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1. Introduction

In the last 20 years, the amount of atmospheric carbon dioxide $(CO₂)$, a potent greenhouse gas, has exhibited a steady increase, as documented by Goel (2021).¹ This increase can be attributed to multiple factors, including the repercussions of the industrial revolution, characterized by an annual increment of 0.17 °C, and burning of fossil fuels. Industries such as those involved in energy production from coal, oil, and natural gases; cement manufacturing; chemical synthesis; metal production; and refineries significantly contribute to $CO₂$ emissions. Consequently, the atmospheric $CO₂$ concentration reached 423 parts per million (ppm) in 2023, surpassing the anticipated level of around 400 ppm or 0.04% of $CO₂$ in the air by volume.² This surge represents an approximately 50% increase since the preindustrial era and an additional 13% since the turn of the millennium. The escalating rate of $CO₂$ emissions raises

A comprehensive review of enhanced $CO₂$ capture using activated carbon derived from biomass feedstock

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The increasing level of atmospheric $CO₂$ requires the urgent development of effective capture technologies. This comprehensive review thoroughly examines various methods for the synthesis of carbon materials, modification techniques for converting biomass feedstock into carbon materials and pivotal factors impacting their properties. The novel aspect of this review is its in-depth comparison of how these modifications specifically affect the pore structure and surface area together with the exploration of the mechanism underlying the enhancement of $CO₂$ adsorption performance. Additionally, this review addresses research gaps and provides recommendations for future studies concerning the advantages and drawbacks of $CO₂$ adsorbents and their prospects for commercialization and economic feasibility. This article revealed that among the various strategies, template carbonization offers a viable option for providing control of the material pore diameter and structure without additional modification treatments. Optimizing the pore structure of activated carbons, particularly those activated with agents such as KOH and ZnCl₂, together with synthesizing hybrid activated carbons using multiple activating agents, is crucial for enhancing their $CO₂$ capture performance. Cost-benefit analysis suggests that biomass-derived activated carbons can significantly meet the escalating demand for $CO₂$ capture materials, offering economic advantages and supporting sustainable waste management. **EXAMPLE THE SECTION CONSULTER AND SURFAINTED CONSULTER THE SECTION CONSULTER CONSULTER CONSULTER AND A COMPREHENSIVE TEXTIFIED AND SURFAINTED CREATED THE UNIT CONSULTER THE CONSULTER THE CONSULTER CONSULTER AND CONSULTER**

concerns on various fronts, primarily because of its pivotal role in climate change and its adverse impact on human health. Manifestations of these impacts include respiratory complications, elevated blood pressure, and accelerated heart rates among individuals. Thus, to lower $CO₂$ emissions, several carbon capture systems have been developed. Numerous initiatives focus on mitigating climate change by capturing and separating $CO₂$, particularly from large power plants and the atmosphere, using different technologies such as membranes, absorption, microbial methods, cryogenics, and chemical looping. However, these technologies often face challenges such as high operational costs and substantial energy consumption. Fortunately, physical processes such as adsorption offer a cost-effective solution and have been widely used to capture various contaminants, including CO₂.³ Despite the initial skepticism, the need to develop green technologies has become increasingly evident, given the incontrovertible link between $CO₂$ emissions, climate change, and the resulting health risks. Therefore, the development and adoption of effective and socially acceptable solutions are imperative. Addressing the challenge of $CO₂$ emissions necessitates innovative approaches and concerted efforts to safeguard both the environment and human well-being.

Accordingly, adsorption has emerged as a viable and costeffective approach for the sequestration and storage of

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carbon. Among the many absorbents used, those that stand out include polymers, carbon compounds, metal oxides, aminebased materials, zeolites, alumina, silica, and metal–organic frameworks (MOFs). In this case, zeolites exhibit sensitivity to moisture, adsorption temperature, and pressure, whereas carbon materials offer notable advantages. Carbon materials, including hydrocarbons, charcoal, carbon nanotubes, graphite, graphene, fullerene, and activated carbon, are simple to prepare and control. Efficient $CO₂$ capture is possible due to their welldeveloped pore architectures and high specific surface area. Unlike MOFs, carbon materials are not moisture sensitive, making them more reliable for industrial applications.⁴ Therefore, it is necessary to develop carbon materials with remarkable capabilities for capturing $CO₂$, given their abundant and economical sources. The process for the preparation of carbon materials is straightforward and easily manageable. Their substantial specific surface area and well-formed pore structures significantly enhance their capacity for capturing $CO₂$ effectively. Furthermore, carbon materials exhibit resistance to moisture, unlike their alternatives. Additionally, activated carbon demonstrates the ability to function across a broad temperature range.

Activated carbons (AC) have emerged as incredibly encouraging adsorbents for $CO₂$ capture due to their significantly enhanced physiochemical, mechanical, thermal, and chemical stability, as depicted in Fig. 1. Although numerous porous carbon-based materials exist, porous carbonaceous materials derived from biomass hold particular appeal as $CO₂$ adsorbents. The preference for biomass as a material is driven by its excellent textural characteristics, high $CO₂$ adsorption capacity, and ease of synthesis, combined with its cost-effectiveness and the renewable nature of biomass resources.⁵ Biomass, which is derived from both plant and animal origins, captures solar energy through photosynthesis, making it a sustainable energy source that can be used for heating or transformed into liquid and gaseous fuels using various methods. Among the diverse biomass sources, traditional options such as bagasse, algae, municipal waste, wet waste, forestry residues, agricultural crop residues, and wood processing by-products stand out due to

their potential to reduce costs. Pyrolysis stands out as the predominant thermochemical technology for biomass conversion, wherein biomass undergoes decomposition into biochar, tar, and gas. Traditionally, biomass is subjected to pyrolysis, a thermal treatment conducted at moderate temperatures without oxygen, resulting in the production of biochar.⁶ Pyrolysis methods include different approaches such as slow or fast pyrolysis, low or high-temperature conditions, and dry or wet pyrolysis.⁷ A variety of biomass sources, including wood, nonwood materials, and agricultural or fruit residues, has been utilized for the production of charcoal. The creation of activated porous carbons, similar to biochar, involves the addition of an activating agent either before or after the biochar is produced. An efficient technique for producing activated porous carbon has been developed using Arundo donax biomass, which was activated in a single step with KOH.⁸ This innovative approach simplifies the conventional two-step activation process, thereby reducing the time constraints. Various activating agents, whether physical or chemical, are utilized in the production of these carbons, including steam, $CO₂$, KOH, ZnCl₂, H₃PO₄, $HNO₃$, and $H₂O₂$.⁹ Changes in the pyrolysis conditions affect the yield, specific surface area, and surface chemistry of biochar. For instance, Ighalo et al. conducted pyrolysis at temperatures ranging from 250 °C to 650 °C, demonstrating an increase in specific surface area of the biochar with an increase in temperature, together with a decrease in its yield, as evidenced by BET data.¹⁰ The composition and quantity of functional groups on the surface of carbon materials are influenced by their preparation techniques and the characteristics of the employed biomass sources. Fig. 2 illustrates the formation of carbon fibers, activated porous carbons, and other byproducts that arise from the combustion of various biomass types, together with their potential applications across different industries.¹¹ **PSC Advances**

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Several studies have shown that unmodified biochar has limited effectiveness in absorbing contaminants, leading to the investigation of various modification techniques, including reactions with steam, acid or alkaline treatments, and doping with heteroatoms.¹² To significantly improve the

Fig. 1 Properties of activated carbon for carbon dioxide capture.

