the area of biorefining processes that incorporate organic solvent extraction, ionic liquids, acidhydrolysis, and steam explosion. 40-43 The obtained lignins may comprise new categories since these alternative processes differ significantly from established pulping systems. Hence, the term "lignin" actually describes a broad category of macromolecules and special attention should be dedicated to clearly identifying their nature or origins and processing conditions (e.g., botanical source and industrial process). Even lignins that originate from the same botanical source and industrial process possess high variability, which is not easily quantified due to their complexity.

2.2 Lignin thermal properties

To date, there is still no systematic understanding of the direct relationship between lignin's chemical and structural features and the corresponding thermal properties. Yet, thermal processing is critical for producing solid-porous adsorbents from lignin. Pyrolysis, the thermal decomposition of a material in the absence of oxygen, is a common lignin treatment method. Under inert conditions, lignin is decomposed at high temperatures, with no combustion, leading to a high carbon yield. This process and others have been discussed in a comprehensive review,⁴⁴ which proposes lignin pyrolysis occurs in three main stages: dehydration ($\sim 30-200$ °C), active pyrolysis (\sim 200–450 °C), and passive pyrolysis (>450 °C) (Fig. 2). In the dehydration stage, moisture and volatiles are released, although it should be noted that significant rearrangement of the lignin structure will also occur below 200 °C. Above 200 °C, decomposition proceeds more rapidly, first cleaving weak ether bonds and eventually condensed C-C bonds, producing depolymerized primary products, including monomeric fragments. In this stage, roughly 40 wt% of the material is lost in gaseous forms. Above ca. 450 °C, the decomposition mechanism transitions to a gradual passive stage dominated by secondary degradation and transformation, including the cleavage or rearrangement of functional groups and depolymerization of carbon species. The pyrolysis degradation mechanism is very complex and includes various reactions and physiochemical phenomena - dependent on both the nature of the lignin structure, processing parameters (such as heating rate and final temperature), and their interactions. Unless otherwise stated, in this review, we consider carbon production from lignin by pyrolysis, which is most often performed under a nitrogen (N_2) atmosphere.

Various thermal analyses can be used to investigate the thermal properties of lignin. Typical approaches include thermogravimetric analysis (TGA) and differential scanning calorimetry (DSC). Along with a decomposition profile, TGA provides information regarding moisture content, volatile matter, fixed carbon, and ash content, thus serving as a valuable tool to guide thermal processing regimes and estimate carbon yield. DSC provides thermodynamic information such as phase transitions, reaction temperatures, and, importantly, the glass transition temperature (T_g) - a critical property for stabilizing the given structures during carbonization.

Kraft lignins are mostly degraded in the range of 250-450 °C (roughly 40 wt%), after which degradation begins to plateau. Most kraft lignins have a relatively low glass transition temperature, between 100 and 170 °C.45 Often, a goal prior to/ during thermal processing is to increase the glass transition temperature of lignin, to facilitate direct decomposition instead of melting. The former mechanism, referred to as thermal stabilization (or oxidative pretreatment), preserves structures such as pores or hierarchical assemblies. Heating lignin at low to moderate temperatures in the presence of oxygen is an effective means to improve the thermal stability of lignin. Not all lignins are equal; in particular, thermostability strongly depends on lignin type. Lignosulfonates have higher molecular weights than most other technical lignins (M_w 10 000-40 000 Da)

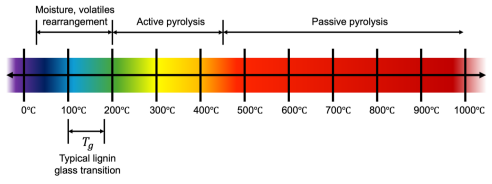


Fig. 2 Key regions for the thermal decomposition and glass transition of typical lignin streams.

and therefore may preserve the inherent morphologies and pore structures at small scales.46 Also, the fractionation of lignin impacts its glass transition, and this can be exploited for thermal stabilization.47,48

Processing of technical lignins

In contrast to tailorable synthetic polymers with uniform structure, the heterogeneity of lignin is a major challenge in processing. Therefore, different strategies have been developed to produce lignin streams with less heterogeneity and selected features, which can be classified mainly as fractionation and assembly (Fig. 3).

Lignins are separated into more chemically uniform streams by using solvent fractionation, ultrafiltration, and/or pH shifting. Increasing lignin uniformity through fractionation may be necessary for developing solid-porous adsorbents from lignin, with tailorable and ordered pore structures. Fractionation is effective in producing fractions of (relatively) narrow molecular weight distribution. Secondary effects of fractionation lead to lignin sub-streams with chemical gradients between each other - a consequence of the inextricably linked molecular weight and functionality-type distributions inherent to lignins. 49 Larger molecular weight fractions typically contain more aliphatic and less phenolic and carboxylic hydroxy groups; differences in the inter-unit linkages among fractions are also typical. Collectively, the solubility of lignin is greatly impacted, together with properties such as thermal stability, reactivity and surface energy/wettability.50

Alternatively, assembly has been applied to transform polymeric lignin into spherical colloidal particles of tailorable size (1 nm-10 μm) and surface charge. They are readily produced in wet or dry forms through solvent exchange, spray-drying, or aerosol flow processes.51 Despite the broad molecular weight distribution and functionality of the starting lignin profile, remarkably narrow particle size distribution is attainable within assembled lignin structures. The degree of uniformity can be further improved with particle size fractionation, offering avenues for highly ordered structures. 52,53

Hierarchical assembly and chemical modification can be applied to tailor both the molecular- and macro-scale properties of lignin-based materials. Lignin supraparticles, which display hierarchical porous structures, have been shown for their potential for CO₂ capture.⁵⁴ Controlled porosity at multiple length scales facilitates molecular and gas transport, producing materials with macrostructures that meet the technical requirements for scaled processes.

The abundant functional groups of lignin, including aliphatic, phenolic, carboxyl hydroxy, and methoxy groups, as well as ether bonds, offer convenient possibilities for chemical modification and to impart properties matching given applications (Fig. 4).55 Chemical modification can take the form of depolymerization, modification of its functional groups, or graft copolymerization.⁵⁶ Depolymerization of lignin macromolecules to platform chemicals enables bottom-up approaches for materials synthesis, vet is not a trivial task due to its structural complexity and recalcitrant C-C and C-O bonds. 57 Functional group modification can include the synthesis of new active sites or the modification of the present hydroxy groups.⁵⁸ Graft copolymerization can be performed to attach new polymer chains to the lignin structure via its reactive hydroxy groups.

While the processing strategies covered in this section have been discussed individually, more sophisticated efforts involve the combination of two or more of these strategies.⁵⁹ The compositional profiles (and associated properties) of the derived lignins must be considered for its impact on the final product. This can be addressed through early-stage technoeconomic analyses, which are recommended to identify viable pathways for valorization.

2.4 Sustainability aspects relevant to technical lignin valorization

Emerging carbon capture technologies must be critically evaluated in terms of their sustainability. Ideally, sustainable solutions should consider the entire environmental impact over their full life cycle, from raw material sourcing to recycling or disposal. While bio-based materials provide renewable carbon sources, they must also meet these sustainability standards. In this context, standardized life cycle assessments (LCAs) are essential.60

Generally, lignin valorization approaches have the potential to reduce the environmental impact of kraft pulp mills by decreasing stack emissions from the recovery boiler.61 The utilization of lignin for carbon capture applications further

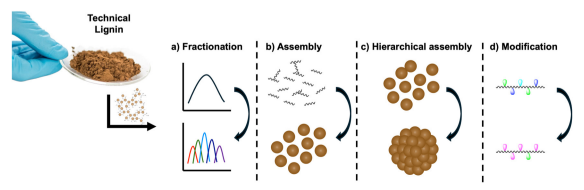


Fig. 3 Typical lignin processing strategies include (a) fractionation, (b) assembly, (c) hierarchical assembly, and (d) chemical modification.

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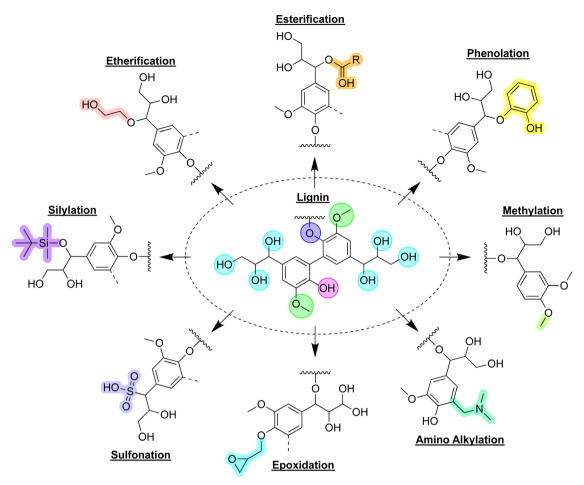


Fig. 4 Lignin derivatization pathways: the central panel depicts a segment of a lignin macromolecule, surrounded by various derivatives corresponding to specific reactions, as indicated

expands the potential for reducing net CO2 emissions and the associated environmental impacts of lignin valorization. A critical assessment of 42 peer-reviewed LCAs regarding lignin and lignin-derived products found that they often offer better environmental performance compared to their fossil-based counterparts, particularly concerning their impact on climate change.⁶² Moreover, when considering the displacement (substitution) of petroleum-derived products by lignin, this impact can be even more significant. Life cycle assessments involve assumptions, boundary conditions, and standardized methods that must be taken into account when interpreting their results, as these factors can significantly influence decision-making.⁶²

In Section 2.3, we introduced various techniques typical for technical lignin processing aimed at reducing heterogeneity and tailoring its properties. While these approaches have gained popularity in the literature, they should be approached with caution. Excessive or redundant processing steps can significantly increase the complexity of the system, as well as capital and operational costs, and may also heighten environmental impacts.

Chemical modification of lignin must adhere to the principles of green chemistry and sustainability. 63,64 Molecular fractionation of lignin can be employed to reveal structure-property

relationships. However, if not properly integrated, these schemes can become complex and expensive, often providing only incremental improvement in lignin uniformity and properties. A practical approach is assembling lignin into spherical particles, which addresses multiple technical challenges and produces relatively homogeneous and structured materials. However, in the most common methods for obtaining colloidal lignin particles—such as pH shift and the use of antisolvents—considerable expenses arise due to the large volumes of organic and aqueous solvents (used for solvent exchange) and the energy-intensive drying processes.

In Section 4 to follow, we will discuss the processing of lignin to porous carbon adsorbent materials. These operations are often energy and chemical-intensive. Thermal processing incurs a significant energy penalty during the carbonization and activation steps. Chemical activation, which is a common approach because of its simplicity and efficacy, requires the use of an activation reagent typically at amounts of at least twice that of the carbon source by mass. Further, a washing step is typically necessary for post-treatment to remove the remaining activation agent and expose the formed pore structure. Assembling porous structures prior to carbonization are promising approaches to bypass or reduce the requirement for chemical

activation and associated washing steps, but this is often at the cost of increased processing time, complexity, and energy for procedures such as thermal stabilization - often required to maintain those structures at the high temperatures necessary for carbonization.

In lignin valorization efforts, processing steps often drive costs and environmental impact, ultimately affecting sustainability. However, in the context of carbon capture technologies, the sustainability of lignin-based adsorbents will critically depend on their adsorption performance. In the following section, we discuss the key performance indicators for ligninbased carbon adsorbents, which ultimately determine the success of these emerging technologies.

3. Properties and performance of lignin-based carbon adsorbents

The performance indicators of lignin-based carbon adsorbents are essential for advancing their commercial-scale implementation in adsorption systems. These indicators encompass both material properties and adsorption performance, which are interrelated. Regarding material properties, standard characterization of carbon materials typically involves assessing the specific surface area, pore structure, surface chemistry, purity, and degree of graphitization. Adsorption capacity, selectivity, and kinetics are key factors that influence adsorption performance.

One of the key challenges for the scaleup adoption of solid adsorbents is the lack of standardization in the characterization and evaluation. To adequately compare various ligninbased adsorbents and benchmark them against competing technologies, it is essential to align experimental methods and subsequent analyses. Therefore, this section will first introduce the properties of lignin-based carbons, which dictate adsorption performance, with a focus on methods for their determination. Subsequently, we will discuss adsorption performance in terms of the most critical performance indicators.

3.1 Carbon properties

The specific surface area of a solid material can be determined using gas physisorption experiments based on equilibrium van der Waals interactions between the gas molecules and the solid surface.65 The BET (Brunauer-Emmett-Teller) theory is the most common approach to obtain the specific surface area (SSA) of carbonized lignin.⁶⁶ Adsorption isotherms can also be used to calculate pore size distributions. Extreme caution is necessary for applying the BET theory to materials with a high degree of microporosity.⁶⁷ Further, the choice of adsorptive is also of critical importance, adsorption of CO2 at 273 K (0 °C) has become the accepted method for carbonaceous materials with very narrow micropores.⁶⁷ The influence of porosity on CO₂ uptake will be discussed in Section 5.1.1.

Activated carbons typically contain disordered arrangements of aromatic carbon structures that stack to create local order.⁶⁸ Their degree of graphitization is therefore dependent on many

factors of the precursor material and thermal processing. Carbonizing lignin produces hard carbon, which is a disordered material that is not easily graphitized, even at very high temperatures (>1500 °C). 69 The degree of graphitization or order can provide insights into the structure and performance of carbon materials. Powder X-ray diffraction (XRD) is a typical approach to gain insights into the order of the carbon and can be used to estimate the degree of graphitization by comparing peak intensities of the (002) and (100) reflections. Raman spectroscopy can also be utilized to provide information on order and crystallite or domain size. A stretching vibration in aromatic layers is known as the G (graphite) band $\sim 1580 \text{ cm}^{-1}$, which can be compared to the D (disorder or defect) band ~ 1350 cm⁻¹ attributed to the breakdown of translational and local lattice symmetries.70

Carbon materials derived from technical lignins can also exhibit diverse surface chemistries, yet they all share in common considerable sp²-hybridized carbons with two-dimensional order in addition to many heteroatoms (mainly oxygen).71 Minor differences in surface chemistry can play an important role in dictating the adsorbent-adsorbate interactions, and therefore, characterization of these groups is critical. Common approaches to identify and attempt to quantify these groups include spectroscopy/spectrometry techniques such as Infrared spectroscopy (IR), Raman, X-ray photoelectron spectroscopy (XPS), and temperature-programmed desorption (TPD). Boehm titration can also be employed, with caution, to measure the concentration of functional groups on the carbon surface.⁷²

Other important material properties include their structural topology, thermal stability, and mechanical stability. The structural topology can be evaluated with microscopy, in particular, electron microscopy is typical for evaluating the structural features at smaller length scales. Thermal stability, of which can be decisive under the various operating conditions necessary considering feed-gas temperatures and regeneration, is typically evaluated with TGA. Mechanical stability of structured adsorbents is essential for determining their suitability for withstanding compression forces in packed bed systems and can be evaluated with compression tests, for example with dynamic mechanical analysis.54

3.2 Adsorption performance

Adsorption capacity (equilibrium capacity and working capacity) is a primary performance indicator for solid gas adsorbents. Equilibrium capacity refers to the amount of CO2 adsorbed at equilibrium under specified conditions, while working capacity represents the amount of CO2 captured during a complete adsorption/desorption cycle, calculated as the difference in capacity between adsorption and desorption conditions.73 Working capacity is more relevant at the industrial scale; however, equilibrium adsorption capacity is often the only metric reported in the literature, particularly for adsorbents associated with low technology readiness levels (TRLs). Equilibrium adsorption capacity is typically determined using volumetric methods (gas adsorption isotherms) or gravimetric methods (TGA).

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Dynamic column breakthrough experiments are important indicators of adsorbent performance. The breakthrough response of an adsorption column refers to the signal observed at the column's outlet stream resulting from a step change in the concentration of one or more adsorbate components.⁷⁴ Notably, breakthrough experiments provide both single- and multi-component equilibrium information, in addition to enabling the characterization of adsorption kinetics. Dynamic column breakthrough experiments are not widely reported, particularly in material development labs that do not specialize in adsorption systems, where sample production is often limited to the milligram scale. Advances in microscale dynamic column breakthrough measurements may accelerate the screening of solid adsorbents, especially from materials that exhibit high variability, such as lignin.⁷⁵

Additional indicators of the performance of solid adsorbents include selectivity, heat of adsorption, and their recyclability/ regeneration. In real processes, feed gas streams consist of multicomponent mixtures that compete for adsorption on the surfaces of the adsorbent. Therefore, evaluating CO₂ selectivity is essential.⁷⁶ Of the various models proposed for predicting multi-component gas adsorption equilibria from purecomponent isotherms, the Ideal Adsorbed Solution Theory (IAST) has remained a preferred choice due to its simplicity and reliability. 77,78 The isosteric enthalpy of adsorption—defined as the heat released upon binding to a surface—governs the local adsorbent temperature and, consequently, the local adsorption equilibria, kinetics, and resulting separation efficiency.⁷⁹ In particular, a large isosteric enthalpy of adsorption can delay equilibrium and indicate stronger adsorbent-adsorbate interactions, which can increase the costs associated with adsorbent regeneration.80-82 The isosteric enthalpy of adsorption is typically determined indirectly by applying the Clausius-Clapeyron equation to adsorption isotherms obtained from volumetric gas adsorption measurements.83 To assess the regeneration stability (mechanical, chemical, and thermal stability) of the adsorbent, cycling experiments under the relevant operating conditions are performed.

4. Processing of lignin-based porous materials

There are several methods for producing solid porous adsorbents from lignin. Traditional approaches involve two main processes: carbonization and activation, which can be carried out either simultaneously in a one-step process or sequentially in a two-step process. In carbonization, thermal treatment is applied to induce a complex reaction that results in a material with a higher carbon content. Thermal treatment is most often performed *via* pyrolysis, while hydrothermal carbonization and microwave treatment are also promising alternatives. In contrast to carbonization, activation aims to introduce or augment the porosity of the material, creating cavities, which facilitate gas transport or create sites for molecular adsorption. This is typically achieved *via* chemical or physical activation. Chemical

activation creates pores by reacting the material with an acid, base, or salt at elevated temperatures. In contrast, physical activation uses a reactive gas environment at high temperatures to etch the material's surface.

The above-mentioned approaches are effective in producing highly porous lignin-based materials but lack control over pore sizes, leading to disordered morphologies and non-uniform pore-size distributions. However, ordered porosity is known to promote molecular transport and adsorption.⁸⁴ Thus, recent approaches have advanced morphology control at multiple scales, reducing or eliminating the severity of the activation steps. Hierarchically structured materials are those with welldefined and reliably controlled combinations of structural features at various scales and typically exhibit multiple advantages compared to their more disordered counterparts. In particular, control over macroscopic dimensions has technical advantages for industrial adoption, for example, in the case of packed columns typical for adsorption processes, where powdery material can induce undesired pressure drop due to insufficient void volume for gas transport. In terms of pore structure, mesopores (2–50 nm) facilitate gas transport and molecular diffusion, forming scaffolds for and access to micropores (<2 nm), which in turn greatly enhance the surface area, offering adsorption sites and molecular sieving capacity. Ultramicropores (<0.7 nm) are critical in determining the dimensions of the binding sites and provide molecular sieving function.

In the following section, we examine lignin processing into solid porous ${\rm CO_2}$ adsorbents and the various strategies used to increase the apparent surface area and optimize porous structure at multiple length scales. We begin with traditional processing to produce irregular porous carbon, simple one-step carbonization, and multi-step or hybrid techniques with various carbonization/activation/doping schemes. Subsequently, we cover 3D networked structures produced by crosslinking chemistries. Finally, we examine the approaches for hierarchically structured materials with more sophisticated designs and greater control over the multi-scale pore structure and macroscopic dimensions of the material, through the formation of superstructures or by hard or soft templating.

4.1 Traditional approaches to CO₂ adsorbents

4.1.1 One-step carbonization/chemical activation. The most convenient approach for producing activated carbon from lignin is a one-step process that simultaneously combines carbonization and activation. Here, lignin is pre-mixed with an activation agent. During carbonization, a pore structure is developed via two simultaneous reactions: (i) the degradation of lignin and the activation agent, and (ii) the reactions between lignin and the added reagents. Carbonization of low-grade lignin streams in one step, harnesses the activation effect of the salts already present in the sourced lignin. Cao *et al.* obtained lignin-based biochar from high ash content ($\sim 46\%$) alkali lignin via pyrolysis at 750 °C in a muffle furnace for 3 hours. The biochar exhibited a maximum CO_2 adsorption capacity at 0 °C of ~ 4 mmol g^{-1} with a specific surface area of 1134 m² g^{-1} , total pore volume of 0.84 cm³ g^{-1} and micropore

volume of 0.49 cm³ g⁻¹. The authors demonstrated the effective role of acid washing with a short ultrasonic treatment time in a post-treatment stage, which significantly increased CO2 adsorption by improving the material purity and unobstructing the pore structure.

Pan *et al.* investigated the effect of pressure during pyrolysis of high ash-content alkali lignin.86 They carbonized lignin at 10 °C min⁻¹ to 800 °C for 3 hours and under three different pressures: negative pressure (-0.1 MPa), atmospheric pressure (0 MPa), and elevated pressure (0.1 MPa). It was found that CO₂ adsorption capacity, specific surface area, and micropore volume positively correlated with lower pressure during carbonization, with a trade-off in carbon yield. The optimized carbon material obtained under negative pressure had a specific surface area of 1577 m² g⁻¹ and micropore volume of 0.695 cm³ g⁻¹, leading to a CO₂ adsorption capacity of ~ 3.6 mmol g⁻¹ (at 0 °C). The vacuum reduced the boiling point of volatiles and reduced vapor residence time, facilitating their removal and minimizing undesired secondary reactions during pyrolysis.

The impurities of high ash-content lignin are resourcedependent and unpredictable, limiting the applicability of one-step carbonization protocols and compromising quality control over the activation process. More consistent carbon production can be achieved by adding activation agents to purified lignin. Sun et al. used a one-step procedure by soaking corn straw lignin in 2:1 w/w phosphoric acid (H₃PO₄), followed by heating at 30 $^{\circ}$ C min⁻¹ to 300–600 $^{\circ}$ C for 120 min.⁸⁷ Mesopores were formed below 500 °C with the presence of H₃PO₄, after which the pore volume decreased sharply, potentially due to the decomposition of phosphorous esters. Their approach could produce specific surface areas and pore volumes of 820 m² g⁻¹ and 0.8 cm³ g⁻¹, respectively. However, the dynamic adsorption performance in a packed bed reactor indicated a relatively poor separation performance of the asproduced carbon pellet, despite a promising selectivity between CO2 and methane (CH4) from equilibrium adsorption results. The authors suggest that further modification of the functional groups on the carbon surface is necessary to improve the selectivity towards CO₂.

Li et al. utilized the one-step method by using wet impregnation with KOH solutions at a 1:1 or 2:1 w/w activation ratio to process kraft lignin.88 After drying, the mixtures were carbonized at a rate of 5 °C min⁻¹ to 600, 700, or 800 °C and holding for one hour. Regardless of the activation ratio or final temperature, the resulting carbons primarily possessed microporous structures (84-93%). Mild carbonization/activation conditions produced lignin-based adsorbents that favor adsorption at lower CO2 partial pressures, while those from high temperatures and high activation ratio conditions performed better at high CO₂ partial pressures. In general, CO₂ uptake depends on ultra-microporosity at low partial pressures, while total micropore volume is more important at high CO2 partial pressures. For this work, the uptake capacities can go up to 2 mmol g^{-1} (at 25 °C) at low CO2 partial pressures, whereas the CO2/N2 selectivity can reach 38 at 15 kPa CO₂. Importantly, Li et al. provide strong evidence that KOH activation can leave intercalated potassium (K+) ions under the carbon surface that are not leached by washing in water. These ions can significantly increase favorable interactions with CO₂, improving uptake by up to 50%.

In the above cases, technical lignins (such as kraft lignins) already have abundant oxygen-containing functional groups that play a role in the surface interactions with CO₂, although the impact on CO2 adsorption is still a topic that remains unclear.89 Aside from harnessing the heteroatoms within the lignin, doping is the most common strategy to produce carbon materials from lignin with favorable surface chemistry for CO₂ adsorption. Heteroatoms, such as nitrogen or sulfur, can increase the affinity to CO2. In the discussion that follows, it is also shown that the surface chemistry is shown to be adjustable according to the technical lignin type, i.e., its functional groups (for example in the case of lignosulfonate or enzymatic hydrolysis lignins), which contain varying quantities of sulfur or nitrogen, respectively.

Saha et al. extended the one-step approach to include Sdoping by combining lignin, KOH, and sodium thiosulfate (Na₂S₂O₃), followed by pyrolysis at 10 °C min⁻¹ to 800 °C and holding time of 2 minutes. 90 A high surface area was generated through activation, while simultaneously introducing sulfur content (1-12.6%) to the carbon surface to improve the surface interactions with CO2. The highest surface area (surface area and pore volume of 3626 m² g⁻¹ and 1.7 cm³ g⁻¹, respectively) was achieved from the largest ratio of KOH and the lowest ratio of Na₂S₂O₃, attributed to a large percentage of microporosity and, consequently, the best CO₂ adsorption capacity (11 mmol g⁻¹ at 25 °C).

N-doping has become popular in the literature, since it can significantly promote the Lewis acid-base interactions. Tkachenko et al. combined nitric acid (as the chemical activator) with urea (as the N-doping agent) to produce N-doped activated carbon from softwood kraft lignin (Fig. 5).⁹¹ In their synthesis, lignin was combined with urea and nitric acid (300 mg urea/ 2.5 mL 40% Nitric acid/1 g lignin) and heated at 6 $^{\circ}$ C min $^{-1}$ to 800 °C. The resulting carbon material was significantly microporous with 3.5% N content and achieved a specific surface area of up to 1000 m² g⁻¹. The CO₂ adsorption capacity boosted to 1.4 mmol g^{-1} at 15 kPa and 20 °C.

4.1.2 Hybrid carbonization, chemical and physical activation. In the typical one-step method (as shown in Section 4.1.1), chemical activation has been established as an effective means to improve porosity. However, a high activation ratio is often necessary to reach the desired high surface areas. Large amounts of activation agents drive up costs, including that of the agents themselves, as well as costs associated with the subsequent washing steps. To address such issues, physical activation is a common alternative to replace or reduce the chemical activation agent. Further, since physical activation is performed using gas, it can be implemented into existing onestep procedures simply by switching gases from the inert pyrolytic atmosphere to the physical agent of choice.

Chen et al. used microwave treatment and combined physical/chemical activation with humidified nitrogen and KOH to prepare carbon from enzymatic hydrolysis lignin (Fig. 6).92 **Review Article** Chem Soc Rev

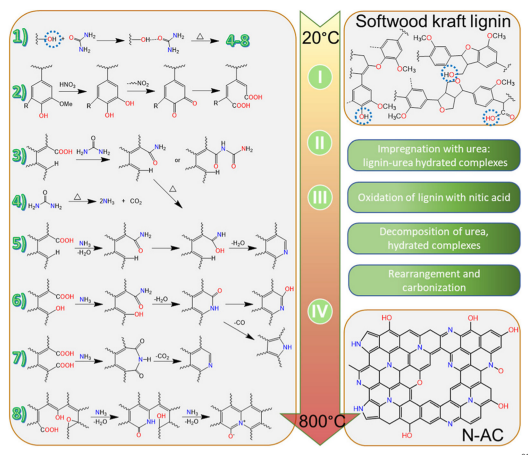


Fig. 5 Schematic illustration of the transformation of lignin to N-doped carbon. Reproduced under terms of the CC-BY 4.0 license. 91 Copyright 2024, American Chemical Society.

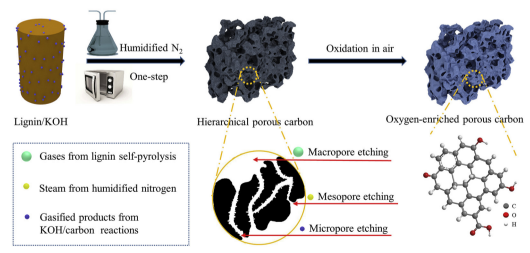


Fig. 6 Schematic illustration of the one-step microwave synthesis in the presence of humidified nitrogen and oxygen-enriched porous carbon. Reproduced with permission. 92 Copyright 2019, Elsevier.

Microwave heating can be advantageous because of its low energy consumption, short heating duration, and uniform heating gradient. Lignin powder was mixed with KOH at a 1:3 ratio and dried, then placed in a microwave under a humidified nitrogen atmosphere. The humidified nitrogen

could act not only as a microwave absorber to increase temperature, but also, with steam, as a physical activator. In an optimum processing condition (30-minute microwaving), the porous carbon showed a specific surface area of 2870 m² g⁻¹ and 2.02 cm3 g-1 pore volume, associated with high oxygen

content. Collectively, these properties enable CO2 uptake of 1.31 mmol g⁻¹ at 30 °C. Most remarkably, the rapid carboniza-

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tion time during microwave heating is the major advantage of this approach.

In another approach, Saha et al. synthesized a carbon with hierarchical porosity from lignin via a hybrid one-step chemical activation, followed by physical activation/nitrogen doping with ammonia (NH₃).93 Lignin was first mixed at a 1:1 ratio with KOH, pyrolyzed at 10 °C min⁻¹ to 800 °C, and held for two minutes before washing and drying. The dried carbon product was later physically activated with NH₃ at 800 °C. The NH₃induced physical activation increases the surface area and pore volume, as well as grafted a variety of nitrogen groups on the carbon scaffold, such as pyridinic, amino (combined primary, secondary, tertiary), pyrrolic (in combination with pyridone), graphitic (or quaternary amine), nitro and nitroso. It is noteworthy that the physical activation process would reduce the carbon yield and cause undesired over-activation effects if not optimized. Nevertheless, CO2 selectivity increased monotonically with nitrogen content, indicating that N-doping can improve the CO₂ selectivity of carbon materials. Surface area was not directly correlated with selectivity towards CO2 over N2. Their method achieved specific surface areas of up to 2922 m² g⁻¹ and CO₂ adsorption capacities of 5.5 and 8.6 mmol g^{-1} at 25 °C and 0 °C, respectively.

4.1.3 Two-step carbonization/activation. A typical two-step approach separates the degradation and porosity development steps. Compared to the one-step approach, it features better control over pore formation and reduced dosage of the activation agent. Therefore, certain hierarchical porosities may only be attainable through this approach. Further, the two-step approach can significantly reduce the requirement of activation chemicals. However, two-step carbonization may suffer from the significant time and energy requirements of multiple thermal treatments.

Slow pyrolysis is the typical initial step in the two-step approach. Gong and Bao investigated a two-step carbonization and activation process to produce porous carbon from lignin.⁹⁴ They first pre-carbonized the lignin at a heating rate of 5 °C min⁻¹ to three different temperatures (350, 450, 550 °C) for 2 hours. The resulting char was ground with KOH at ratios 1:1, 1:2, and 1:3, followed by activations at 5 °C min⁻¹ to 600 °C or 800 °C under N₂. The authors found that a lower final activation temperature favored the development of pores. Their approach could achieve gas adsorption performance of 3.98 and 5.82 mmol g^{-1} at 100 kPa, 25 °C and 0 °C, respectively.

Hydrothermal carbonization is a promising alternative for the thermal conversion of wet organic streams such as industrial pulping residues, especially since the vast majority of technical lignins are extracted as a soluble residue in the form of black or brown liquor. The hydrothermal process can effectively avoid the energy-intensive drying and demineralization steps. Interestingly, the concept of hydrothermal processing was first conceived as a simulation of coal formation. 95,96 In such a process, biomass is subjected to an aqueous treatment at low to moderate temperatures (\sim 180–250 °C), whereby

various chemical reactions (e.g., hydrolysis, dehydration, decarboxylation, polymerization, aromatization, condensation) result in a solid carbonaceous material.⁹⁷ While the carbon product of biomass pyrolysis is commonly known as biochar, the product of hydrothermal carbonization is known as "hydrochar", as will be denoted in the following paragraphs.

Sangchoom et al. synthesized porous carbon from lignin without demineralization via a two-step process. 98 Their approach involved hydrothermal carbonization at 300-390 °C, followed by chemical activation with KOH at ratios 4:1 or 2:1 *via* heating at a rate of 3 $^{\circ}$ C min⁻¹ to 600–900 $^{\circ}$ C and holding for 1 hour. While the original lignin contained a 20 wt% noncombustible ash content, the hydrochar contained only organic material without minerals. This result indicates that the hydrothermal carbonization successfully subverted the demineralization step. Remarkably, the as-produced carbon products achieved high surface area and pore volumes of 1157-3235 m² g⁻¹ and 0.59-1.77 cm³ g⁻¹, respectively, allowing excellent CO₂ adsorption capacity of 4.6 mmol g⁻¹ at 25 °C and 100 kPa, 17.3 mmol g⁻¹ at 25 $^{\circ}$ C and 2000 kPa, and 7.4 mmol g^{-1} at 0 $^{\circ}$ C and 100 kPa. The authors denoted that pore size, particularly micropores ~ 7 Å, is key for CO2 uptake at ambient conditions.