Fig. 2 Illustration of the process of producing biochar and activated carbons from biomass together with their potential applications across different industries. Reproduced from ref. 11.

ability of carbon sorbents to capture $CO₂$, it is essential to introduce basic sites and increase the porosity of the material. Nitrogen doping, a prevalent modification technique, involves the incorporation of nitrogen-containing chemicals such as polyethyleneimide (PEI) and triethylenephosphoramide (TEPA) into carbon materials, resulting in the introduction of pyridinic N or pyrrolic N groups, which both play pivotal roles in $CO₂$ capture.^{13,14} For example, incorporating chitosan into carbonized materials has been shown to significantly improve their CO₂ adsorption capacity, achieving 5.83 mmol g⁻¹ at 0 ° C.¹⁵ Additionally, metal oxides or salts are used to modify carbon materials, allowing the introduction of alkaline metal oxides or hydroxides. This modification increases the alkalinity of carbon materials, thereby enhancing their ability to adsorb acidic $CO₂$. Research has demonstrated that biochar derived from walnut shells and impregnated with metal nitrates, particularly magnesium–biochar combinations, exhibits superior $CO₂$ adsorption capabilities compared to other metal–biochar composites (such as that with aluminum, iron, nickel, calcium, or sodium).^{1,16} However, despite the progress in modification techniques and their application in $CO₂$ capture, there is still a lack of comprehensive reviews on the methods for the preparation and modification of biomassbased carbon materials. Thus, this review provides a critical analysis of the preparation technologies and conversion conditions that influence the properties of carbon materials derived from biomass. Also, the emerging techniques for modifying these carbon materials, their applications in $CO₂$ capture, and the underlying capture mechanisms are explored. Finally, the potential future directions and challenges for research in this field are highlighted.

2. Navigating thermochemical conversion technologies for biomass

A wide range of thermochemical conversion methods designed for utilizing biomass has been investigated, including pyrolysis, gasification, and hydrothermal processes. This study assesses their effectiveness in converting biomass into valuable energy resources, with a focus on environmental sustainability and resource optimization.

2.1 Conventional pyrolysis

Pyrolysis, a thermochemical process devoid of oxygen, is instrumental in converting biomass into valuable products such as char, oil, and syngas through high-temperature heating.¹⁷ This process is highly versatile, encompassing variations such as fast, flash, and slow pyrolysis, which are categorized based on the temperature and heating rate. Typically, natural biomass derived from plants primarily consists of three key biopolymers, i.e., cellulose (making up 35–50% of the biomass), hemicellulose (comprising 25–30%), and lignin (constituting 10–30%), with a small fraction of mineralized inorganic components.18,19 These components differ in their chemical structures and undergo distinct, complex reactions during pyrolysis. Among the various pyrolytic conditions, temperature is regarded as the most critical factor, given that it greatly impacts the final textural properties and surface functionalities of biochar. Fig. 3 depicts the thermal decomposition behavior of these components over a wide temperature range, highlighting the specific temperature intervals at which each component undergoes thermal breakdown. The intricate mechanism of pyrolysis involves sequential decomposition stages of biomass constituents. Initially, water evaporation occurs, followed by the degradation of hemicellulose at around

Fig. 3 Decomposition temperatures for lignin, cellulose, and hemicellulose in biomass during thermal processing

200–350 °C. Subsequently, cellulose undergoes decomposition with an increase in temperature, while lignin is decomposed at temperatures exceeding 400 °C. These temperature-dependent reactions are pivotal in determining the final properties and functionalities of the resulting biochar. Among the pyrolytic conditions, temperature stands out as a critical parameter influencing the texture and surface characteristics of biochar. Slow pyrolysis, which operates at temperatures between 300– 600 °C and with low heating rates of 0.1-1 °C s⁻¹, primarily produces biochar, with the typical yields in the range of 30– 50%.²⁰ In contrast, fast and flash pyrolysis methods are designed to efficiently produce bio-oil. These methods are characterized by extremely high heating rates, such as 1000 °C $\rm s^{-1}$, and very short residence times. 21 The pyrolysis cycle can be comprehensively understood through four distinct stages, as elucidated by the thermogravimetric analysis (TGA) of biomass. These stages involve moisture removal, hemicellulose degradation, cellulose decomposition, and lignin degradation, occurring in specific temperature ranges. Thus, a comprehensive understanding of these stages is crucial for optimizing the properties of biochar and activated porous carbons. This knowledge allows better control of the product selectivity, particularly in balancing the production of char, bio-oil, and syngas.²²

The initial phase of pyrolysis involves removing moisture entirely at temperatures up to 220 °C, followed by the disintegration of hemicellulose in the temperature range of 220–315 ° C. Subsequently, cellulose decomposes in the range of 315– 400 °C, and lignin degradation occurs beyond 400 °C.²³ Understanding the transformations these biopolymers undergo during pyrolysis is crucial for gaining deep insight into the overall mechanism. This deep understanding not only enables an improvement in the inherent properties of biochar and precise control of the selectivity for products such as char, biooil, and syngas, as well as activated porous carbons, but it also

aids in selecting the most suitable biomass sources from a diverse range of naturally occurring feedstocks.²⁴ Fig. 4 illustrates the quantities of biochar, syngas, and bio-oil produced under various pyrolysis conditions. Fast and flash pyrolysis techniques yield significantly more bio-oil than slow pyrolysis, mainly because the shorter residence time minimizes secondary cracking reactions, thereby increasing the bio-oil yield. Maintaining the optimal pyrolysis temperature is crucial for maximizing bio-oil production, given that higher temperatures typically favor its generation. However, conventional pyrolysis methods have inherent drawbacks, such as low heat transfer efficiency and prolonged heating times, resulting in high costs. In contrast, microwave pyrolysis offers rapid heating rates and shorter reaction times, thereby enhancing the productivity and reducing expenses. Additionally, water, a strong absorber of microwaves, enables the direct microwavepyrolysis of high-moisture biomass without pretreatment. Comparative studies indicate that microwave pyrolysis results in biochar with a greater specific surface area and higher

Fig. 4 Proportional breakdown of char, bio-oil, and syngas produced through various pyrolysis methods.

Table 1 Influence of pyrolysis parameters on the generation of biochar, bio-oil, and syngas

porosity compared to conventional pyrolysis under similar temperature conditions. However, conventional pyrolysis tends to produce a higher yield of char. This is because conventional methods often lead to reduced yields of biochar and bio-oil due to the increased occurrence of biomass cracking and gas formation.