Hydrothermal carbonization can be used to produce hydrochar but can also be modified for other objectives, such as the co-production of bio-oils.99 Hao et al. used the solid byproducts from the conversion of lignin to bio-oil, and incidentally, they produced a magnetic-activated carbon from eucalyptus enzymatic hydrolysis lignin and softwood kraft lignin. 100 These hydrochars were considered as a side product of a ligninto-liquid process when performing hydrothermal treatment together with formic acid. Briefly, 200 g of lignin, 200 mL of formic acid, 500 g of water, and the catalyst were added to a 5 L stainless-steel reactor. A temperature of 380 °C was used to produce hydrochars from the kraft lignin from Norway spruce and 365 °C when using the lignin derived by enzymatic hydrolysis of eucalyptus biomass. The as-produced biochars were mixed with KOH solutions and heated for 5 hours under 200 °C, followed by activation at 700 °C or 800 °C for 4 hours. The magnetic properties of the porous carbon were likely a consequence of corrosion of the stainless-steel reactor during hydrothermal treatment with formic acid, resulting in hydrochars of up to 12.3% iron by weight. These lignin-based magnetic carbons exhibited high surface area and adsorption capacities (up to 2975 m² g⁻¹ and 6 mmol g⁻¹ at 101 kPa and 0 $^{\circ}$ C), but it is perhaps also a good example of the potential challenges for hydrothermal carbonization treatment, being affected by corrosion. It is further not clear the advantage of such a magnetic material.

Although the most attractive feature of hydrothermal carbonization is the energy savings (from bypassing the drying step compared to traditional pyrolysis), it has also been employed to treat dry feedstocks. Demir et al. produced N-doped porous carbon from organosolv lignin via a two-step hydrothermal carbonization and chemical activation scheme. 101 Hydrothermal carbonization was performed in water at 300 °C and 10 MPa, together with probe ultrasonication. Later, the

hydrochar was heated to 700-1000 °C for activation with the coaddition of KOH and adenine (N-doping source). With their approach, a high surface area was achieved (1788–2957 m² g⁻¹), together with up to 4.8 mmol g⁻¹ in CO₂ adsorption capacity (100 kPa, 25 °C). Heteroatom doping increased the CO2 capture performance slightly, compared to the pristine carbon. CO2 adsorption selectivity was dependent on the synergism of heteroatom doping and microporosity, rather than microporosity alone.

Atta-Obeng et al. used a two-step hydrothermal carbonization and activation with additional polyethyleneimine (PEI) functionalization. 102 Hydrothermal treatment was performed at 350 °C, activation at 4:1 KOH to char ratio 10 °C min⁻¹ to 800 °C for 1 hour. The activated samples were further functionalized with PEI at 5-25 wt%. Activation increased the surface area from 2.8 to 1341 m² g⁻¹, and PEI functionalization increased CO₂ adsorption capacity from 1.53 mmol g⁻¹ to 2.0 mmol g⁻¹ at 30 °C. However, the authors also found that increasing PEI impregnation beyond 5% leads to severe pore blockage.

While typical two-step approaches separate carbonization and activation, alternative methods maintain simultaneous carbonization/activation treatments followed by a separate doping step. Dong et al. produced porous carbon from lignin using a KOH carbonization/activation step, followed by N-doping through hydrothermal treatment or pyrolysis. 103 Pyrolysis was performed at 5 °C min⁻¹ to 700 °C for 2 h at KOH: lignin mass ratios of 0.25, 0.5, 1, and 2. The resulting porous carbon was N-doped by mixing with 5 g of urea and subjected to pyrolysis under the same conditions used for carbonization/activation or by hydrothermal treatment in a reactor at 220 °C for 2 hours. The authors found that pore volumes in the range of 0.6-0.8 nm played a critical role in determining the adsorption capacity at 25 °C and 101 kPa, consistent with some density functional theory (DFT) simulation results that indicated an optimal pore size for CO2 adsorption of 0.62-0.72 nm, based on the optimal adsorption distance for CO₂ molecules (0.31-0.36 nm). However, at lower adsorption pressures or higher adsorption temperatures, pore size became less significant, and nitrogen-containing functional groups gained importance for enhancing chemical and physical interactions with CO2, thereby increasing adsorption capacity. The chemisorption of CO₂ by surface nitrogen groups at 50 °C was indicated by in situ diffuse reflectance infrared Fourier transform (DRIFT) spectra. Furthermore, the N-doped

samples exhibited higher selectivity at lower adsorption pressures, suggesting that under those conditions, nitrogencontaining functional groups had a stronger affinity for CO₂ than narrow micropores. Overall, their approach achieved high surface areas of up to 2126 m² g⁻¹ and high adsorption capacities of up to 4.79 mmol g^{-1} at 25 $^{\circ}$ C and 101 kPa.

Bai et al. employed a two-step method involving simultaneous carbonization and N-doping treatment, followed by chemical activation. 105 Lignin was mixed with melamine at a 1:1 mass ratio and pyrolyzed at a rate of 10 °C min⁻¹ to 500 °C for 1 h under an Argon atmosphere. The resulting carbon powder was then mixed with CuCl2·2H2O at a weight ratio of 1:2 for activation and heated at 10 $^{\circ}$ C min⁻¹ to 800, 850, or 900 °C for 2 h. The authors found that melamine not only served its N-doping function but also introduced mesoporosity and enhanced micropore development during CuCl₂ activation. Notably, the selected lignin contained approximately 2% sulfur, which contributed oxidized sulfur functionalities to the carbon surface, increasing its basic character. The analysis revealed that narrow micropores (<1 nm) exhibited a strong linear correlation with adsorption capacity; meanwhile, nitrogen and sulfur content only showed a weak correlation. However, when comparing the IAST selectivity of the prepared adsorbents, it was found that heteroatom content (nitrogen and sulfur) was critical for enhancing selectivity towards CO2 over N2. Overall, the optimized method achieved high surface areas (1678 m² g⁻¹) and adsorption capacities (6.78 and 3.57 mmol g⁻¹ at 100 kPa and 0 °C and 30 °C, respectively), along with remarkable CO₂/N₂ IAST selectivity (132 and 81 at 100 kPa and 0 °C and 30 °C, respectively).

4.1.4 Other multi-step carbonization/activation routes. Multi-step schemes have been used to introduce heteroatoms or to tune the carbon porosity. In certain scenarios, additional thermal processing steps could be incorporated on top of the typical two-step carbonization and activation protocols, such as physical activation for increasing surface area or heteroatom doping to improve surface interactions with CO2. The primary objective of these extra steps is to improve the performance of the adsorbent, while, obviously, at the expense of increased operational complexity and costs. Liu et al. used a hydrothermal treatment with melamine, followed by mechanochemical processing of enzymatic hydrolysis lignin with KOH into pellets, and then proceeded with pyrolysis to produce an N-doped porous carbon material (Fig. 7). 106 In step 1, melamine was added to enzymatic hydrolysis lignin at a 1:1 mass ratio, and dispersed in water for thermal treatment (150 °C, 20 hours).

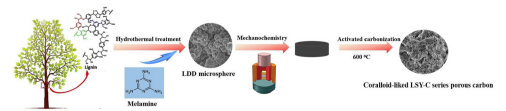


Fig. 7 Multistep process for mechanochemical processing of enzymatic hydrolysis lignin to nitrogen-doped activated carbon. (LDD: lignin-based hydrothermal carbon; LSY: lignin-based N-doped porous carbon). Reproduced with permission. 106 Copyright 2024, Elsevier.

In the second stage, the dried material was combined with KOH at a 4:1 ratio and compacted into 1.5 cm pellets at different pressures (10, 20, 30 MPa) and residence times (10, 20, 40 min). Finally, the resulting pellets (\sim 3 mm thick) were pyrolyzed in a tube furnace at a heating rate of 3 °C min⁻¹ to 500-800 °C and held for 1 hour. The advantage of introducing mechanochemical processing (step 2) is its operational simplicity and scale-up capability. Mechanical treatments also bypass more intensive chemical reaction schemes, such as high temperatures. After the multi-step treatment process, both the microporosity and pore volume of the lignin-based carbon material increased. In particular, CO2 uptake is measured to be improved with the introduction of narrow micropores (<1 nm) and N-containing groups. The resulting material could achieve large surface area (1233.2 m² g⁻¹), micropore volume $(V_{(d<1.0 \text{ nm})} 0.27 \text{ cm}^3 \text{ g}^{-1})$ and CO_2 adsorption capacity (5.00 mmol g⁻¹ at 0 °C).

Lignosulfonates represent a special case of lignin resources due to their sulfonated structure, higher average molecular weight distribution, and solubility in water, as previously introduced in Section 2.1. Yet, lignosulfonates are, to date, the most commercially relevant lignin resource. Gao et al. produced porous carbon from sodium lignosulfonate at various initial pH using a three-step approach, that sequentially involves hydrothermal carbonization, pyrolysis, and CO2 physical activation. 107 Sodium lignosulfonate (7.5 g) was first added to 100 mL aqueous solutions and titrated by sulfuric acid at a 1:7 ratio. Hydrothermal carbonization was then conducted at 260 °C for 24 hours. The resulting char was further carbonized at 10 °C min⁻¹ to 900 °C for 1.5 hours and, subsequently, physically activated using CO2 gas at 800-900 °C. The first step of acid treatment increased the degree of graphitization of the resulting carbon. Also, increasing acid dosage raised the carbon yield and decreased the oxygen content, reflected in a rise of carboxyl and a decrease of hydroxy and carbonyl groups.

The structural and chemical differentiations resulted in various CO2 adsorption capacities at different temperatures. The authors found that adsorption is primarily influenced by CO₂-substrate interactions at low temperatures (e.g., electrostatic and van der Waals forces), whereas the diffusion resistance of the pore channel becomes the dominating factor at higher temperatures, when CO₂ diffuses fast. With a vast pore size and functional group selection from the multi-step production of lignin-based carbon materials, Gao et al. investigated their corresponding interactions towards CO₂ molecules by quantum DFT (Fig. 8). It was shown that smaller pores and oxygen-containing functional groups increased the favorable adsorption energy between CO2 and the carbon surface, with carboxylic groups displaying the strongest attraction to CO₂. This is because the oxygen-containing groups act as a Lewis base to interact with the acidic CO₂ molecule. Ultimately, these lignin-derived carbons could achieve an adsorption capacity of up to 5.10 mmol g^{-1} at 0 °C and 100 kPa.

Park et al. synthesized an N-doped porous carbon from sodium lignosulfonate via the multi-step approach, which included (i) hydrothermal carbonization, (ii) KOH activation and (iii) post-doping with urea at high temperatures and under

Ar gas flow. 108 Hydrothermal carbonization was performed at 200 °C for 12 hours. The resulting char was activated with KOH at a 1:3 ratio at 5 °C min⁻¹ to 700 °C. The activated carbon was doped with urea at a 1:1-1:8 ratio, and subjected to a heating rate of 5 °C min⁻¹ to 800 °C. The KOH activation step significantly increased the surface area from 125 to 3170 $\text{m}^2\text{ g}^{-1}$. However, the doping step could introduce undesired blocking of micropores. The highest CO₂ adsorption capacity was characterized to be 13.6 mmol g⁻¹ at 1 MPa, achieved with the 4:1 mass ratio of urea to activated carbon, originating from the optimal balance of the high surface area and nitrogencontaining surface functional groups.

Following the concept of RCC, Zhao et al. produced a porous carbon CO2 reduction electrocatalyst from sodium lignosulfonate. ZnCl₂ activation was applied to pre-carbonized samples (pyrolysis at 500 °C for 1 hour), followed by the N-/P-doping with melamine or triphenylphosphorus. 109 The activation was operated with ZnCl₂ at a ratio of 2:1 at 800 °C for 2 hours. Then, the carbon materials were mixed with melamine (N) or triphenylphosphorus (P) at a 5:1 doping agent to lignin ratio, and pyrolyzed at 700–1000 °C for 2 hours under N₂. Both N and P doping are beneficial for CO₂ reduction and hydrogen evolution. However, P doping was more favorable towards hydrogen evolution than CO2 reduction, compared to the N-doped samples. The authors found that the enhanced CO2 reduction performance from their N-doped porous carbon was actually the result of a combination of multiple factors. For example, the textural properties, degree of graphitization, pore structure, and surface functionalities were highly dependent on the carbonization temperature, and the resulting electrocatalytic performance of the material was determined by the combined effect of these variables. Overall, their material was reported with surface areas of up to 1459 m² g⁻¹ and adjustable syngas H_2/CO ratios from 2.3 to 8.0.

4.2 Structured porosity via crosslinking

Amorphous hyper-crosslinked polymers are a type of porous materials with high surface area and pore volume, as well as adjustable surface chemistry. They can be synthesized with lignin via Friedel-Crafts reactions, which proceed from electrophilic aromatic substitution. Hyper-crosslinking is a promising approach for constructing highly rigid crosslinked bridges between the aromatic repeat units of lignin. These hypercrosslinked polymeric materials may not necessarily require carbonization steps, as they are designed to exhibit pore structures from crosslinking. These crosslinked samples are typically more mechanically and thermally stable than the raw material, giving them a higher tolerance to harsh environments. The major drawback of the crosslinking approach is the extra cost of chemical reagents and the complex chemistry of the reactions.

Meng and Weber crosslinked organosolv lignin with formaldehyde dimethyl acetal (FDA). The resulting materials were evaluated by their CO2 capture performance, either with or without further low-temperature pyrolysis (550 °C). In their scheme, organosolv lignin, anhydrous iron chloride (FeCl₃),

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(a) Micropore: Macropore: Nonporous: $\Delta E=-24.70 \text{ kJ/mol}$ 46 87 k.I/mol -30.57 kJ/mol **(b)** Pure carbon sheet: C=O: ΔE =-212.56 kJ/mol ΔE=-234.12 kJ/mol ΔE =-224.45kJ/mol $\Delta E = -225.32 \text{ kJ/mo}$

Fig. 8 Carbon sheets with varying pore sizes represented by ball-and-stick model and electron cloud density with bond lengths between the oxygen atoms of CO_2 , as well as calculated adsorption energy specified. Here, (a) depicts the comparison between interactions with different pore structures (non-porous, microporous, macroporous) and (b) shows the comparison of various functional groups on the carbon sheets (pure carbon, carbonyl, hydroxy, and carboxylic acid group). Reproduced with permission. 107 Copyright 2024, Elsevier.

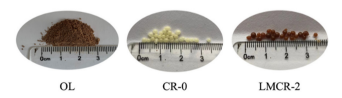


Fig. 9 Lignin-modified hyper-crosslinked porous resins using from poplar sawdust organosolv lignin. (OL: organosolv lignin; CR-0: resin without lignin; LMCR-2: resin with lignin). Reproduced with permission. (112 Copyright 2024, Elsevier.

and FDA were mixed in 1,2-dichloroethane (DCE) and added to a round bottom flask equipped with a condenser under an Ar atmosphere. The mixture was stirred for 1 hour at room temperature, followed by heating at 45 $^{\circ}$ C for 5 hours to form a crosslinked network. Subsequently heated at 80 $^{\circ}$ C for 19 hours to complete the reaction. Both materials were found to be primarily ultra-microporous. Notably, the crosslinked material was capable of maintaining its shape during carbonization, in addition to having increased CO_2 uptake, despite a reduced selectivity. 110

Shao *et al.* used six model lignin monomer units to prepare a hyper-crosslinked polymer in a one-pot Friedel–Crafts reaction, also utilizing the FDA as the crosslinker. ¹¹¹ These six phenols were first dissolved in DCE at 35 $^{\circ}$ C. Catalysts (FeCl₃, or AlCl₃) were added to the mixture and then heated to 80 $^{\circ}$ C under reflux for 24 hours. The crosslinking reaction took place in the porous materials, while some of the samples could preserve

micro-scale spherical morphologies. However, the materials exhibited relatively low specific surface areas for all six monomer types (7.8–246.9 $\text{m}^2\text{ g}^{-1}$). Of the monomers studied, 4-ethyl phenol produced the highest surface area by far (246.9 m² g⁻¹). The authors hypothesized that this higher surface area is due to more highly reactive sites leading to increased crosslinking and more abundant micro/mesopores. They proposed that for the other phenol monomers, the Friedel-Crafts reaction suffer from steric hindrance. Further, they speculated that methoxy groups have an adverse impact on crosslink formation via Friedel-Crafts reaction due to catalyst deactivation by complexation with FeCl3 and the oxygen atoms. Their hypercrosslinked polymers could achieve up to ~ 1.5 mmol g⁻¹ at 0 °C.