Table 1 shows the influence of the pyrolysis parameters on the generation of biochar, bio-oil, and syngas. Additionally, these insights will aid in identifying suitable biomass sources from the diverse array of naturally occurring biomass feedstocks. In summary, elucidating the mechanism of pyrolysis and its intricate stages not only improves the understanding of biochar production but also facilitates the development of efficient thermochemical conversion processes for the utilization of biomass, contributing to sustainable resource management and energy production.

2.2 Microwave pyrolysis

Microwave pyrolysis has emerged as a promising strategy for carbonizing raw materials, offering solutions to the challenges associated with conventional pyrolysis methods. Analogous to commercial microwave applications in daily life, this approach transfers electromagnetic energy to the precursor material, effecting rapid and targeted heating. Microwave pyrolysis offers distinct advantages compared to traditional furnaces, primarily due to its capability for precise and rapid heating, which results in a highly energy-efficient system. Additionally, carbon-based compounds with high microwave absorbance enable the selective absorption of microwave radiation, allowing the targeted heating of the reactants and promoting specific chemical reactions. Consequently, microwave pyrolysis is expected to

result in higher char yields and maintain a more stable temperature profile.²⁵ These attributes contribute to enhanced production rates and optimal production costs, underscoring the potential of microwave pyrolysis as a transformative approach in biomass conversion processes. Table 2 summarizes the key differences between conventional and microwave pyrolysis based on insights gathered from studies conducted by various research groups.^{26,27} The key advantage of microwave pyrolysis is its unique heating mechanism. Unlike traditional methods, which rely on convection and conduction through heated coils to transfer heat to materials, often facing limitations due to the thermal properties and surface temperature of materials, microwave pyrolysis uses the interaction between microwave energy and the dipoles in the target material to generate heat internally. This internal heating process makes the system more efficient than traditional surface heating, given that it facilitates uniform heat distribution and consistent heating throughout the material. Consequently, microwave pyrolysis is recognized for its energy efficiency, rapid processing, and time-saving procedures, facilitating high heating rates and temperatures.²⁸ These advantages contribute to the superior reputation of microwave pyrolysis compared to traditional pyrolysis methods. Review Workers are the proposition of the characteristic on 18 2024. The method of the common control of the common common

Pecan nutshells were used as precursors for producing activated carbon through a one-step process. The nutshells were treated and mixed with two different chemical agents, potassium carbonate and potassium hydroxide, in a 1:1 ratio. Subsequently, this mixture was subjected to microwave pyrolysis at power levels of 300 W and 400 W for periods ranging from 2 to 6 min. Notably, microwave pyrolysis significantly improved the formation of microporous structures, especially

in the samples activated with potassium hydroxide. For example, the samples pyrolyzed at 300 W for 6 min exhibited more than 73% ultra-microporosity. Additionally, these samples demonstrated an excellent $CO₂$ adsorption performance, achieving 5.3 mmol g^{-1} at 1 bar and 0 °C. This level of performance was comparable to that of the activated carbon synthesized through conventional methods reported in the literature, underscoring the significant potential of microwave pyrolysis for future application in the production of activated carbon. Vairaja et al. investigated the performance disparities between conventional and microwave pyrolysis in producing AC.²⁹ Both methods were employed to fabricate carbon activated from the husk of sunflower seeds. Following washing and drying, the precursor material was impregnated with varying concentrations of $ZnCl₂$ and subjected to microwave carbonization at power levels ranging from 200 to 1000 W for durations spanning 5 to 40 min. Subsequently, the samples underwent conventional treatment. The results indicated that the highest the activated carbon surface area was achieved at 600 W microwave power. In a separate study, Ali and Idris examined the influence of activation processes on the characteristics of synthesized activated carbon.³⁰ Pistachio shells were combined with different concentrations of K_2CO_3 , KOH, ZnCl₂, and $H₂SO₄$, and the impact of the microwave power level in the range of 450 to 800 W was assessed. The optimal yields were obtained using a microwave power of 600 W with 15 min of irradiation when K_2CO_3 was used as the activation agent. Additionally, the surface area of the activated carbon increased by nearly 600% when the power was increased from 450 W to 600 W. However, this increase was followed by a decline of 27% and 58% in surface area as the power was further increased to 700 W and 800 W, respectively. Microwave pyrolysis shows promising results in biochar production due to its high yield, reduced energy requirements, and shorter activation times. Despite these advantages, the use of microwave technology for the synthesis of activated carbon is still limited to laboratoryscale applications, largely due to the technical challenges associated with scaling up to industrial levels. **PSC** Advances

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2.3 Mechanism of pyrolysis

In elucidating the mechanism of pyrolysis, it is crucial to delineate the sequential transformations that three key biopolymers undergo throughout the pyrolysis cycle. Understanding the intricate processes involved not only sheds light on the fundamental principles governing pyrolysis but also aids in identifying the optimal biomass sources for extraction. By comprehending the underlying mechanisms, researchers can discern the most suitable biomass substrates from the vast array of naturally occurring sources, thereby facilitating the efficient conversion of biomass into valuable products. This elucidation of the pyrolysis mechanisms not only enhances our understanding of bioenergy production but also contributes to sustainable resource utilization.

2.3.1 Cellulose. Cellulose is a polymer formed through the linking of p-glucose units through β -(1-4) glycosidic bonds, creating a continuous network of chains that can range from

intramolecular and intermolecular hydrogen bonds between the –OH groups results in linear, parallel, and crystalline structures known as microfibrils. These microfibrils are aligned longitudinally within the cell walls of biomass materials, providing rigidity and strength to their cellular structures. Thus, understanding the pyrolysis mechanism of cellulose into biochar is crucial for leveraging this thermochemical process for the effective conversion of biomass. During pyrolysis, cellulose undergoes a sequence of intricate thermal decomposition reactions, ultimately yielding biochar as the primary product. Initially, cellulose undergoes dehydration, liberating water vapor as it disintegrates into smaller molecular fragments. Subsequently, these fragments undergo depolymerization and cross-linking reactions, leading to the generation of a complex mixture of volatile compounds, including gases and organic vapors, together with solid char residues. The pyrolysis process is initiated with the slow pyrolysis of cellulose at temperatures below 250 °C, accompanied by a reduction in the polymerization degree and the release of H_2O , CO_2 , and CO. Above 250 °C, cellulose transforms into tar, which is predominantly comprised of organic compounds, while leaving behind a charred black residue. Mechanistically, the process begins with solid cellulose depolymerizing through the breaking of the glucosidic bond, resulting in the formation of a bicyclic compound, levoglucosan (LGA).³¹ Subsequent dehydration and isomerization of LGA produce levoglucosenone (LGO), as well as $1,4:3,6$ -dianhydro- β -D-glucopyranose (DGP) and $1,6$ -anhydro- β -D-glucofuranose (AGF).³² Both DGP and AGF undergo conversion to LGO through a dehydration reaction. LGA undergoes successive dehydration and rearrangement reactions, resulting in the generation of furan derivatives such as 5-methylfurfural, 2,3-butanedione, furfural, hydroxymethylfurfural, glycolaldehyde, and glyceraldehyde.³³ Further rearrangements, including aromatization, condensation, and polymerization, lead to the formation of a network of carbon matrix commonly known as biochar. Accordingly, understanding these intricate mechanisms sheds light on optimizing the production of biochar and its potential applications in various fields.

several hundred to thousands of units. The formation of

The presence of hydroxyl groups (–OH) plays a pivotal role in the formation of biochar, particularly influencing its hydrophilic surface properties, which are crucial for facilitating the dispersion of metal species. This investigation specifically addresses the relationship between the functionalities of biochar and its polarities by subjecting cellulose to pyrolysis at temperatures in the range of 200 °C to 700 °C, and subsequently characterizing the resulting biochar. The findings revealed that -OH, rather than the carbon–oxygen $(C=O)$ and carbon–oxygen–carbon $(C=O)$ O–C) functionalities, signicantly contribute to the formation of biochar with a hydrophilic surface. Furthermore, this study identified the temperature of 440 $\mathrm{^{\circ}C}$ as crucial, marking the peak abundance of hydroxyl groups (–OH) and aliphatic carbon– hydrogen (C–H) bonds. Beyond this temperature threshold in the range of 460 °C to 700 °C, a notable transition from oxygen-rich functionalities to carbon-rich functionalities occurs, with the dominant aromatization process leading to a substantial increase in the hydrophobicity of the biochar. The hydrophilic

Fig. 5 Schematic showing the relationship between hydrophilicity and functional development of biochar. Reproduced from ref. 34.

nature of the biochar surface is vital in its utility as a support material for enhancing the dispersion of metals, such as copper (Cu), as demonstrated by the creation of bonding sites conducive to chelating with Cu^{2+} (Fig. 5).³⁴

2.3.2 Hemicellulose. Hemicellulose, the second most abundant component in biomass-based materials, is crucial for maintaining the structural integrity of cell walls. It consists of a variety of heteropolymerized saccharides, including xylan, glucoxylan, glucomannans, and glucuronoxylan. Its diverse composition arises from monomer units such as xylose, glucose, galactose, mannose, and arabinose.³⁵ This heterogeneity gives rise to the amorphous nature of hemicellulose, providing essential support to cellulose. Upon pyrolysis, the decomposition of hemicellulose is initiated through depolymerization, yielding oligosaccharides similar to cellulose. Glucomannans and xylan, for instance, breakdown into various

intermediates, including levoglucosenone, 1,6-anhydro-a-Dgalactopyranose, levomannosan, 1,6-anhydro-b-D-mannopyranose, and 1,4-anhydro-p-xylopyranose.³⁶ These intermediates undergo additional changes into syngas and bio-oil products or directly participate in chemical reactions, leading to the formation of biochar networks. Aromatization, dehydration, and decarboxylation reactions dominate the conversion process, facilitating the formation of stable biochar structures (Fig. 6). Understanding the intricate mechanism of hemicellulose pyrolysis is crucial for optimizing the biomass conversion processes and harnessing biochar as a sustainable resource for various applications. Moreover, elucidating the pathways involved in the decomposition of hemicellulose sheds light on the fundamental principles governing biomass pyrolysis, paving the way for advancements in bioenergy production and environmental sustainability.

Fig. 6 Illustration of the breakdown mechanism of hemicellulose during pyrolysis and subsequent biochar formation.

2.3.3 Lignin. Lignin is an intricate organic polymer composed of p-hydroxyphenol, syringyl, and guaiacyl units, exhibiting a highly variable three-dimensional amorphous structure due to the differences in relative proportions and crosslinkages through -O- and C-C linkages.³⁷ Functional groups such as $-OCH_3$, OH, COOH, and C=O confer high polarity to lignin molecules. The pyrolysis of lignin is a complex process occurring over a wide temperature range owing to its diverse structure. The majority of interconnections in lignin involve β -O-4 linkages, constituting about 60% of its structure.³⁸ Upon pyrolysis, these linkages undergo cleavage, generating free radical species. Subsequent radical reactions lead to chain polymerization, ultimately leading to the production of a biochar network.³⁹ Moreover, methane gas is released during pyrolysis when the methoxy groups are broken. The intricate nature of biomass, combined with the distinct behavior of its constituent biopolymers at varying temperatures during pyrolysis, poses a signicant challenge in elucidating the precise mechanism of the entire process. Furthermore, understanding the process by which activated porous carbons are derived from biomass presents an even greater challenge. Despite these complexities, previous review articles have attempted to describe the pyrolysis process, highlighting the ongoing efforts to unravel the intricate mechanisms underlying biomass conversion and biochar formation.³⁶ **PSC Advances** Article consistents Article is an invitate on the interact of properties are consistent and interact the consistent are consistent at the interact of the common and interact the common and interact the comm

2.4 Hydrothermal carbonization

Hydrothermal carbonization (HTC), recognized as hydrous pyrolysis, stands out as cost-effective and sustainable thermochemical conversion technology. It operates through exothermic reactions, primarily involving dehydration and decarboxylation reactions aimed at reducing the oxygen (O) and hydrogen (H) content in biomass. In a sealed reactor system using water as the reaction solvent, biomass undergoes a sequence of chemical transformations, including hydrolysis, dehydration, decarboxylation, polymerization, and aromatization. HTC is executed in the temperature range of 180 °C to 260 °C, under pressures of approximately 2–6 MPa, and for durations spanning 5–240 min.⁴⁰ This process encompasses two main techniques, *i.e.*, hydrothermal vaporization and liquefaction, which occur at temperatures exceeding 260 °C, leading to the substitution of gaseous and liquid fuel generation for solid fuel.⁴¹ A distinctive advantage of HTC lies in its ability to process high moisture content feedstock and wet biomasses, eliminating the need for pre-drying as required in traditional pyrolysis methods. This not only streamlines the process but also significantly reduces the energy consumption, thereby enhancing the overall efficiency.⁴² The resulting products from HTC include hydrochar (solid carbon matter), bio-oil (liquid), and small amounts of gases, with their properties and percentage distribution influenced by the process conditions.⁴³

Hydrocar has emerged as a crucial product due to its versatile applications, displaying properties such as hydrophobicity and low ash content, which improve the mass, energy density, and combustion performance compared to the original feedstock.⁴⁴ Alternatively, biochar offers advantages in terms of porosity and surface area, making it suitable for applications such as adsorption and soil conditioning. HTC involves hydrothermal carbonization at both high and low temperatures (>220 °C, HHTC and \leq 220 °C, LHTC, respectively). Although HHTC produces materials similar to coal under natural formation processes, LHTC, conducted under milder conditions, is more commonly employed in laboratories.⁴⁵ Notably, current LHTC technology has been employed for the production of useful carbon compounds in a variety of shapes, including honeycomb and fibrous structures.⁴⁶ HTC has significant utility in treating high-water-content biomass, similar to that seen in municipal sludge, resulting in considerable energy savings and positioning HTC as technology for green thermochemical conversion.