Liu et al. produced lignin-modified hyper-crosslinked porous resins, using extracted ethanol organosolv lignin from poplar sawdust (Fig. 9). 112 A free-radical copolymerization step was first conducted within the organic phase of lignin, VBC (monomers), DVB (crosslinker), ethyl acetate, and 2,2-azobisisobutyronitrile (AIBN) (initiator) in a 1 wt% aqueous poly(vinyl alcohol) (PVA) solution at 45 °C. Continuous stirring was applied while the mixture was gradually heated to 75 °C for 3 hours, 85 °C for 3 hours, and 95 °C for 2 hours, respectively. After purifying and drying the lignin-modified polymer spheres, Friedel-Crafts reactions were performed to crosslink the resins. 113 While their results showed that lignin could be incorporated in the hyper-crosslinked composites, most of the lignin-modified resins became broken particles after Friedel-Crafts reactions. It is

most likely that lignin destabilizes and disrupts the network of copolymers during suspension polymerization. They also found that the control resin (without the addition of lignin) had superior CO₂ adsorption capacity. Therefore, the lignin addition was predominantly not beneficial for the formation of micropores, potentially because of reduced crosslinking in the presence of lignin. The authors found that CO2 adsorption was highly dependent on both specific surface area and micropore volume. Notably, at reduced pressure (25 °C and 15 kPa), one of the ligninmodified resins could achieve better performance than the control sample, potentially the results of weaker surface chemistry without the addition of lignin. Their lignin-modified hypercrosslinked porous resins could achieve up to ~ 2 mmol g⁻¹ at 0 °C and 100 kPa.

Karaaslan et al. used epoxide chemistry to react kraft lignin in alkaline solutions with epichlorohydrin and then crosslinked into hydrogels using diamine hardeners (Fig. 10).114 The hydrogels were subsequently solvent exchanged into organogels and then supercritically dried with CO2 to keep the internal pore structure. Carbonization produced a high surface area structured carbon aerogels that contained both mesopores and micropores. Activation resulted in an increase in surface area (up to 1609 m² g⁻¹) and micropore volume and 5.4 mmol g⁻¹ of CO₂ adsorption capacity at 0 °C was achieved at 100 kPa. 115

4.3 Hierarchical-structured adsorbents

The approaches discussed so far produce materials with multiscale porosity, but most of them do not attempt to control the

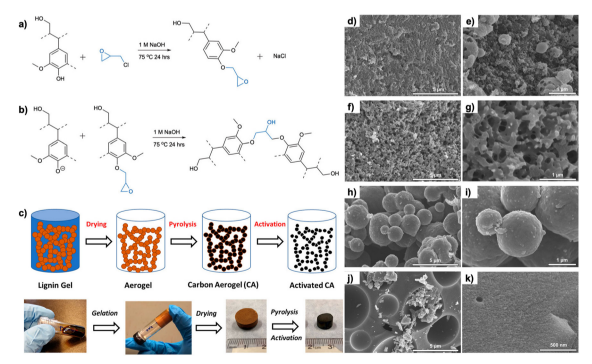


Fig. 10 Epoxy chemistry for lignin treatment and carbonization. Reaction mechanism between lignin and epichlorohydrin: (a) Epoxidation of hydroxyl groups of lignin, followed by (b) formation of crosslinks. (c) Schematic and photography of the processing steps of lignin carbon aerogels. (d)-(k) SEM images of the lignin carbon aerogels, showing how surface morphologies were affected by the lignin-to-crosslinker ratio: (d), (e) 55% lignin, (f), (g) 68% lignin, (h), (i) 81% lignin, (j), (k) 87% lignin. Reproduced under terms of the CC-BY 4.0 license. 114 Copyright 2022, Frontiers Media S.A.

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hierarchical structure at multiple length scales. More recently, controlled synthesis of multiscale architectures has become increasingly desirable because it directly impacts CO_2 capture applications. These approaches offer improved integration in scalable processes, improved gas phase transportation efficiency, and facilitate the accessibility of CO_2 gas molecules to surface adsorption sites. In this section, we cover the approaches that attempt to control the hierarchical architecture of the material.

4.3.1 Direct formation of carbon structures. Graphitic carbon materials with a hollow core are known as carbon nanocages. Carbon nanocages have generated increased attention due to their unique geometric (3D nanostructure), high specific surface area, and tunable electronic structure. Qin *et al.* reported the direct synthesis of hollow graphitic carbon from lignin at low pyrolysis temperatures. The synthesis involved first reacting formalin/lignin/NaOH (3/7/0.5 w/w) at 70 °C for 1 hour, followed by fixing with ferric nitrate and polymerization at 10 °C for 12 hours. Finally, pyrolysis was performed at a slow heating rate (1 °C min⁻¹) to 600 °C for 3 hours under Ar. The process is both time- and energy-consuming, and, unfortunately, the resulting material had a relatively low specific surface area and CO₂ adsorption capacity of 192.8 m² g⁻¹ and \sim 0.65 mmol g⁻¹, respectively (25 °C and 101 kPa).

The conversion of lignin to carbon fiber materials has been discussed for at least 55 years, 118 since carbon fibers have excellent functional properties potentially useful for $\rm CO_2$ capture applications, such as high strength-to-weight ratio, stiffness, conductivity, and excellent thermal and chemical resilience. 119 Particularly, lignin is a promising feedstock for carbon fibers because of its low cost, high carbon content, and thermoplastic properties. Lignin-based carbon fibers are typically produced νia extrusion method, such as melt-spinning, wet-spinning, dryspinning, gel-spinning, and electrospinning.

Calvo-Muñoz et al. produced lignin-based carbon fibers from organosolv lignin with almost exclusively ultra-micropores (<0.7 nm) and reported promising breakthrough adsorption capacities of 1.3 mmol g^{-1} or 5.7 wt% (in a column system, 25 °C, 15 kPa). 120 The electrospun lignin/ethanol fibers were treated with an intensive oxidative thermostabilization under an air atmosphere with a ramp of 0.08 °C min⁻¹ to 200 °C, followed by a 48-hour hold. The materials were then carbonized at 900 °C. Carbon fibers from this protocol feature narrow micropores, which are likely to be responsible for the superior CO₂ uptake. A general and major challenge of this approach to utilize lignin-based carbon fibers is the requirement for a timeand energy-intensive thermostabilization step. Further, the density of the material, specifically of the carbon fibers, is much less than that of regular granulated lignin-based activated carbon, which has implications for their translation to scalable processes.

4.3.2 Templating methods. In the field of pore structure engineering, ordered porous materials are a class of materials that exhibit long-range order and size homogeneity in their pore structure.⁸⁴ Templating methods are the most common approaches for the production of ordered porosity. The shaping of the targeted material is guided by additional sacrificial

material of a controlled or defined morphology.¹²¹ These shaping methods can be classified into hard and soft templating. Soft templating exploits the assembled structures of soft matter, such as polymers, surfactants, foams, and emulsions, including the subsequent removal of the template. Similarly, hard templates employ solid templates to construct the desired pore structure.¹²²

4.3.2.1 Hard templating. Sani et al. produced ordered mesoporous amine-functionalized lignin-based carbons, using a siliceous mesostructured cellular foam as a hard template in a solvothermal process. In their approach, 2.4 g of lignin was mixed with 1.2 g of template in 15 mL tetrahydrofuran (THF). 123,124 After drying, the composite was carbonized (700-900 °C) for 1 hour, and the carbon product was subsequently stirred in 1 M NaOH for 24 hours to remove the hard silica template. The resulting carbons were demonstrated to be mainly mesoporous high BET surface areas (up to 960 m² g⁻¹) and mesopore volumes of 1.50 cm³ g⁻¹. Further, the carbon preserved a quasi-spherical porous morphology after carbonization, inherited from the mother template. While the pristine mesoporous carbon demonstrated low CO2 uptake, the mesoporous carbon could be used as a support for polyethylenimine (PEI) to enhance its CO2 adsorption properties. Given the mesoporous structure, the material could accommodate high PEI loadings of up to 60 wt%, at which point the CO₂ adsorption was highest at 2.95 mmol g⁻¹ at 75 °C and 15 kPa. Due to this combination, the PEI-modified carbons exhibited enhanced adsorption kinetics and good cycle stability.

Zhao et al. developed a templating method with basic magnesium carbonate (BMC) or magnesium oxide (MgO) for the production of activated carbon from acid-precipitated alkali lignin, and compared the obtained results with those of samples chemically activated with ZnCl₂ or KOH. 125 In the templating approach, lignin was dispersed in water with BMC or MgO at a 1:3 or 2:1 ratio, respectively. The dispersion was later heated to 80 °C, followed by drying and calcination at 5 °C min⁻¹ to 600 °C for 2 hours. The templated carbons appeared rough with slit-like pores, whereas relatively flat with etched pores were found with the chemically activated samples (Fig. 11). However, the templated samples exhibited weaker performance than the chemical activation counterparts, in terms of specific surface area, total pore volume, and CO2 adsorption capacity. The latter reached specific surface areas of 1336.5 $\mathrm{m^2\,g^{-1}}$ and adsorption capacity of 3.60 mmol $\mathrm{g^{-1}\,CO_2}$ at 25 °C. Apparently, chemically activated samples had a more pronounced microporosity (89.1-93.2%) compared to the templated samples (58.7-44.6%). This is probably because hard templating methods mainly influence macroscale morphologies. Besides, CO₂ interaction relies more on microporosity, whereas macroporosity is more critical for transport phenomenon.

4.3.3.2 Soft templating. The amorphous 3D nature of the lignin macromolecular structure limits its potential for producing well-defined structures such as fibers. In efforts to fully unleash the potential of multiscale design, new lignin morphologies have

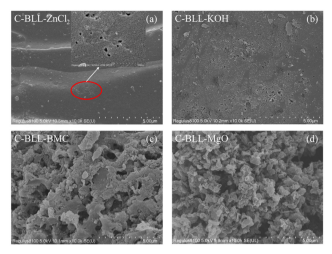


Fig. 11 Porous carbon materials prepared from lignin by using four different activation strategies: (a) chemical activation with ZnCl₂ and (b) KOH. Templating method with (c) BMC and (d) MgO. (C-BLL: carbonized black liquor lignin; BMC: basic magnesium carbonate) Reproduced with permission. ¹²⁵ Copyright 2023. Elsevier.

garnered interest in combination with available bio-based additives. Such lignified hierarchical assemblies of plants found in nature can inspire the construction of new hierarchically structured materials with engineering porosity.

Geng *et al.* used an ice-templating approach to produce carbon aerogels from kraft lignin. 126 Lignin was pre-mixed with TEMPO-oxidized cellulose nanofibers (TOCNF) of \sim 8–12 wt%. The high aspect ratio of TOCNF can establish a robust fibrous network that forms the aerogels during ice-templating and freezedrying. The as-produced aerogel was carbonized with a 2-hour hold at 500 °C to stabilize the material, followed by heating to 1000 °C. This ice-templating approach could produce an anisotropic macroporous structure, while the carbonization process could introduce *meso*- and micropores, demonstrating hierarchical porosity (Fig. 12). The carbon aerogels at 12 wt% TOCNF exhibited a high surface area (1101 m² g $^{-1}$) and adsorption capacity of 5.23 mmol g $^{-1}$ at 0 °C and 100 kPa.

In most literature, a structural polymer would be required to interconnect spherical lignin particles during soft templating since lignin itself is difficult to form a mechanical-robust interconnected network. Under such scenarios, the most common

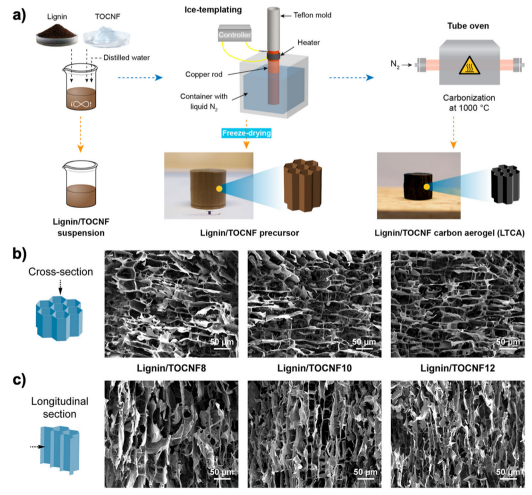


Fig. 12 (a) Synthesis of lignin/TOCNF carbon aerogels, and SEM images of (b) cross section and (c) longitudinal section of the lignin/TOCNF precursors with different TOCNF contents. Reproduced under terms of the CC-BY 4.0 license. (Copyright 2022, American Chemical Society.)

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choice is cellulose. Cellulose naturally assembles in the form of fibrils and can be liberated from the cell wall matrix, in the form of cellulose nanofibrils (CNF) or nanocrystals (CNC). Both CNFs and CNCs have garnered significant attention as functional agents in nanocomposite materials due to their attractive physical and chemical properties, not least their universal availability. For example, it has been demonstrated that the topology of nanonetworks formed from CNFs induces interparticle cohesion and enables universal and robust superstructure of nanoscale particles for the lay-in of spherical lignin particles. Hurther, the cellulosic network acts as a sacrificial template during carbonization due to its lesser relative thermal stability and carbon content. These features are particularly attractive given the increasing momentum for spherical lignin particles as promising and versatile assemblies.

The rate of CO₂ adsorption/desorption in lignin-based carbon materials is often limited by the diffusion rate of CO₂ gases in amorphous carbon networks. To address such bottlenecks, the superstructuring of spherical lignin particles can be employed to tailor the molecular transport. In this context, Zhao et al. produced multiscale carbon supraparticles, utilizing the drying cohesion of cellulose nanofibrils to bind spherical lignin particles (Fig. 13).54 Softwood kraft lignin was applied to produce spherical particles at 200 and 420 nm via solvent exchange. Supraparticles were formed via evaporation-induced self-assembly of aqueous lignin particles and CNF over various ratios (CNF solid fraction in the dried supraparticles ranging from 0.5-15 wt%) by casting on a superhydrophobic Tefloncoated surface at 60 °C. The dry supraparticles required oxidative thermostabilization (0.01-5 °C min⁻¹) at 250 °C for 2 hours under an air atmosphere. After thermo-stabilization, the supraparticles could be carbonized at 10 °C min⁻¹ to 600-900 °C for 1 hour. Physical activation was also performed with N2 gas

saturated with water (steam activation) or N_2 gas saturated with aqueous ammonia vapor (ammonia-steam activation).

The properties of lignin supraparticles are largely dependent on their specific preparation protocol. The thermal stability of lignin supraparticles could be improved by a slow oxidative thermo-stabilization, transforming lignin from a fusible thermoplastic to a thermosetting material. The rate of pre-oxidation is also critical to preserve the particles' hierarchical properties. In general, smaller particles require slower heating rates, which requires more time and energy investment. Since the preparation of supraparticles relies on drop casting, the droplet size is critical to ensure the roundness of the as-produced supraparticles. Too large droplets would be deformed by gravity into more elliptic shapes. CNF is also critical for providing mechanical robustness and avoiding structural failure; without CNF, a ring-shaped superstructure was obtained. CNF loadings > 2.5 wt%, and particle size in the range of 200-500 nm turned out to be the optimal processing conditions for the coassembly with the fibers. 128 The incorporation of CNF was also critical for preserving the suprastructure during thermal treatment. After oxidative thermo-stabilization and carbonization, the resulting carbon supraparticles with CNF were preserved, while the CNFfree particles were reduced to a powder.

The excellent mechanical strength and controlled macrostructure of the spherical supraparticles are well-suited for packed bed adsorption systems, since they are able to withstand the load of an entire packed column and reduce pressure drop (*i.e.*, enabling sufficient gas transport during operation), especially compared to smaller materials. The specific surface area and CO_2 uptake of the carbon supraparticles ranged from 348 to 405 m² g⁻¹ and ~1.6 to 1.3 mmol g⁻¹ (40 °C and 101 kPa), respectively, depending on the final carbonization temperature. In an attempt to further improve the CO_2 capacity

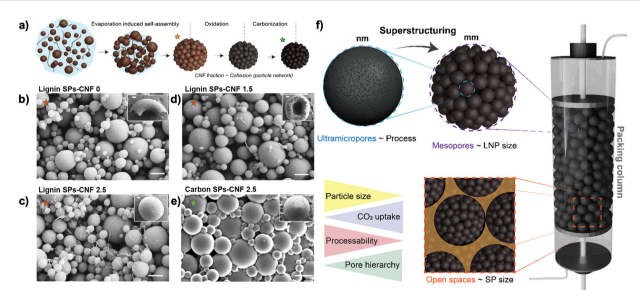


Fig. 13 (a) Process of carbon supraparticle formation *via* evaporation-induced self-assembly followed by oxidation and carbonization. (b)–(e) Electron microscope images of the lignin-based supraparticles with different contents of CNF (insets are at lower magnification). (f) Super structuring of lignin particles (nanometer-scale) into millimeter-scaled carbon supraparticles (SPs) for use in CO₂ capture in a packed column. Reproduced under terms of the CC-BY 4.0 license.⁵⁴ Copyright 2021, American Chemical Society.

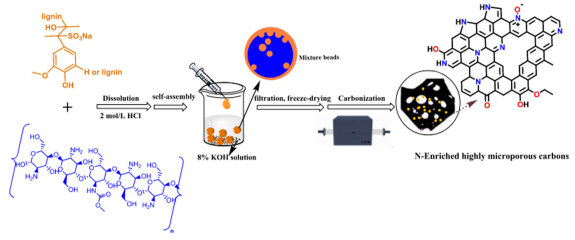


Fig. 14 Preparation process of N-doped highly microporous carbon derived from lignin/chitosan composites beads. Reproduced with permission. 130 Copyright 2023, Elsevier.

of the supraparticles and also to investigate the potential synergism of ultramicropores and N-doping on CO₂ uptake, the effect of N-doping on CO2 content was analyzed with the same structure but different nitrogen content with steam and ammonia-steam activation. It was found that steam could enhance the surface area to up to 967 m² g⁻¹ and ammonia 1152 m² g⁻¹ with N-content of 4.78 wt%. Further, the maximum CO2 uptake for steam activation and ammonia-steam activation was 1.8 and 1.7 mmol g^{-1} , respectively. Comparing the samples with steam vs ammonia-steam physical activation, showed similar porosity, surface area, graphitic structure, and CO2 uptake, suggesting incorporation of nitrogen heteroatoms may not have a significant influence on CO2 uptake. In summary, it was found that the carbon supraparticles with larger surface area or doped with nitrogen actually did not show higher CO₂ uptake, highlighting the complexity and multivariate nature of CO2 uptake of porous carbon materials.

Shao et al. produced a naturally N-doped porous carbon adsorbent through the self-assembly of sodium lignosulfonate and chitosan to gel microbeads in the presence of KOH for chemical activation (Fig. 14). 130 In their protocol, sodium lignosulfonate and chitosan were mixed at different mass ratios (3:1, 2:2, 1:3) in water followed by the addition of 2 M hydrochloric acid (HCl) to dissolve the chitosan with dispersed lignosulfonate. The dispersion was subsequently dropped into an 8 wt% KOH solution to form the gel beads. After drying, the beads were carbonized at 5 °C min⁻¹ to 600 °C for 2 hours. If the beads were washed to remove KOH prior to carbonization, the beads could maintain their structure to produce an aerogel, compared to those reduced into powder forms if no washing. Nonetheless, the powdered material (i.e., no washing treatment) exhibited a high surface ($\sim 892-1308 \text{ m}^2 \text{ g}^{-1}$) area, whereas the washed sample had a very low surface area (13.4 m² g⁻¹). XPS measurements showed that the N-content of the carbon materials ranged from $\sim 3-5\%$ according to chitosan content. The porosity and N content of N-doped microporous carbon could be effectively regulated by adjusting the ratio of lignin to chitosan in the precursor gel microbeads. Surface area, total pore volume, micropore volume, and N-content increased as the lignin:chitosan ratio decreased.

5. Perspectives on the rational design of lignin-based adsorbents

In the previous section, we reviewed the state-of-the-art of lignin-to-carbon processing and its implications in CO2 capture. Table 1 provides a summary of the lignin-based porous adsorbents covered in this review, from traditional approaches to structured porosity via crosslinking and hierarchically structured materials. However, one has to admit that the development of lignin-based materials is mostly in the early stages, and rational design will become essential to improve CO2 capture performance. In this section, we provide several perspectives on the future development of lignin-based carbon materials, that are design-driven following (i) a fundamental understanding, and (ii) computational simulations. Lastly, we introduce recent developments that use artificial intelligence (AI).

Fundamental design-driven carbon capture with lignin 5.1

5.1.1 Physisorption. The most critical factor affecting the CO₂ uptake potential of lignin-based adsorbents is the porous structure, which is also the most prioritized feature in material development. 133,134 CO2 capture in a porous medium is primarily controlled by physisorption, which originates from van der Waals interactions. Due to the nature of non-covalent bonding, the physisorption process of CO2 adsorption occurs spontaneously without the requirement of activation energies, but is also reversible at the same time. 135 Therefore, an optimized porous structure could offer more interaction points for effective CO₂ capture via physisorption, of which the surface area and pore volumes should characterize the properties. It is worth mentioning that the presence of functional groups (e.g., nitrogenous groups) could significantly enhance the

 Table 1
 Summary of lignin-based porous adsorbents^a (processing methods, surface area and adsorption performance)

Feedstock	Other chemicals	Processing technique	$\frac{\mathrm{BET}}{\mathrm{SSA}^b}$	$\begin{array}{c} {\sf Adsorption} \\ {\sf capacity}^c \end{array}$	Selectivity ^d	Ref.
Traditional approaches High-ash content alkali lignin	ı	Pyrolysis, negative/atm./elevated	1577	3.6 (0 °C)	I	98
Corn straw lignin Kraft lignin	Chemical activation (H_3PO_4) Chemical activation (KOH)	pressure Pyrolysis Pyrolysis	820 2121	$0.4~(0~^{\circ}\mathrm{C}) \ 3.3,~2.0~(0, 25~^{\circ}\mathrm{C})$	IS: \sim 15 (20 °C) IAST: 38 (25 °C & 15 PPa)	88
Dealkaline lignin (TCI America) Softwood kraft lignin (UPM	Chemical activation (KOH), S-doping $(Na_2S_2O_3)$ Chemical activation (HNO_3) , N-doping $(Urea)$	Pyrolysis Pyrolysis	3626 1000	11 (25 °C) 11.4 (20 °C &	IAST: $\sim 60~(25~^{\circ}\text{C})$ IAST: $\sim 18~(25~^{\circ}\text{C})$	90 91
BioPiva) Enzymatic hydrolysis lignin	Chemical activation (KOH), physical activation (humidified N.)	Microwave	2870	15 KFa) 1.3 (30 °C)	I	92
Dealkaline lignin (TCI America)	(indimental 182) Chemical activation (KOH), physical activation/ N-doning (NH3)	Pyrolysis	2922	$8.6,5.5(0,2.5^{\circ}\mathrm{C})$	IAST: $\sim 100~(25~^{\circ}\mathrm{C})$	93
Dealkalized lignin	Chemical activation (KOH)	Pyrolysis	2604	$\begin{array}{cccccccccccccccccccccccccccccccccccc$	IAST: $\sim 16~(25~^{\circ}\text{C}, \text{CO}_2/\text{N}, 10:90)$	94
High ash content lignin (Borre-	Chemical activation (KOH)	Hydrothermal carbonization,	3235	$7.4, 4.6 (0, 25 {}^{\circ}C)$		86
gaard right of the Eucalyptus enzymatic hydrolysis lignin Norway spense braff lignin	Bio-oil production (formic acid, catalyst), chemical	Eyrotysis Hydrothermal carbonization,	2875	6.0 (0°C)	IAST: $\sim 4~(0~^{\circ}C)$	46
ngun, youway spiroc man ngun Organosolv lignin (lignol innova- tion company)	Chemical activation (KOH), N-doping (adenine)	Pyroysis Hydrothermal carbonization, pyrolysis	2957	4.8 (25 °C)	IS: 18 (0 °C, CO ₂ /N ₂ 10: 90) IAST: 22 (0 °C, CO ₂ /N ₂ 10: 90)	101
Not specified	Chemical activation (KOH), functionalization (PEI)	Hydrothermal carbonization,	1341	2.0 (30 °C)	(CC - CT	102
Dealkalized lignin (TCI Shanghai)	Chemical activation (KOH), N-doping (Urea)	Pyrozyza Pyrolysis, hydrothermal carbonization	2126	4.79, 1.44 (25 °C at 101, 15 kPa) 2.97, 0.63 (50 °C at 101, 15 kPa)	IAST: 36 (25 °C)	103
Lignin (Rizhao Huatai Paper Industry)	N-Doping (melamine), chemical activation (CuCl $_2$ · 2H $_2$ O)	Pyrolysis	2703	6.78, 3.57 (0, 30 °C at 100 kPa)	IAST: 132, 81 (0, 30 °C)	105
Model lignin (methylene bridged mesitylene)	Polymer synthesis (mesityylene, HCl, HCHO, NaCl, ZnCl ₂ , CH ₂ Cl ₂ , FeCl ₃ , CH ₃ OH), chemical activation (KOH)	Oxidative thermostabilization, pyrolysis	1899	6.16 (0 °C)	IAST: 24, 28 (0, 25 °C)	131
Enzymatic hydrolysis lignin	N-Doping (melamine), chemical activation (KOH)	Hydrothermal carbonization, mechanochemical processing,	2030	5.0 (0 °C)	IAST: 525 (0 $^{\circ}$ C) Henry's Law: 29 (0 $^{\circ}$ C)	106
Sodium lignosulfonate	pH adjusting (H_2SO_4), physical activation (CO_2)	Hydrothermal carbonization,	1336	5.1 (0 °C)	IAST: 27 (25 °C, CO ₂ /N ₂	107
Sodium lignosulfonate	Chemical activation (KOH), N-doping (Urea)	Pytotysis Hydrothermal carbonization,	3064	2.7, 13.0 (25 °C	10:30) -	108
Sodium lignosulfonate	Chemical activation (ZnCl_2), N-/P-doping (melamine/triphenylphosphorus)	pyrotysis Pyrotysis	1459	& 101, 1013 M a)	${\rm CO_2RR}$ performance: ${\rm H_2/CO}$ ratios 2.3–8.0	109
Poplar organosoly lignin Model lignin monomers	E Crosslinking (FDA, FeCl ₃ , DCE) Crosslinking (FDA, FeCl ₃ , AlCl ₃ , DCE)	Pyrolysis —	140 247	$\begin{array}{c} 2.2 - 2.5 \; (25 \; ^{\circ}\text{C}) \\ 1.5 \; (0 \; ^{\circ}\text{C}) \end{array}$	60 (30 °C) IAST: 257 (0 °C) Henry's I our. 25 (0 °C)	110
Poplar sawdust ethanol organosolv lignin	Crosslinking (VBC, DVB, ethyl acetate, AIBN, PVA)	I	696	2.0 (0 °C)	——————————————————————————————————————	112
Southern pine softwood kraft lignin (BioChoice TM)	Crosslinking (NaOH, epichlorohydrin)	Supercritical-drying, pyrolysis	1609	5.4 (0 °C)	I	114

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Table 1 (continued)						
Feedstock	Other chemicals	Processing technique	$\overset{\text{BET}}{\operatorname{SSA}^b}$	$\begin{array}{ll} \mathrm{BET} & \mathrm{Adsorption} \\ \mathrm{SSA}^b & \mathrm{capacity}^c \end{array}$	Selectivity ^d	Ref.
Hierarchically structured-adsorbents	ts					
Not specified	Formalin, NaOH, ferric nitrate	Pyrolysis	193	$0.7~(25~^{\circ}\mathrm{C})$		117
Alcell $^{\circledR}$ lignin	Electrospinning (ethanol)	Oxidative thermostabilization, pyrolysis	850	$1.3~(25~^{\circ}{ m C}$ & $15~{ m kPa})$	I	120
Lignin (Sigma-Aldrich)	Hard template (siliceous mesostructured foam, THF), functionalization (polyethyleneimine)	Pyrolysis	096	$3.0~(75^{^\circ}\mathrm{C}~\&$ $15~\mathrm{kPa})$	1	123,124
Acid precipitated lignin (alkali pulping company, Hunan, China)	Templating (BMC, MgO), chemical activation (ZnCl ₂ , KOH)	Pyrolysis/calcination	1337	$5.2,3.6,2.2$ $(0,25,50^{\circ}\mathrm{C})$	IAST: $14.2, 12.7 (25 ^{\circ}\text{C}, \text{CO}_2/\text{N}_2, 10: 90, 15: 85)$	125
Kraft lignin (Sigma-Aldridge)	TEMPO-oxidized cellulose nanofibers	Ice-templating, thermal stabilization, pyrolysis	1101	5.2 (0 °C)		126
Softwood kraft (BioPiva 100), pine kraft lignin (indulin AT)	Particle formation (THF/DMF), superstructuring (CNF), physical activation (ammonia-steam)		1152	1.75 (40 °C)	1	54
Sodium lignosulfonate	Bead formation & N-doping (chitosan), chemical activation (KOH)	Pyrolysis	1308	$4.9,3.2\ (0,25^{\circ}\mathrm{C})$	IAST: 146 $\left(0~^{\circ}\mathrm{C}\right)$ Henry's Law: 24 $\left(0~^{\circ}\mathrm{C}\right)$	132

^a Performance metrics reported are maximums achieved and do not necessarily correspond to the same sample. ^b SSA reported in units of m² g⁻¹, ^c CO₂ adsorption capacity in mmol g⁻¹, assumed to be measured at 101 kPa and 15% CO_2/N_2 unless otherwise absorption capacity can be assumed to be measured at 101 kPa and 15% CO_2/N_2 unless otherwise specified

material affinity towards CO2 molecules, attributed to much stronger chemisorption mechanisms, which will be discussed later in Section 5.1.2.

The pore size distribution requires specific attention, when aiming at improving the carbon capture efficiency of biomassderived porous materials. The significance of pore size and size distributions lies in their impact on the accessibility of reaction sites and the accommodation of CO2 molecules within the material matrix. For well-defined synthetic nanomaterials (e.g., MOFs, zeolites, porous organic polymers), their CO₂ capture capability can be precisely controlled by tuning the microporosity and mesoporosity during material synthesis. 136,137 For heterogeneous bio-based materials, pore sizes could exhibit a broad distribution, ranging from a few nanometers to micrometers, depending on the pretreatment and activations. Such a hierarchical porosity is generally considered to be beneficial for CO2 capture efficiency with a sacrifice in selectivity. 138 Macroporosity facilitates the gas flow and transportation within a torturous porous medium. Mesopores could promote local mass transports and act as local constraints to enhance gas-solid interactions, being suitable for highpressure CO2 adsorption, while improving microporosity is the key to generating active adsorption sites in both low- and high-pressure conditions.

Accordingly, tailoring pore sizes within lignin-based materials affects the kinetics of CO₂ molecule adsorption. In general, faster CO2 uptake kinetics is favorable. The early-stage diffusion of CO₂ gas into microporous regions is the rate-limiting step for adsorption kinetics. Increasing the pressure could facilitate the accessibility of CO₂ molecules to micropores that are smaller than 0.33 nm to allow strict physical adsorption. 139,140 In real practice, functional groups are introduced to improve the CO2 affinity with adsorbent substrates, accelerate their early-stage diffusion, and enhance low-pressure adsorption performance. Templating and controlled activation processes are the most common methodologies for producing consistent and well-dispersed pore structures. 17,54,126,141 These endeavors seek to strike a balance between pore size and distribution, aiming for materials that offer enhanced adsorption kinetics and selectivity in CO2 capture applications. Recently, Keffer et al. 142,143 utilized computational approaches to investigate the effect of N-doping in lignin-derived carbon quantum dots (CQDs) for carbon capture. They found that both N-doping within the aromatic structure of the CQDs and the introduction of amine groups positively influenced CO2 capture efficiency and selectivity over N2, O2, and H2O.