The primary byproducts of HTC treatment are gas, bio-oil, and hydrochar, which exhibit distinct characteristics. Hydrochar is comprised of matrix and coke microparticles, with the matrix formed through condensation, dehydration, and decarboxylation of unhydrolyzed biomass. In contrast, the coke microparticles originate from the aggregation of organic molecules and are rich in functional groups such as –OH, $C=O$, and –COOH, facilitating their reactivity with organic molecules in the liquid phase. Overall, HTC represents a promising avenue for biomass conversion, offering a range of valuable products with diverse applications and demonstrating potential for both laboratoryscale research and industrial implementation (Table 3).⁷¹

3. Key factors impacting the characteristics of carbon materials

Considering that pyrolysis stands out as the dominant method for biomass conversion, this section delves deeply into the diverse array of factors that influence the initiation and characteristics of carbon materials during pyrolysis. Besides the inherent qualities of biomass itself, significant emphasis is placed on manipulating the variables involved in the reaction. Parameters such as pyrolysis temperature, duration of residence, rate of heating, and ambient conditions all play pivotal roles in determining the outcomes of carbon materials. These variables not only govern the yield of char but also intricately mold its properties, encompassing aspects such as surface area and pH levels. Given the profound interplay between adsorption capacity and the attributes of sorbents, it is essential to meticulously scrutinize the full spectrum of factors that impact carbon materials.

3.1 Biomass-derived feedstocks for the production of carbon materials

Exploring the diverse array of biomass-derived feedstocks for the production of carbon materials reveals a promising avenue for sustainable resource utilization. The categorization of feedstock for AC production plays a pivotal role in evaluating the nutrient

content and determining appropriate conversion techniques. Carbon-rich biomass, derived from both plant and animal sources, stands as the primary feedstock for AC production, with the objective of maximizing the carbon content in solid form.⁷² Activated carbon encompasses carbon materials with a robust carbon framework and a highly developed internal pore structure and surface area. The properties of activated carbon are influenced by the structural polymers found in biomass, including cellulose, hemicelluloses, and lignin. A higher cellulose and hemicellulose content contributes to increased volatile vapor production, while lignin serves as the principal precursor for the synthesis of AC.⁷³ The characteristics of biomass, including mineral matter content, significantly impact the pyrolysis properties and resultant product yield.⁷⁴ Classification based on moisture content delineates between wet (above 30%) and dry (below 30%) biomass, aiding in the selection of suitable conversion methods.⁷⁵ Biomass can be further categorized as 'purpose-grown' and 'waste biomass'. Purpose-grown biomass, cultivated specifically for AC production, exhibits a low moisture content (5%) , high yield, and energy density, finding applications in the energy and biofuel sectors. Conversely, waste biomass, lacking economic value, serves as an alternative feedstock for char production, often undergoing hydrothermal carbonization due to its elevated moisture content.⁷⁶ Biomass sources encompass a wide range, including animal sources (such as pig dung, pigskin, wool, and fishbone), plant sources (such as straw, rapeseed, sawdust, and peanut shell), and municipal waste (such as sludge).⁷⁷ The main elements in biomass are carbon, hydrogen, and oxygen, with minor amounts of nitrogen and sulfur. The appropriate treatment of biomass enables its conversion into carbon materials with commendable adsorption capabilities. However, certain types of biomass, such as rice husk, which contains high levels of silicon, are commonly used in the production of silicon carbide (SiC).⁷⁸ Thus, the selection of suitable biomass constitutes the initial step in preparing highquality carbon-based adsorbents. This chapter primarily reviews the biomass thermochemical conversion technologies, including pyrolysis, gasification, and hydrothermal carbonization, together with the factors influencing the properties of carbon materials such as the feedstock, temperature, residence time, and heating. Review Wext (1982)

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3.2 Pressure

In the domain of carbon material science, the influence of pressure stands as a pivotal yet often underestimated determinant. Pressure serves as a dynamic force that profoundly impacts the synthesis, structure, and properties of carbon materials across various manufacturing processes. Whether in the realm of crafting carbon nanomaterials or producing activated carbons, the application of varying pressure levels plays a central role in shaping the final product. Under high-pressure conditions, structural transformations are induced, porosity is altered, and mechanical properties are enhanced, thereby offering avenues for precise material tailoring. Conversely, low-pressure environments may foster the formation of specific carbon allotropes or facilitate unique chemical transformations, adding to the

intricacy of material design. The significance of pressure is particularly notable in the production of biochar and hydrochar. The reactions occurring within the reaction chamber adhere to Le Chatelier's principle, highlighting the critical role of pressure modulation. Increasing the pressure within the chamber, either through the introduction of non-reactive gases or elevating reaction temperatures, leads to notable enhancements in the yield and quality of both hydrochar and biochar.⁴⁵ This improvement is attributed to the accelerated decomposition of biomass, resulting in a superior end-product. In processes such as HTC, surpassing the critical pressure threshold enhances both the hydrolysis process and biomass decomposition, underscoring the multifaceted impact of pressure.⁷⁹ Although the physical structure of hydrochar remains relatively unchanged under increasing pressure, finer and more uniformly distributed particles emerge, accentuating the role of pressure in morphology control. Similarly, in pyrolysis, an increase in pressure prolongs the residence time, facilitating the formation of secondary carbon and augmenting the char quality. Moreover, pressure exerts a tangible influence on the carbon content and energy density of biochar, with high-pressure pyrolysis yielding superior outcomes in these aspects.⁸⁰ Understanding the subtle interplay between pressure and carbon materials is paramount for optimizing their performance across an extensive array of applications, spanning from energy conversion and storage to environmental remediation and beyond. **PEC Advances** Articles on the significance of presume is entimated on the frequential on the control and proposed under products are controlled to the frequential of the control and the significance of the control and the

3.3 Temperature

Temperature stands as a pivotal determinant in shaping the characteristics of carbon materials, exerting a profound influence on their synthesis, structure, and intrinsic properties. Varied temperature regimes dictate the ultimate outcomes of carbon materials, exerting notable effects on parameters such as porosity, surface area, and chemical reactivity. In addition to feedstock considerations, the significance of operating conditions cannot be overstated. Among these factors, the synthesis temperature holds particular prominence. As the pyrolysis temperature increases, discernible trends emerge, where the specific surface area and pH levels tend to increase, while the char yield experiences a decline. For instance, investigations by Singh et al. illustrated a notable reduction in biochar yield together with an increase in pH and surface area across the temperature range of 250 °C to 650 °C during the pyrolysis of rice husk.¹¹ However, excessive temperatures can lead to detrimental effects, such as the destruction of surface and pore structures due to melting and collapse. This phenomenon can be ascribed to the high temperature-induced disintegration and reorganization of the biomass components. Moreover, pH levels exhibit a positive correlation with temperature due to the enrichment of alkaline minerals such as sodium and potassium under hightemperature pyrolysis conditions. Elevated pyrolysis temperatures also facilitate the expulsion of volatile substances, thereby contributing to the formation of more developed pore structures. The research by Lahijani et al. underscored the impact of pyrolysis temperature on the $CO₂$ capture efficiency, revealing that the biochar produced at higher temperatures exhibited

enhanced $CO₂$ adsorption capabilities.⁸¹ This enhancement can be attributed to the fragmentation of the carbon particles, reduction in average pore size, and formation of micropores induced by increasing the carbonization temperatures. The presence of a microporous structure signicantly augments the adsorption of $CO₂$, underscoring the critical role of temperature in tailoring carbon materials for diverse applications, including environmental remediation and carbon capture technologies.