5.1.2 Chemisorption. Apart from the physisorption, CO₂ uptake can be largely facilitated through the introduction of chemical binding sites, i.e., chemisorption. Common functional groups for chemisorption include various N-containing molecules, alkoxides, and carboxyl groups, as successfully demonstrated on a variety of MOFs 102,144,145 and biomassbased porous carbon materials. 146 In terms of amino groups, the DFT results indicate that the neutral amino group can only weakly adsorb a CO2 molecule through non-covalent interactions, as evidenced by the N-C distance of 2.82 Å. 147 Nitrogen atoms in the bulk materials exhibit a Lewis basic characteristic,

indicating an affinity towards CO_2 and the potential to serve as CO_2 adsorption sites.¹⁴⁸ This is because the deprotonated amine group $R_1R_2N^-$ easily catches a dissolved CO_2 molecule to produce carbamate, through the following reactions:¹⁴⁷

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 $2R_1R_2NH \rightleftharpoons R_1R_2NH_2^+ + R_1R_2N^-$ (1)

$$R_1R_2N^- + CO_2 \rightarrow R_1R_2NCOO^-$$
 (2)

Additionally, the nitrogen atoms from tertiary amines $(R_1R_2R_3N)$ tend to undergo protonation to generate hydroxide, which serves as the Lewis base to the capture the acidic CO_2 gas:^{149,150}

$$CO_2 + H_2O \rightarrow H_2CO_3 \rightarrow HCO_3^- + H^+$$
 (3)

$$HCO_3^- \to CO_3^{2-} + H^+$$
 (4)

$$R_1R_2R_3N + H^+ \rightarrow R_1R_2R_3NH^+$$
 (5)

Alkoxides are another highly promising group of substances for adsorbing CO_2 . They are less sensitive to O_2 and have the ability to directly interact with CO_2 to create a carbonate compound.

$$RO^- + CO_2 \rightarrow ROCO_2^-$$
 (6)

Even though the above-mentioned functional groups are likely to provide chemical reaction sites for ${\rm CO_2}$ capture, the steric effects of the applied macromolecules cannot be ignored. The exchange-repulsion from the non-polar groups could cancel the attractive effects between ${\rm CO_2}$ and reaction sites. ¹⁵¹ Due to the propensity of lignin to induce coil-globule transitions at high temperatures or in a cosolvent environment, ^{152–155} it is more probable that it would augment the exposed surface area, hence improving the accessibility of lignin for ${\rm CO_2}$ chemisorption.

Cui et al. grafted the active amine group on a lignin-derived compound vanillin and alkaline lignin by Mannich reaction. 156 The ratio of material $n({\rm vanillin}):n({\rm formaldehyde}):n({\rm amine reagent})$ is 1:1:1 in the protocol, all of which are 0.06 mol. The vanillin modified by acrylamide obtained a CO $_2$ uptake capacity of 0.114 g CO $_2$ per gram (2.6 mmol g $^{-1}$) of the material under 25 °C and 100 kPa. In-depth analysis identified the multisite synergistic adsorption process, in which CO $_2$ double interacts with the amide group and single interacts (hydroxy oxygen on the adsorbent's benzene ring) due to a large local difference in electronegativity and a small steric hindrance.

 ${\rm CO_2}$ adsorption mechanisms can involve a complex interplay of both physisorption and chemisorption. It is crucial to approach fundamental interpretations carefully, as an oversimplified acid-base interaction model may not adequately account for all interactions. Sun *et al.*¹⁵⁷ synthesized four conjugated azacyclo-copolymers that self-assembled into two-dimensional layered structures with well-defined and uniform amino and azacyclo groups. Experimental and computational results revealed that the negative electrostatic potential around the azacyclo-N-doping sites could induce strong van der Waals interactions, facilitating selective ${\rm CO_2}$ adsorption. Although the reported ${\rm CO_2}$ uptake efficiency was below the theoretical limit

(0.255 mmol g $^{-1}$ at 0 $^{\circ}$ C and 1 bar, compared to the 1 mmol g $^{-1}$ predicted for N-doping), this advancement represents a significant shift in the design of highly effective porous carbon adsorbents through chemical and surface modification. ^{158,159}

PEI impregnation is another common practice for lignin chemisorption requires amine modification/grafting, in which PEI is commonly applied with alkaline combined high-temperature treatment. An example has been given in previous sections, 124 where the derived mesoporous carbon material demonstrates a high $\rm CO_2$ adsorption capacity, above 2 mmol $\rm g^{-1}$. Some mechanistic analysis was also performed to identify the $\rm CO_2$ interactions with amide and hydroxy groups.

5.2 Computational design-driven carbon capture with lignin

Computational chemistry and molecular modeling approaches offer fundamental insights into complex physiochemical phenomena that would otherwise be inaccessible with experiments alone. These approaches have been employed in efforts to address a range of challenges in lignin chemistry, from its polymerization, linkages and supramolecular interactions in the plant cell wall, 160–162 to its functional properties such as solubility and glass transition temperature. 152,163,164

Computational methods, such as DFT and quantum mechanics/molecular mechanics (QM/MM), can be utilized to investigate molecular-level phenomena and support the development of modeling approaches at larger length scales, such as those conducted with molecular dynamics (MD) and Monte Carlo (MC) simulations. Of particular relevance for the production of carbon from lignin include reactive force field approaches that enable the modeling of thermochemical reactions (i.e., pyrolysis). 165 Yet, the variability and multiscale complexity of lignin offers a significant challenge for modeling and simulation approaches. 166 To address the challenge of lignin heterogeneity and variability in molecular simulations, tools have been developed for the generation of libraries of lignin polymer topologies suitable for classical and quantumlevel studies. 167,168 With the introduction of these tools, new levels of topological complexity may be studied, offering an improved representation of broader lignin populations.

Insights on the molecular level provide an in-depth understanding of the interaction of CO2 and porous carbon and guidelines on further materials design and optimization. Theoretical analysis has shown that there is physisorption between CO2 and cyclo[18]carbon, 169 as well as acyclic iminebased functional molecules and a range of functional molecules that contain multi-N-containing superbases and heteroaromatic ring complexes (Fig. 15).170,171 In order to gain a deeper comprehension and to distinguish the significant and insignificant factors of non-covalent interaction between CO2 and the lignin molecules, the utilization of energy decomposition analysis (EDA) is crucial. This method effectively decomposes the overall interaction energy into distinct physical components that are readily comprehensible. Briefly, EDA can be categorized into two main methods, one of which involves Kitaura–Morokuma, 172 LMOEDA, GKS-EDA, 173 NEDA, 174 and ALMO-EDA. 175 Another representative one is

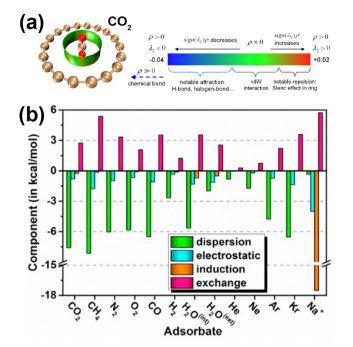


Fig. 15 Non-covalent interactions involved in CO_2 capture mechanisms. (a) Optimized structures and non-covalent interaction maps of CO_2 molecule containing cyclo[18]carbon. (b) Components of interaction energy between cyclo[18]carbon and various chemical species obtained from symmetry-adapted perturbation theory (SAPT) calculations. Reproduced with permission. 169 Copyright 2021, Elsevier.

the symmetry-adapted perturbation theory (SAPT) method. 176 Lu et al. employed SAPT theoretical calculations to forecast the physical adsorption of CO₂ on cyclo[18]carbon, incorporating a London Dispersion interaction energy of -7.6 kcal mol⁻¹, resulting in a total attraction ratio of 87.36%. 169 Neutral compounds, like aromatic systems, can exhibit a similarly high ratio of London Dispersion interactions. 171 On the other hand, this ratio will exhibit a bias towards electrostatic interaction (Ele) dominance, when the physical absorption occurs between CO₂ and ionic solvents.¹⁷⁷ Remarkably, the interaction strength between COO- and CO2 is nearly equal to that of pyrrole and indole, with a similar magnitude of higher than -10 kcal mol⁻¹. This result could provide a theoretical insight for surface modification on lignin molecules, such as aminefunctionalization and carboxylation, 178,179 to enhance noncovalent interaction with CO2.

5.3 Artificial intelligence (AI) design-driven carbon capture with lignin

In recent years, the development of AI has created a paradigm shift in material design strategies. Compared to the conventional bottom-up (*i.e.*, fundamental-driven design) or top-down (*i.e.*, performance screening) approaches, AI-driven design relies on the collection of large databases (*e.g.*, materials' structural, properties) through literature, experiments and/or simulations, and then using a nonlinear pathway within high-dimensional geometric-variable space¹⁸⁰ (*i.e.*, the machine learning algorithm) to optimize and predict performances.

The most critical criterion for successful machine learning research is the access to large amounts of reliable data. For example, machine-learning or deep-learning methods have been successful in determining the reaction pathways for catalyst designs, CO₂ adsorption, and particle analysis, among many others, ^{181–184} owing to the abundant data availability.

In the regime of lignin research, state-of-the-art research is mostly still in the stage of data exploration, collection, and validation. Establishing a more comprehensive lignin knowledge library would be beneficial to boost the AI-driven designs for lignin-based materials in general. In this regard, it is essential for researchers to provide a comprehensive description of the feedstock properties, including lignin supplier, biomass source, extraction process, chemical structure characteristics and purity indicators. Apart from the experimental results, simulations are less expensive for data collection without the need for materials gathering and consumption, despite a debatable extent of reliability. Nevertheless, current simulation means can be efficient in predicting intra-/intermolecular forces, colloidal interactions, particle assembly, *etc.*, 186–188 all of which can be the resources for machine learning research.

Conclusion and outlook

With growing concerns about climate change, developing efficient and sustainable CO_2 capture materials has become a matter of high relevance, including the exploration of related technologies for their upstream and downstream applications (e.g., CO_2 storage, utilization, and adsorbent regeneration). Despite its advancements, the aminated liquid-based carbon capture technologies face challenges due to the energy intensity required for solution regeneration, which limits the sustainability of its carbon-negative life cycle. In this context, carbon adsorbents are highly promising materials due to their costeffective synthesis, customizable properties, and potential to lower costs associated with adsorbent regeneration.

Bio-derived residues can be effectively converted into carbon materials with high surface area and multi-scale porosity. In particular, lignin stands out as a promising source due to its high carbon content, versatile functional structure, and low cost. This work reviewed recent developments in lignin-based carbon adsorbents for CO2 capture. Unlike petroleum-based chemicals, which are commercially upgraded into homogeneous compounds, the biomass processing industry, particularly in the context of lignin utilization, has not yet reached full maturity. We began this review by examining the origins of lignin feedstocks and their industrial varieties, thermal properties, processing strategies, and sustainability aspects. We follow with an overview of the key performance indicators for ligninderived carbon adsorbents, including their carbon properties and adsorption performance. We then covered state-of-the-art processing of technical lignins to produce carbon adsorbents and offered a critical discussion on this emerging interdisciplinary field, focusing on the potential of rational designs in future developments.

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The field of lignin-based materials for CO₂ capture is still in its early stages. Several challenges need to be addressed for future development. The primary challenge of lignin valorization lies in its natural variability, heterogeneity, and chemical complexity. The impact of lignin's origins on the final products is not yet fully understood, and there is a lack of consistency and broad applicability across various processing techniques. Further investigation into the structure–property relationships for carbon development is necessary. Despite these challenges, the field is advancing and aligning with current trends in the broader lignin community, such as fractionation, chemical modification, and the assembly into particles.

As discussed in this review, the diversity of lignin has significant implications for processing. This serves as both a caution and a reminder of the importance of selecting the appropriate lignin type. For instance, different grades of lignin can be utilized for specific purposes, such as using pulping salts as chemical activators. Additionally, the presence of heteroatoms in various technical lignins can enable novel processing strategies or provide options for additive-free doping, such as with water-soluble sulfur-containing lignosulfonates or nitrogen-containing enzymatic hydrolysis lignins. Thermal processing also varies with lignin selection due to differences in molecular weight and thermal stability. Furthermore, there are ongoing efforts to use advanced protocols, such as assembly and supraparticle formation, to address challenges related to disordered porosity and technological scalability, crucially, these approaches are highly dependent on lignin source and grade.

Due to the variety of new competing technologies, technoeconomic assessments should accompany lignin-based technologies to evaluate their feasibility at an early stage, identify key process variables, and establish benchmarks. New technologies require knowledge of various thermodynamic and kinetic parameters for design and optimization that go beyond the typical metrics such as adsorption capacity and selectivity. 189 Alternative technologies, such as MOFs and COFs, are also rapidly advancing, often exhibiting high adsorption capacities relative to carbon adsorbents. Membranes of various types of COFs have achieved up to 22.6 mmol g⁻¹ for TAPB-PDA COF aerogels. 190,191 While these materials have typically faced challenges with moisture, mechanical stability and scalable production, many of these challenges are also being addressed. 192 Furthermore, emerging functional materials such as photoresponsive adsorbents offer additional opportunties. 193,194 In this context, a systems approach to the rational design and comprehensive assessment of lignin-based adsorbents is essential to facilitate development in this active field.

Finally, current research on lignin carbonization is predominantly driven by top-down approaches, where experimental screening is used to identify optimal performance. Future developments could shift towards bottom-up designs, utilizing fundamental principles or AI-driven methods, which involve synthesizing designed structures followed by performance validation. This shift is due to the long-standing challenges in understanding the complex carbonization process. We anticipate

that in-depth mechanistic studies will enhance the rational design of lignin-based carbon materials, ^{195,196} and advance the broader field of carbon capture. Additionally, leveraging existing experimental and simulation data, we expect that combining machinelearning techniques with materials synthesis will enable data-driven optimization of the carbonization process.

Author contributions

D. B. R.: conceptualization, visualization, writing – original draft, writing – review & editing; J. C., Z. W.: writing – original draft; S. R.: writing – review & editing; I. B.: conceptualization, writing – review & editing; Y. D.: conceptualization, writing – original draft, writing – review & editing; Y. L.: conceptualization, supervision, writing – original draft, writing – review & editing; O. J. R.: conceptualization, supervision, funding acquisition, writing – review & editing.

Data availability

No primary research results, software or code have been included and no new data were generated or analyzed as part of this review.

Conflicts of interest

There are no conflicts to declare.

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References

- 1 W. H. Organization, Ecosystems and human well-being: health synthesis: a report of the Millennium Ecosystem Assessment, World Health Organization, 2005.
- 2 J. D. Shakun, P. U. Clark, F. He, S. A. Marcott, A. C. Mix, Z. Liu, B. Otto-Bliesner, A. Schmittner and E. Bard, *Nature*, 2012, **484**, 49–54.
- 3 P.-S. Wei, Y.-C. Hsieh, H.-H. Chiu, D.-L. Yen, C. Lee, Y.-C. Tsai and T.-C. Ting, *Heliyon*, 2018, **4**, e00785.

- 4 H. S. Kheshgi, S. J. Smith and J. A. Edmonds, *Mitigation and Adaptation Strategies for Global Change*, 2005, vol. 10, pp. 213–220.
- 5 J. W. Rae, Y. G. Zhang, X. Liu, G. L. Foster, H. M. Stoll and R. D. Whiteford, *Annu. Rev. Earth Planet. Sci.*, 2021, 49, 609–641.
- 6 T. M. Letcher, Future energy, Elsevier, 2020, pp. 3-17.
- 7 P. Friedlingstein, M. O'Sullivan, M. W. Jones, R. M. Andrew, D. C. Bakker, J. Hauck, P. Landschützer, C. Le Quéré, I. T. Luijkx, G. P. Peters and W. Peters, *Earth System Science Data*, 2023, 15, 5301–5369.
- 8 P. Fennell, J. Driver, C. Bataille and S. J. Davis, *Nature*, 2022, 603, 574–577.
- 9 R. E. Siegel, S. Pattanayak and L. A. Berben, ACS Catal., 2022, 13, 766–784.
- 10 T. C. Drage, C. E. Snape, L. A. Stevens, J. Wood, J. W. Wang, A. I. Cooper, R. Dawson, X. Guo, C. Satterley and R. Irons, J. Mater. Chem., 2012, 22, 2815–2823.
- 11 L. Joos, J. M. Huck, V. Van Speybroeck and B. Smit, *Faraday Discuss.*, 2016, **192**, 391–414.
- 12 L. Zheng, Oxy-fuel combustion for power generation and carbon dioxide (CO2) capture, Elsevier, 2011.
- 13 M. Pardakhti, T. Jafari, Z. Tobin, B. Dutta, E. Moharreri, N. S. Shemshaki, S. Suib and R. Srivastava, ACS Appl. Mater. Interfaces, 2019, 11, 34533–34559.
- 14 D. G. Boer, J. Langerak and P. P. Pescarmona, ACS Appl. Energy Mater., 2023, 6, 2634–2656.
- 15 M. Maluszynska-Hoffman, N. Repond, J. A. Wurzbacher and C. Gebald, WO 2017/009241 A1, 2017.
- 16 F. Baraka and J. Labidi, Sci. Total Environ., 2024, 912, 169093.
- 17 Y. Lu, M. Kamkar, S. Guo, X. Niu, Z. Wan, J. Xu, X. Su, Y. Fan, L. Bai and O. J. Rojas, *Small*, 2023, 19, 2300686.
- 18 T. Guo, Z. Wan, M. Panahi-Sarmad, G. Banvillet, Y. Lu, S. Zargar, J. Tian, F. Jiang, Y. Mao, Q. Tu and O. J. Rojas, *ACS Nano*, 2024, **18**, 14954–14967.
- E. Gomez-Delgado, G. V. Nunell, A. L. Cukierman and P. R. Bonelli, Carbon Lett., 2020, 30, 155–164.
- 20 J. Y. Lai, L. H. Ngu, S. S. Hashim, J. J. Chew and J. Sunarso, Carbon Lett., 2021, 31, 201–252.
- 21 A. Alabadi, S. Razzaque, Y. W. Yang, S. Chen and B. Tan, *Chem. Eng. J.*, 2015, **281**, 606–612.
- 22 A. E. Ogungbenro, D. V. Quang, K. Al-Ali and M. R. M. Abu-Zahra, Activated carbon from date seeds for CO₂ capture applications, 13th International Conference on Greenhouse Gas Control Technology (GHGT), Lausanne, Switzerland, 2016.
- 23 J. Serafin, M. Ouzzine, O. F. Cruz, J. Srenscek-Nazzal, I. C. Gómez, F. Z. Azar, C. A. R. Mafull, D. Hotza and C. R. Rambo, *Waste Manage.*, 2021, **136**, 273–282.
- 24 M. Wu, W. Chen, J. Hu, D. Tian, F. Shen, Y. Zeng, G. Yang, Y. Zhang and S. Deng, *Sci. Total Environ.*, 2019, **695**, 133898.
- Y. Lu, Y. Chun, X. Shi, D. Wang, F. Ahmadijokani and
 O. J. Rojas, *Adv. Mater.*, 2024, 36, 2400311.
- 26 C. Z. Li, X. C. Zhao, A. Q. Wang, G. W. Huber and T. Zhang, Chem. Rev., 2015, 115, 11559–11624.

- 27 D. S. Bajwa, G. Pourhashem, A. H. Ullah and S. G. Bajwa, Ind. Crops Prod., 2019, 139, 11.
- 28 O. Musl and A. Potthast, Lignin Chemistry, 2024, pp. 61-83.
- 29 J. K. Weng and C. Chapple, New Phytol., 2010, 187, 273–285.
- 30 W. Boerjan, J. Ralph and M. Baucher, Annu. Rev. Plant Biol., 2003, 54, 519–546.
- 31 Y. Mottiar, R. Vanholme, W. Boerjan, J. Ralph and S. D. Mansfield, *Curr. Opin. Biotechnol*, 2016, 37, 190–200.
- 32 M. Balakshin, E. A. Capanema, X. Zhu, I. Sulaeva, A. Potthast, T. Rosenau and O. J. Rojas, *Green Chem.*, 2020, 22, 3985–4001.
- 33 M. Y. Balakshin and E. A. Capanema, RSC Adv., 2015, 5, 87187–87199.
- 34 A. Berlin and M. Balakshin, in *Bioenergy Research: Advances and Applications*, ed. V. K. Gupta, M. G. Tuohy, C. P. Kubicek, J. Saddler and F. Xu, Elsevier, Amsterdam, 2014, pp. 315–336.
- 35 M. Y. Balakshin, E. A. Capanema, I. Sulaeva, P. Schlee, Z. E. Huang, M. Feng, M. Borghei, O. J. Rojas, A. Potthast and T. Rosenau, *ChemSusChem*, 2021, 14, 1016–1036.
- 36 P. Tomani, Cellul. Chem. Technol., 2010, 44, 53.
- 37 L. Kouisni, P. Holt-Hindle, K. Maki and M. Paleologou, J. Sci. Technol. For. Prod. Processes, 2012, 2, 6–10.
- 38 M. Hubbe, R. Alén, M. Paleologou, M. Kannangara and J. Kihlman, *BioResources*, 2019, 14, 2300–2351.
- 39 L. Dessbesell, M. Paleologou, M. Leitch, R. Pulkki and C. Xu, *Renewable Sustainable Energy Rev.*, 2020, **123**, 11.
- 40 N. Izaguirre, E. Robles, R. Llano-Ponte, J. Labidi and X. Erdocia, *Ind. Crops Prod.*, 2022, 175, 114251.
- 41 S. K. Singh, Bioresour. Technol. Rep., 2022, 17, 100958.
- 42 D. M. de Carvalho and J. L. Colodette, *BioResources*, 2017, **12**, 6907–6923.
- 43 Q. He, I. Ziegler-Devin, L. Chrusciel, S. N. Obame, L. Hong, X. Lu and N. Brosse, *ACS Sustainable Chem. Eng.*, 2020, **8**, 5380–5392.
- 44 E. Leng, Y. Guo, J. Chen, S. Liu, E. Jiaqiang and Y. Xue, *Fuel*, 2022, **309**, 122102.
- 45 L. A. Riddell, J. P. B. Lindner, P. de Peinder, F. Meirer and P. C. Bruijnincx, *ChemSusChem*, 2024, 17, e202301464.
- 46 A. Berlin and M. Balakshin, *Bioenergy Res.: Adv. Appl.*, 2014, 315–336.
- 47 M. A. Karaaslan, M. Cho, L.-Y. Liu, H. Wang and S. Renneckar, ACS Sustainable Chem. Eng., 2021, 9, 458–470.
- 48 M. Cho, M. Karaaslan, H. Wang and S. Renneckar, *J. Mater. Chem. A*, 2018, **6**, 20973–20981.
- 49 O. Musl, S. Galler, G. Wurzer, M. Bacher, I. Sulaeva, I. Sumerskii, A. K. Mahler, T. Rosenau and A. Potthast, *Biomacromolecules*, 2022, 23, 1413–1422.
- 50 M. Gigli and C. Crestini, Green Chem., 2020, 22, 4722-4746.
- 51 M. Österberg, M. H. Sipponen, B. D. Mattos and O. J. Rojas, Green Chem., 2020, 22, 2712–2733.
- 52 J. Chen, J. Tian, N. Feng, L. Ning, D. Wang, B. Zhao, T. Guo, J. Song and O. J. Rojas, *Small*, 2024, 2309756.
- 53 J. Liu, M. Nero, K. Jansson, T. Willhammar and M. H. Sipponen, *Nat. Commun.*, 2023, **14**, 3099.

54 B. Zhao, M. Borghei, T. Zou, L. Wang, L. S. Johansson, J. Majoinen, M. H. Sipponen, M. Österberg, B. D. Mattos and O. J. Rojas, *ACS Nano*, 2021, **15**, 6774–6786.

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- 55 J. Carrier, C.-Y. Lai and D. Radu, *ACS Environ. Au*, 2024, 4, 196–203.
- 56 P. Figueiredo, K. Lintinen, J. T. Hirvonen, M. A. Kostiainen and H. A. Santos, *Prog. Mater. Sci.*, 2018, **93**, 233–269.
- 57 Z. Sun, B. Fridrich, A. de Santi, S. Elangovan and K. Barta, *Chem. Rev.*, 2018, **118**, 614–678.
- 58 S. Laurichesse and L. Avérous, *Prog. Polym. Sci.*, 2014, 39, 1266–1290.
- 59 T. Zou, M. H. Sipponen, A. Henn and M. Österberg, ACS Nano, 2021, 15, 4811–4823.
- 60 I. Standard, Environmental management-Life cycle assessment-Requirements and guidelines, ISO, 2006.
- 61 C. Culbertson, T. Treasure, R. Venditti, H. Jameel and R. Gonzalez, *Nord. Pulp Pap. Res. J.*, 2016, **31**, 30–40.
- 62 C. Moretti, B. Corona, R. Hoefnagels, I. Vural-Gürsel, R. Gosselink and M. Junginger, *Sci. Total Environ.*, 2021, 770, 144656.
- 63 P. T. Anastas and J. C. Warner, *Green chemistry: theory and practice*, Oxford university press, 2000.
- 64 C. Libretti, L. Santos Correa and M. A. R. Meier, *Green Chem.*, 2024, **26**, 4358–4386.
- 65 R. Bardestani, G. S. Patience and S. Kaliaguine, *Can. J. Chem. Eng.*, 2019, 97, 2781–2791.
- 66 S. Brunauer, P. H. Emmett and E. Teller, *J. Am. Chem. Soc.*, 1938, **60**, 309–319.
- 67 M. Thommes, K. Kaneko, A. V. Neimark, J. P. Olivier, F. Rodriguez-Reinoso, J. Rouquerol and K. S. W. Sing, *Pure Appl. Chem.*, 2015, 87, 1051–1069.
- 68 S. Zhang, M. Zheng, Y. Tang, R. Zang, X. Zhang, X. Huang, Y. Chen, Y. Yamauchi, S. Kaskel and H. Pang, Adv. Funct. Mater., 2022, 32, 2204714.
- 69 M. Khosravi, N. Bashirpour and F. Nematpour, *Adv. Mater. Res.*, 2014, 829, 922–926.
- 70 M. B. Vázquez-Santos, E. Geissler, K. László, J.-N. Rouzaud, A. Martínez-Alonso and J. M. D. Tascón, J. Phys. Chem. C, 2012, 116, 257–268.
- 71 A. Carlos, D. L. y Leon and L. R. Radovic, *Chemistry & Physics of Carbon*, 2023, pp. 213–310.
- 72 R. B. Fidel, D. A. Laird and M. L. Thompson, *J. Environ. Qual.*, 2013, **42**, 1771–1778.
- 73 F. Raganati, R. Chirone and P. Ammendola, *Ind. Eng. Chem. Res.*, 2020, **59**, 3593–3605.
- 74 N. S. Wilkins, A. Rajendran and S. Farooq, *Adsorption*, 2021, 27, 397–422.
- 75 N. S. Wilkins, J. A. Sawada and A. Rajendran, *Ind. Eng. Chem. Res.*, 2022, **61**, 7032–7051.
- 76 T. Lu, J. Bai, J. Huang, Q. Yu, M. Demir, M. Kilic, B. N. Altay, L. Wang and X. Hu, *Energy Fuels*, 2023, 37, 3886–3893.
- 77 A. L. Myers and J. M. Prausnitz, *AIChE J.*, 1965, **11**, 121–127.
- 78 K. S. Walton and D. S. Sholl, AIChE J., 2015, 61, 2757-2762.
- 79 S. Sircar and D. V. Cao, *Chem. Eng. Technol.*, 2002, 25, 945-948.

- 80 L. Tian, Y. Zhi, Q. Yu, Q. Xu, M. Demir, S. G. Colak, A. A. Farghaly, L. Wang and X. Hu, *Energy Fuels*, 2024, 38, 13186-13195.
- 81 D. Levesque and F. D. Lamari, Mol. Phys., 2009, 107, 591-597.
- 82 C. Ma, J. Bai, M. Demir, Q. Yu, X. Hu, W. Jiang and L. Wang, Sep. Purif. Technol., 2022, 303, 122299.
- 83 A. Nuhnen and C. Janiak, *Dalton Trans.*, 2020, 49, 10295–10307.
- 84 Y. Chun, Y. Zhu, C. Stubenrauch, Y. Lu and O. J. Rojas, Curr. Opin. Colloid Interface Sci., 2024, 73, 101822.
- 85 W. Cao, H. Xu, X. Zhang, W. Xiang, G. Qi, L. Wan and B. Gao, *Chem. Eng. J.*, 2023, **471**, 144523.
- 86 Z. Pan, G. Qi, X. Zhang, Q. You, Y. Zheng, W. Xiang, Y. Zhao and B. Gao, Sep. Purif. Technol., 2024, 345, 127398.
- 87 Y. Sun, G. Yang, J.-P. Zhang, Y. Wang and M.-S. Yao, *Chem. Eng. Technol.*, 2012, 35, 309–316.
- 88 M. B. Li, X. Liu, C. G. Sun, L. Stevens and H. Liu, *J. Environ. Chem. Eng.*, 2022, **10**, 11.
- 89 G. Yin, Z. Liu, Q. Liu and W. Wu, *Chem. Eng. J.*, 2013, **230**, 133–140.
- 90 D. Saha, G. Orkoulas and D. Bates, Materials, 2023, 16, 12.
- 91 O. Tkachenko, A. Nikolaichuk, N. Fihurka, A. Backhaus, J. B. Zimmerman, M. Strømme and T. M. Budnyak, *ACS Appl. Mater. Interfaces*, 2024, **16**, 3427–3441.
- 92 W. M. Chen, X. Wang, Z. Hashisho, M. Feizbakhshan, P. Shariaty, S. Niknaddaf and X. Y. Zhou, *Microporous Mesoporous Mater.*, 2019, 280, 57–65.
- 93 D. Saha, S. E. Van Bramer, G. Orkoulas, H. C. Ho, J. H. Chen and D. K. Henley, *Carbon*, 2017, **121**, 257–266.
- 94 L. Z. Gong and A. Bao, J. CO2 Util., 2023, 68, 11.
- 95 T. Wang, Y. Zhai, Y. Zhu, C. Li and G. Zeng, Renewable Sustainable Energy Rev., 2018, 90, 223–247.
- 96 F. Bergius, in *Die Anwendung hoher Drucke bei chemischen Vorgängen und eine Nachbildung des Entstehungsprozesses der Steinkohle*, ed. W. Knapp, 1913.
- 97 H. Wikberg, T. Ohra-aho, F. Pileidis and M.-M. Titirici, ACS Sustainable Chem. Eng., 2015, 3, 2737–2745.
- 98 W. Sangchoom and R. Mokaya, *ACS Sustainable Chem. Eng.*, 2015, 3, 1658–1667.
- 99 M. O. Bengoechea, A. Hertzberg, N. Miletić, P. L. Arias and T. Barth, *J. Anal. Appl. Pyrolysis*, 2015, **113**, 713–722.
- 100 W. Hao, F. Björnerbäck, Y. Trushkina, M. Oregui Bengoechea, G. Salazar-Alvarez, T. Barth and N. Hedin, *ACS Sustainable Chem. Eng.*, 2017, 5, 3087–3095.
- 101 M. Demir, T. D. Tessema, A. A. Farghaly, E. Nyankson, S. K. Saraswat, B. Aksoy, T. Islamoglu, M. M. Collinson, H. M. El-Kaderi and R. B. Gupta, *Int. J. Energy Res.*, 2018, 42, 2686–2700.
- 102 E. Atta-Obeng, B. Dawson-Andoh, E. Felton and G. Dahle, Waste Biomass Valorization, 2019, vol. 10, pp. 2725–2731.
- 103 J. Dong, R. Wang, X. Wang, S. Tan, Z. Zhao, Q. Yin and X. Lu, *Carbon*, 2024, 229, 119530.
- 104 J. Zhang, D. Huang, J. Shao, X. Zhang, H. Yang, S. Zhang and H. Chen, *J. Cleaner Prod.*, 2022, 355, 131642.
- 105 Y. Bai, Y. Li, M. Li, X. Lin, C. Cai, H. He and S. Liu, *Ind. Crops Prod.*, 2024, 220, 119183.