3.4 Residence time

Residence time, which denotes the duration in which the feedstock undergoes thermal decomposition in a reactor at a predefined temperature, has emerged as a critical parameter influencing the characteristics of carbon materials. Particularly in pyrolysis processes, the residence time exerts a substantial influence on both the yield and physicochemical attributes of biochar, primarily by fostering pore formation and structural refinement.⁸² Prolonged reaction durations often yield positive outcomes, enhancing the energy density and facilitating the production of secondary hydrochar, distinguished by its polyaromatic structure. Conversely, shorter residence times tend to yield a higher hydrochar content, a phenomenon notably observed in HTC processes.⁸³ Manipulation of the residence time induces the release of volatile compounds, consequently affecting the composition of the resulting materials, with a lower soil product content observed under increased residence times.⁸⁴ Additionally, the residence time significantly impacts the texture of hydrochar, modulating processes such as polymerization and hydrolysis, and influencing the micropore diameter. Co-pyrolysis experiments involving agricultural residues and sewage sludge further elucidated the role played by the residence time, demonstrating a pronounced decrease in biochar yield with extended durations, suggesting the completion of organic compound degradation within 150 min.⁸⁵ Correspondingly, surface area exhibits a notable increase with longer residence times, which is attributed to the enhanced thermal decomposition and subsequent porous structure development, potentially augmenting the $CO₂$ adsorption capacity.⁸⁶ However, a further increase beyond 150 min resulted in a decline in surface area due to structural collapse. These insights underscore the intricate relationship between residence time and properties of carbon materials, emphasizing the need for meticulous control and optimization to tailor the characteristics of materials for diverse applications, including carbon sequestration and environmental remediation.

3.5 Heating rate

The rate at which heating occurs plays a crucial role in shaping the characteristics of carbon materials, influencing parameters such as carbonization efficiency, surface morphology, and chemical composition. Typically, accelerated heating rates result in higher carbonization temperatures, fostering rapid pyrolysis and potentially inducing structural alterations, whereas slower rates afford a more controlled decomposition process, yielding materials with distinctive surface functionalities. This aspect of the heating rate significantly impacts the

pyrolysis carbonization mechanism and the characteristics of the resultant biochar. Notably, an increase in heating rate is often correlated with enhanced biomass weight loss. For instance, the research conducted by Li et al. scrutinized the pyrolysis of lignin at varying heating rates, revealing a decrease in biochar yield from 55.5% to 50.1% as the heating rate increased from 5 to 20 $^{\circ}$ C min⁻¹, together with a concomitant rise in gas yield from 18.2% to 22.2%.⁸⁷ Moreover, increased heating rates facilitate the cracking of organic constituents and volatiles within biochar, expediting the generation of gas. Despite the prevailing assumptions that higher heating rates uniformly augment the specific surface area of biochar, contradictory findings have been documented. Hu et al. observed a decrease in specific surface area from 362 to 146 $m²$ $\rm g^{-1}$ with an increase in the heating rate from 2 to 15 °C min $^{-1}$, followed by an increase to 327 m^2 g^{-1} at 25 °C min^{−1}.⁸⁸ This phenomenon is attributed to the intense decomposition and volatile diffusion occurring at steep heating rates, underscoring the interplay between the heating rate and biochar characteristics.

4. Activated carbon

Activated carbon, commonly known as AC, encompasses carbonaceous materials featuring a sturdy carbon framework together with a fully developed surface area and interior pore structure. Various organic materials possess the potential for conversion into activated carbon, with petroleum coke, peat, and wood being the common precursors utilized in its development.⁸⁹ Therefore, comprehending the composition and structure of the precursor material is imperative in determining the most suitable activating agent for AC production. The constituent structural polymers within biomass, including cellulose, hemicelluloses, and lignin, significantly influence the characteristics of the resulting AC. Typically, biomass is comprised of approximately 40% cellulose, 30% hemicellulose, and 30% lignin, with lignin requiring higher temperatures for complete decomposition compared to cellulose.⁹⁰

Consequently, materials rich in lignin content tend to yield AC with a lower surface area at lower activation temperatures.⁹¹ Additionally, non-structural components such as proteins, ash, waxes, pectin, tannins, and chlorophylls contribute to the raw material composition.⁹² Micropores, among the various pore structures present, play a pivotal role in adsorption due to their small void space, facilitating the trapping of organic molecules. The micropores formed during the activation of carbon are crucial for the adsorption of carbon dioxide, given that they allow localized van der Waals forces to overlap, promoting strong electrostatic attraction and physisorption.⁹³ A pore size of less than 1.0 nm is deemed effective for $CO₂$ capture due to the small molecular size of $CO₂$.⁹⁴ The adsorption process exhibits low activation energy, enabling reversible adsorption and the potential for multiple-layer $CO₂$ adsorption on AC, thereby enhancing the adsorption capacity. Thus, optimizing the synthesis process is paramount for producing AC with a high surface area, which is conducive to efficient $CO₂$ adsorption. However, despite its stability and superior surface area, activated carbon faces challenges regarding economic and environmental viability.⁹⁵ Recent advancements highlight the equivalent $CO₂$ adsorption capabilities of activated carbon derived from various waste material precursors compared to their commercial counterparts, offering promising avenues for the sustainable production of AC.⁹⁶ Review Were

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4.1 Carbon material modification techniques

In the field of carbon materials, investigating techniques for their modification is a crucial area of study driving research and innovation forward. This review offers a comprehensive examination of the various methodologies intended to customize the properties and functions of carbon materials, elucidating a wide array of modification techniques and their applications across different fields. Carbon materials derived from biomass in their original state exhibit inherent limitations in chemical composition and pore structure, restricting their ability to adsorb $CO₂$ effectively. Therefore, modifying carbon materials becomes exceedingly significant, particularly in the realm of $CO₂$ capture.

Fig. 7 Schematic illustrating the two primary approaches for modifying carbon materials: (a) pre-treatment and (b) post-treatment methods.