106 D. Liu, L. Shao, P. Zhan, L. Zhang, Z. Wu, J. Wang, X. Ma and J. Huang, *Sep. Purif. Technol.*, 2024, 347, 127657.

Chem Soc Rev

- 107 J. Gao, Z.-Q. Wang, B. Li, W. Zhao, Z.-R. Ba, Z.-Y. Liu, J.-J. Huang and Y.-T. Fang, *Bioresour. Technol.*, 2024, 393, 130171.
- 108 S. Park, M. S. Choi and H. S. Park, *Carbon Lett.*, 2019, 29, 289–296.
- 109 J. Zhao, Q. Wang, W. Zhang, D. Shen and C. Cheng, *Int. J. Hydrogen Energy*, 2024, 67, 127–135.
- 110 Q. B. Meng and J. Weber, *ChemSusChem*, 2014, 7, 3312–3318.
- 111 L. S. Shao, N. Liu, L. Z. Wang, Y. F. Sang, H. A. Wan, P. Zhan, L. Zhang, J. H. Huang and J. N. Chen, *Chemosphere*, 2022, 288, 11.
- 112 N. Liu, L. S. Shao, C. Wang, F. B. Sun, Z. P. Wu, P. Zhan, L. Zhang and H. A. Wan, *Int. J. Biol. Macromol.*, 2022, 221, 25–37.
- 113 N. Liu, J. Chen, Z. Wu, P. Zhan, L. Zhang, Q. Wei, F. Wang and L. Shao, *ACS Appl. Polym. Mater.*, 2021, 3, 2178–2188.
- 114 M. A. Karaaslan, L. T. Lin, F. Ko and S. Renneckar, Front. Mater., 2022, 9, 894061.
- 115 M. A. Karaaslan, L. C. Lin, F. Ko and S. Renneckar, presented in part at the American Chemical Society, San Diego, CA, USA, March 20-24, 2022, 2022.
- 116 Z. Li, B. Li, C. Yu, H. Wang and Q. Li, Adv. Sci., 2023, 10, 2206605.
- 117 H. Qin, S. Kang, Y. Huang, S. Liu, Y. Fang, X. Li and Y. Wang, *Mater. Lett.*, 2015, **159**, 463–465.
- 118 S. Otani, Y. Fukuoka, B. Igarashi and K. Sasaki, US3461082A, 1969.
- 119 W. Fang, S. Yang, X.-L. Wang, T.-Q. Yuan and R.-C. Sun, *Green Chem.*, 2017, **19**, 1794–1827.
- 120 E. M. Calvo-Muñoz, F. J. García-Mateos, J. M. Rosas, J. Rodríguez-Mirasol and T. Cordero, *Front. Mater.*, 2016, 3, 14.
- 121 Y. Lu, M. Mehling, S. Huan, L. Bai and O. J. Rojas, *Chem. Soc. Rev.*, 2024, **53**, 7363–7391.
- 122 L. Zhang, L. Jin, B. Liu and J. He, Front. Chem., 2019, 7, 22.
- 123 S. Sani, X. Liu, M. B. Li, L. Stevens and C. G. Sun, *Micro-porous Mesoporous Mater.*, 2023, 347, 11.
- 124 S. Sani, X. Liu, L. Stevens, H. M. Wang and C. G. Sun, *Fuel*, 2023, 351, 11.
- 125 J. Zhao, W. J. Zhang, D. K. Shen, H. Y. Zhang and Z. H. Wang, *J. Energy Inst.*, 2023, **107**, 9.
- 126 S. Geng, J. Wei, S. Jonasson, J. Hedlund and K. Oksman, *ACS Appl. Mater. Interfaces*, 2020, **12**, 7432–7441.
- 127 C. Salas, T. Nypelö, C. Rodriguez-Abreu, C. Carrillo and O. J. Rojas, Curr. Opin. Colloid Interface Sci., 2014, 19, 383–396.
- 128 B. D. Mattos, B. L. Tardy, L. G. Greca, T. Kämäräinen, W. Xiang, O. Cusola, W. L. E. Magalhães and O. J. Rojas, *Sci. Adv.*, 2020, **6**, eaaz7328.
- 129 H. Wang, C. Zeng, C. Wang, J. Fu, Y. Li, Y. Yang, Z. Du, G. Tao, Q. Sun and T. Zhai, *Nat. Mater.*, 2024, 23, 596–603.
- 130 L. Shao, H. a Wan, L. Wang, J. Wang, Z. Liu, Z. Wu, P. Zhan, L. Zhang, X. Ma and J. Huang, Sep. Purif. Technol., 2023, 313, 123440.

- 131 S.-C. Qi, Y. Liu, A.-Z. Peng, D.-M. Xue, X. Liu, X.-Q. Liu and L.-B. Sun, *Chem. Eng. J.*, 2019, 361, 945–952.
- 132 L. S. Shao, H. A. Wan, L. Z. Wang, J. J. Wang, Z. H. Liu, Z. P. Wu, P. Zhan, L. Zhang, X. C. Ma and J. H. Huang, Sep. Purif. Technol., 2023, 313, 15.
- 133 G. Singh, J. Lee, A. Karakoti, R. Bahadur, J. B. Yi, D. Y. Zhao, K. AlBahily and A. Vinu, *Chem. Soc. Rev.*, 2020, 49, 4360–4404.
- 134 C. Goel, S. Mohan and P. Dinesha, *Sci. Total Environ.*, 2021, 798, 21.
- 135 V. K. Singh and E. A. Kumar, *Appl. Therm. Eng.*, 2016, 97, 77–86.
- 136 N. Planas, A. L. Dzubak, R. Poloni, L. C. Lin, A. McManus, T. M. McDonald, J. B. Neaton, J. R. Long, B. Smit and L. Gagliardi, J. Am. Chem. Soc., 2013, 135, 7402–7405.
- 137 G. K. Cui, J. J. Wang and S. J. Zhang, *Chem. Soc. Rev.*, 2016, 45, 4307–4339.
- 138 Y. Q. Chen, X. Zhang, W. Chen, H. P. Yang and H. P. Chen, *Bioresour. Technol.*, 2017, **246**, 101–109.
- 139 A. E. Creamer and B. Gao, *Environ. Sci. Technol.*, 2016, **50**, 7276–7289.
- 140 M. E. Boot-Handford, J. C. Abanades, E. J. Anthony, M. J. Blunt, S. Brandani, N. Mac Dowell, J. R. Fernández, M. C. Ferrari, R. Gross, J. P. Hallett, R. S. Haszeldine, P. Heptonstall, A. Lyngfelt, Z. Makuch, E. Mangano, R. T. J. Porter, M. Pourkashanian, G. T. Rochelle, N. Shah, J. G. Yao and P. S. Fennell, *Energy Environ. Sci.*, 2014, 7, 130–189.
- 141 G. Banvillet, S. Pritchard, J. J. Kaschuk, X. Shi, M. Imani, Y. Lu, A. Takagi, M. Kamkar and O. J. Rojas, *Mater. Today Nano*, 2023, 24, 100424.
- 142 M. Samandari, M. T. Broud, D. P. Harper and D. J. Keffer, *J. Phys. Chem. B*, 2024, **128**, 8530–8545.
- 143 M. T. Broud, M. Samandari, L. Yu, D. P. Harper and D. J. Keffer, J. Phys. Chem. C, 2023, 127, 13639–13650.
- 144 R. W. Flaig, T. M. Osborn Popp, A. M. Fracaroli, E. A. Kapustin, M. J. Kalmutzki, R. M. Altamimi, F. Fathieh, J. A. Reimer and O. M. Yaghi, J. Am. Chem. Soc., 2017, 139, 12125–12128.
- 145 D. Y. C. Leung, G. Caramanna and M. M. Maroto-Valer, Renewable Sustainable Energy Rev., 2014, 39, 426–443.
- 146 X. Ma, Y. Yang, Q. Wu, B. Liu, D. Li, R. Chen, C. Wang, H. Li, Z. Zeng and L. Li, Fuel, 2020, 282, 118727.
- 147 Z. Zhang, A. L. Kummeth, J. Y. Yang and A. N. Alexandrova, *Proc. Natl. Acad. Sci. U. S. A.*, 2022, **119**, e2123496119.
- 148 A. Lancheros, S. Goswami, M. R. Mian, X. Zhang, X. Zarate, E. Schott, O. K. Farha and J. T. Hupp, *Dalton Trans.*, 2021, 50, 2880–2890.
- 149 Y. Lu, D. Sun, J. Ralston, Q. Liu and Z. Xu, J. Colloid Interface Sci., 2019, 557, 185–195.
- 150 M. Zhang, Y. Nan, Y. Lu, Q. You and Z. Jin, Fuel, 2023, 331, 125773.
- 151 T. Lu and Q. Chen, J. Phys. Chem. A, 2023, 127, 7023-7035.
- 152 L. Petridis, R. Schulz and J. C. Smith, *J. Am. Chem. Soc.*, 2011, **133**, 20277–20287.
- 153 A. S. Patri, B. Mostofian, Y. Pu, N. Ciaffone, M. Soliman, M. D. Smith, R. Kumar, X. Cheng, C. E. Wyman, L. Tetard,

A. J. Ragauskas, J. C. Smith, L. Petridis and C. M. Cai, *J. Am. Chem. Soc.*, 2019, **141**, 12545–12557.

Review Article

- 154 S. V. Pingali, M. D. Smith, S.-H. Liu, T. B. Rawal, Y. Pu, R. Shah, B. R. Evans, V. S. Urban, B. H. Davison, C. M. Cai, A. J. Ragauskas, H. M. O'Neill, J. C. Smith and L. Petridis, *Proc. Natl. Acad. Sci. U. S. A.*, 2020, 117, 16776–16781.
- 155 Z. Nezafat, M. Nasrollahzadeh, S. Javanshir, T. Baran and Y. H. Dong, *Green Chem.*, 2023, 25, 9603–9643.
- 156 Y. D. Cui, B. He, Y. Lei, Y. Liang, W. T. Zhao, J. Sun and X. M. Liu, *Chin. J. Chem. Eng.*, 2023, 54, 89–97.
- 157 S.-C. Qi, J.-K. Wu, J. Lu, G.-X. Yu, R.-R. Zhu, Y. Liu, X.-Q. Liu and L.-B. Sun, J. Mater. Chem. A, 2019, 7, 17842–17853.
- 158 S.-C. Qi, X.-J. Lu, Y.-C. Lou, R. Zhou, D.-M. Xue, X.-Q. Liu and L.-B. Sun, *Engineering*, 2022, **16**, 154–161.
- 159 A.-Z. Peng, S.-C. Qi, X. Liu, D.-M. Xue, S.-S. Peng, G.-X. Yu, X.-Q. Liu and L.-B. Sun, *Chem. Eng. J.*, 2019, 372, 656–664.
- 160 S. Beck, P. Choi and S. H. Mushrif, *Phys. Chem. Chem. Phys.*, 2022, **24**, 20480–20490.
- 161 S. Beck and S. H. Mushrif, Org. Biomol. Chem., 2023, 21, 4596–4600.
- 162 R. L. Silveira, S. R. Stoyanov, S. Gusarov, M. S. Skaf and A. Kovalenko, J. Phys. Chem. Lett., 2015, 6, 206–211.
- 163 J. V. Vermaas, M. F. Crowley and G. T. Beckham, *ACS Sustainable Chem. Eng.*, 2020, **8**, 17839–17850.
- 164 D. Vural, J. C. Smith and L. Petridis, *Phys. Chem. Chem. Phys.*, 2018, 20, 20504–20512.
- 165 T. Zhang, X. Li, L. Guo and X. Guo, Energy Fuels, 2019, 33, 11210–11225.
- 166 P. N. Ciesielski, M. B. Pecha, A. M. Lattanzi, V. S. Bharadwaj, M. F. Crowley, L. Bu, J. V. Vermaas, K. X. Steirer and M. F. Crowley, ACS Sustainable Chem. Eng., 2020, 8, 3512–3531.
- 167 M. J. Orella, T. Z. H. Gani, J. V. Vermaas, M. L. Stone, E. M. Anderson, G. T. Beckham, F. R. Brushett and Y. Román-Leshkov, ACS Sustainable Chem. Eng., 2019, 7, 18313–18322.
- 168 J. V. Vermaas, L. D. Dellon, L. J. Broadbelt, G. T. Beckham and M. F. Crowley, *ACS Sustainable Chem. Eng.*, 2019, 7, 3443–3453.
- 169 Z. Liu, T. Lu and Q. Chen, Carbon, 2021, 171, 514-523.
- 170 S. Anila, C. H. Suresh and H. F. Schaefer, III, *J. Phys. Chem. A*, 2022, **126**, 4952–4961.
- 171 H. M. Lee, I. S. Youn, M. Saleh, J. W. Lee and K. S. Kim, *Phys. Chem. Chem. Phys.*, 2015, 17, 10925–10933.
- 172 K. Kitaura and K. Morokuma, *Int. J. Quantum Chem.*, 1976, **10**, 325–340.
- 173 P. Su, Z. Jiang, Z. Chen and W. Wu, *J. Phys. Chem. A*, 2014, **118**, 2531–2542.

- 174 E. D. Glendening, J. Am. Chem. Soc., 1996, 118, 2473-2482.
- 175 P. R. Horn, Y. Mao and M. Head-Gordon, *Phys. Chem. Chem. Phys.*, 2016, **18**, 23067–23079.
- 176 K. Szalewicz, Wiley Interdiscip. Rev.: Comput. Mol. Sci., 2012, 2, 254–272.
- 177 E. I. Izgorodina, J. L. Hodgson, D. C. Weis, S. J. Pas and D. R. MacFarlane, *J. Phys. Chem. B*, 2015, **119**, 11748–11759.
- 178 J.-Y. Chung, U. Hwang, J. Kim, N.-Y. Kim, J. Nam, J. Jung, S.-H. Kim, J. K. Cho, B. Lee, I.-K. Park, J. Suhr and J.-D. Nam, *ACS Sustainable Chem. Eng.*, 2023, 11, 2303–2313.
- 179 J. Sun, C. Wang, L. P. Stubbs and C. He, *Macromol. Mater. Eng.*, 2017, 302, 1700341.
- 180 H. Sidky, W. Chen and A. L. Ferguson, *Mol. Phys.*, 2020, 118, e1737742.
- 181 Y. Wang, J. M. Lamim Ribeiro and P. Tiwary, Curr. Opin. Struct. Biol., 2020, 61, 139–145.
- 182 F. Noé, A. Tkatchenko, K.-R. Müller and C. Clementi, *Annu. Rev. Phys. Chem.*, 2020, **71**, 361–390.
- 183 S. Bhakat, RSC Adv., 2022, 12, 25010-25024.
- 184 R. Jones, M. Khalkhali, Y. Lu, Q. Liu, A. Ghosh and R. Lespiat, US Pat., US20240025807A1, 2024.
- 185 G. Tofani, E. Jasiukaitytė-Grojzdek and B. Likozar, *BioResources*, 2024, **19**, 6967–6969.
- 186 J. Zhang, Y.-K. Lei, Z. Zhang, J. Chang, M. Li, X. Han, L. Yang, Y. I. Yang and Y. Q. Gao, *J. Phys. Chem. A*, 2020, 124, 6745–6763.
- 187 L. Bonati, G. Piccini and M. Parrinello, *Proc. Natl. Acad. Sci. U. S. A.*, 2021, **118**, e2113533118.
- 188 D. Ray, E. Trizio and M. Parrinello, J. Chem. Phys., 2023, 158, 204102.
- 189 A. Rajendran, G. K. H. Shimizu and T. K. Woo, Adv. Mater., 2024, 36, 2301730.
- 190 G. Zhao, Z. Li, B. Cheng, X. Zhuang and T. Lin, Sep. Purif. Technol., 2023, 315, 123754.
- 191 L.-G. Ding, B.-J. Yao, F. Li, S.-C. Shi, N. Huang, H.-B. Yin, Q. Guan and Y.-B. Dong, *J. Mater. Chem. A*, 2019, 7, 4689–4698.
- 192 T. Nguyen, G. Shimizu and A. Rajendran, Chemrxiv, 2024, DOI: 10.26434/chemrxiv-2024-nx3vg.
- 193 P. Tan, Y. Jiang, Q. Wu, C. Gu, S. Qi, Q. Zhang, X. Liu and L. Sun, *Chin. J. Chem. Eng.*, 2022, 42, 104–111.
- 194 Y. Qiao, J. J. Bailey, Q. Huang, X. Ke and C. Wu, *Renewable Sustainable Energy Rev.*, 2022, **158**, 112079.
- 195 Z. Pan, A. Bodi, J. A. van Bokhoven and P. Hemberger, *Phys. Chem. Chem. Phys.*, 2022, **24**, 21786–21793.
- 196 Z. Pan, X. Wu, A. Bodi, J. A. van Bokhoven and P. Hemberger, *Green Chem.*, 2024, **26**, 9899–9910.