The two primary approaches for modifying carbon materials are outlined, as illustrated in Fig. 7, i.e., pre-treatment, which involves directly mixing raw biomass with acids or metal salts before carbonization, and post-treatment, modifying carbonized biomass through activation with substances such as acids, bases, metal salts, and gases. Notably, it is imperative to treat raw biomass with diluted acid prior to its modification to remove its ash content, thus averting pore blockage during the carbonization process. This review thoroughly explores the modification methods, which are classified into three main categories of physical methods (steam/air/ $CO₂$ activation), chemical processes (acid/alkaline activation, metal or metal oxide impregnation, and heteroatom doping), and supplementary techniques (ball milling, microwave, ultrasound, and plasma treatment). Through a detailed analysis of these techniques, this review seeks to offer valuable insights into the latest advancements and future trajectories in the modification of carbon materials, aiming to enhance their performance and widen their range of applications.

4.1.1 Physical treatment. Physical activation presents an eco-friendly, cost-efficient, and straightforward method compared to chemical activation approaches. By employing gases as templates, precise control of the textural characteristics of activated carbon is achievable. During pyrolysis in the temperature range of 300–1100 °C, diverse degradation byproducts including CO_2 , CO , CH_4 , NO, H_2O , and NH_3 are produced, depending on the source of the precursor material. Moreover, external medium sources such as $CO₂$, NH₃, steam, He, Ar, air, and gas mixtures can be utilized to activate the surface of carbon. Extensive endeavors have been devoted to enhancing the pore volume and surface area of carbon materials through physical activation methodologies.

4.1.1.1 Steam activating methods. Steam activation is a significant technique for improving the textural characteristics of carbon materials by facilitating the creation of additional pores.⁹⁷ This process involves the diffusion of steam into the carbon material, which promotes the formation of pores. The underlying mechanisms of steam activation encompass various reactions, including the interaction between carbon and water vapor to generate carbon monoxide and hydrogen, together with the production of methane and other gases.

$$
C + H_2O \rightarrow C(O) + H_2 \tag{1}
$$

$$
C(O) \to CO + C \tag{2}
$$

$$
CO + H_2O \rightarrow CO_2 + H_2 \tag{3}
$$

$$
C + 2H_2O \rightarrow CO_2 + 2H_2 \tag{4}
$$

$$
C + CO_2 \rightarrow 2CO \tag{5}
$$

$$
C + 2H_2 \rightarrow CH_4 \tag{6}
$$

$$
CH_4 + H_2O \rightarrow CO + 3H_2 \tag{7}
$$

Observations demonstrate a notable enhancement in specific surface area and pore volume of biochar following its

steam activation. For instance, Xie et al. employed steam activation to modify biochar derived from tea waste, resulting in a significant increase in specific surface area from 342 to 576 $m²$ g^{-1} and enhancement in pore volume from 0.022 to 0.109 cm³ $\rm g^{-1.}$ ⁹⁸ The enhancement in pore structure resulting from steam activation can be attributed to the elimination of pore-clogging tar, the creation of new micropores, and the enlargement of existing pores. Different steam activation conditions, such as temperature and reaction time, have varying effects on the activation results. Other studies using cellulose fibers as a raw material revealed that steam activation facilitated the formation of new pores and influenced the expansion of the pore sizes, leading to a remarkable $CO₂$ adsorption capacity of 3.78 mmol $\rm g^{-1.99}$ Furthermore, the steam activation of carbon derived from cellulose fibers resulted in the expansion of existing ultramicropores and the generation of additional ultra-micropores (Fig. 8).¹⁰⁰ This activation process disrupted the graphitic structure, resulting in pore sizes exceeding 2 nm and a significant increases in specific surface area and pore volume. Additionally, Chen et al. demonstrated that steam activation of biochar produced from pine sawdust not only enhanced its textural properties but also enriched its surface functional groups.¹⁰¹ The specific surface area increased from 316 to 582 m^2 g⁻¹, and the pore volume increased from 0.17 to 0.25 cm^3 $\rm g^{-1}.$ XPS analysis indicated an increase in the oxygen content of biochar from 9.16% to 11.05% following steam activation, which was attributed to the increase in the content of C–O, C– OH, and O–C $=$ O functional groups. These findings underscore the efficacy of steam activation in improving the textural properties and surface chemistry of carbon materials, thereby expanding their potential applications across various fields. **PSC** Advances Articles Articles

4.1.1.2 Air activating methods. Air activation has emerged as a notable strategy for altering carbon materials, offering an environmentally friendly alternative compared to chemical activation techniques. This method involves utilizing air as the activating agent, eliminating the need for additional chemicals. Oxygen, known for its potent oxidizing properties, plays a crucial role in the activation process. However, due to the prohibitive cost of pure oxygen, air is commonly utilized for activation purposes. The underlying mechanism for air activation can be understood through various reactions,¹⁰² as follows:

$$
2C + O_2(g) \rightarrow 2CO \tag{8}
$$

$$
C + O_2(g) \rightarrow CO_2 \tag{9}
$$

$$
C(O) + O_2(g) \rightarrow CO(g) + CO_2 \tag{10}
$$

Air oxidation has the potential to introduce new functional groups into biochar. However, this oxidation process may lead to a slight decrease in surface area due to the new functional groups obstructing the pores. Kim et al. observed that air oxidation enriched the micropores of biochar and introduced O-groups, albeit resulting in reduced specific surface area.¹⁰³ Furthermore, carbon materials produced at lower temperatures are more susceptible to oxidation, leading to the introduction of additional –COOH and $C=O$ groups. In comparison with steam

Fig. 8 Diagram illustrating the process of activating the surface of carbon with steam.

activation, air oxidation tends to introduce O-containing groups more readily. Another research group reported differing outcomes between steam activation and air oxidation, with steam activation resulting in an increased surface area but decreased functional group concentration, while air oxidation led to a higher functional group concentration in biochar.¹⁰⁴ Lee et al. conducted air activation on biochar at various temperatures and noted a significant enhancement in surface groups such as -OH, $C=O$, and $C-O$.¹⁰⁵ This enhancement is attributed to the reactions between O_2 and the lignocellulosic components or aromatic substances within the biochar. Additionally, they examined the impact of activation time on the process, observing an initial increase in the micropores within the first 15 min, followed by an increase in the content of mesopores. However, a prolonged activation time could potentially lead to micropore destruction, highlighting the importance of selecting an appropriate activation duration. For carbon materials intended for $CO₂$ capture applications, determining the optimal oxygen activation treatment time is essential to maximize their efficacy.

4.1.1.3 $CO₂$ activating methods. $CO₂$ activation has emerged as a signicant technique for modifying carbon materials, providing an eco-friendly avenue to enhance their properties. This method involves employing $CO₂$ as the activating agent, leading to tailored advancements in the pore structures and surface functionalities within carbon materials. Its adoption is gaining recognition due to its environmentally benign nature and the abundance of $CO₂$ as a readily accessible resource. $CO₂$ activation can be carried out either simultaneously with biomass pyrolysis or as post-pyrolysis treatment. Studies suggest that pyrolysis conducted in a $CO₂$ atmosphere is advantageous for preserving C and N in biomass. Moreover, this activation approach significantly improves the porosity and specific surface area of carbon materials. For instance, Tomin et al. illustrated

that CO₂ activation predominantly yields micropores in porous carbon fibers, resulting in a specific surface area of up to 1012 m^2 $\rm g^{-1.106}$ However, the development of mesoporous structures may be observed in certain carbon materials following $CO₂$ activation, which is influenced by factors such as the type of pristine materials and activation conditions. Activation of various biochars with $CO₂$, including wheat straw, softwood, and peach stones, resulted in a more pronounced development of mesoporous structure in wheat straw and softwood biochar, while peach stone biochar exhibited some enhancement in microporosity.¹⁰⁷ CO₂ activation involves the Boudouard reaction, an endothermic process, and thus higher temperatures favor $CO₂$ activation. For instance, $CO₂$ activation of biochar conducted at temperatures in the range of 800 °C to 950 °C showed an escalation in pore improvement with an increase in the activation temperature.¹⁰⁸ Specifically, the specific surface area of biochar increased from 392 m² g⁻¹ to 506, 854, 1059, and 2186 m² g⁻¹, respectively, with micropores less than 2 nm dominating the pore structures for all the samples. These findings underscore the efficacy of $CO₂$ activation in enhancing the porosity and specific surface area of carbon materials, offering promising avenues for their utilization in various applications.

4.1.2 Chemical treatment

4.1.2.1 Alkaline activating methods. Alkaline activation is a crucial method for modifying carbon materials, offering a flexible approach to customize their properties. This process involves the use of alkaline agents such as KOH and NaOH to activate the carbon precursors, leading to the formation of tailored pore structures and surface functionalities. This method is particularly advantageous for enhancing the $CO₂$ adsorption capacity by promoting pore development and incorporating alkaline groups. The common alkaline activators include KOH, NaOH, and amino $(-NH₂)$ groups. Among them, KOH is widely recognized for its effectiveness as a metal hydroxide in activating carbon materials. Typically, the activation process begins by thoroughly mixing KOH with the carbon precursor. The redox reaction between KOH and the carbon precursor starts at around 400 °C, producing K_2CO_3 . As the temperature increases to about 600 °C, KOH is fully converted into K₂CO₃. At temperatures above 700 °C, K₂CO₃ begins to decompose into $CO₂$ and potassium oxide (K₂O). Additionally, at even higher temperatures, both K_2CO_3 and K_2O can be reduced to potassium (K) , which infiltrates the carbon lattice in vapor form, assisting the formation of micropores and enhancing the porous structure.

The reaction sequence for KOH activation is outlined as $follows: ¹⁰⁸$

$$
2C + 6KOH \to 2K + 2K_2CO_3 + 3H_2
$$
 (11)

$$
K_2CO_3 \rightarrow K_2O + CO_2 \tag{12}
$$

$$
C + CO_2 \rightarrow 2CO \tag{13}
$$

$$
2C + K_2CO_3 \rightarrow 2K + 3CO \tag{14}
$$

 $C+K_2O \rightarrow 2K + CO$ (15)

KOH activation enhances the surface functionality of carbon materials by introducing various groups such as $-OH$, $C=O$, $C C$, and $=$ C–H. However, as the synthesis temperature increases, the O-containing groups on the surface of the carbon material decompose, resulting in the production of CO and CO₂.¹⁰⁹ Due to the acidic nature of carbon dioxide, incorporating Lewis bases on the surface of carbon materials has been proven to be advantageous for $CO₂$ capture. An investigation into biochar derived from a mixture of food and wood waste demonstrated that although KOH activation led to superior microporous structures and enhanced $CO₂$ adsorption capacity, activation with KOH–CO₂ did not yield further improvements.¹¹⁰ Researchers explored the impact of KOH activation on the production of porous carbon from biomass for supercapacitors, noting the significant influence of KOH on the microstructure of the resulting char. The optimized carbon materials exhibited a specific surface area of up to 2790 $\mathrm{m^2\,g^{-1}}$, with a graded pore structure enhancing the rate performance and achieving a maximum specific capacitance of 327 F $\rm g^{-1}.$

Alternatively, NaOH activation offers stronger oxidizing properties and is more environmentally friendly. Studies indicate that the redox reaction between NaOH and carbon contributes to forming both micropores and mesopores in carbon materials. During activation, NaOH interacts with carbon materials through reactions as follows:¹¹¹

$$
2C + 6NaOH \rightarrow 2Na + 2Na2CO3 + 3H2
$$
 (16)

$$
2C + Na2CO3 \rightarrow 2Na + 3CO
$$
 (17)

$$
2Na + CO2 \rightarrow Na2O + CO
$$
 (18)

Sodium hydroxide (NaOH) activation of carbon generates active intermediates such as CO , $CO₂$, and $H₂$, simultaneously

facilitating the infiltration of Na and $Na₂CO₃$ into the carbon layer, which expands its atomic structure and enhances the specific surface area and pore diameter of the resulting carbon materials. However, excessive NaOH concentrations can intensify gasification reactions, ultimately reducing the effective area. Elevated NaOH concentrations may also induce a series of reactions (e.g., C-NaOH, C-Na₂CO₃, C-Na₂O, C-Na, C-CO₂, and C–CO), leading to the rupture of the C–C and C–O–C bonds and diminishing the yield of carbon materials.¹¹² Studies by Shin et al. demonstrated the synthesis of carbon materials from macroalgae using NaOH activation, yielding materials with a significant surface area of 1238 $\mathrm{m^2\,g^{-1}}$. Another investigation highlighted the enhancement of aromatic properties in biochar through NaOH activation, promoting $\pi-\pi$ interactions during pollutant adsorption.¹¹³

Incorporating amino groups into carbon materials also significantly enhances their $CO₂$ capture capacity. NH₃ treatment, typically performed at high temperatures, introduces basic functional groups and removes acidic groups from the surface of carbon. This treatment results in the formation of amides, lactams, imides, and heterocyclic compounds such as pyrrole and pyridine.¹¹⁴ Additionally, $NH₃$ activation moderately improves the pore size and specific surface area. Studies by Wu et al. demonstrated an increase in the specific surface area of biochar from coconut fiber from 4, 7, and 541 m² g⁻¹ to 9, 24, and 553 m² g⁻¹, respectively, after NH₃⋅H₂O modification.¹¹⁵ Liu et al. synthesized microporous N-doped carbon materials using $NH₃$ as the activator, highlighting the synergistic effects of pores and N-groups in determining high $CO₂$ capture capacity and selectivity over N_2 ¹¹⁶ **PSC** Advances Articles. Typically, the active-

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4.1.2.2 Acid activating methods. Acid activation serves as a crucial technique for tailoring the characteristics of carbon materials, significantly enhancing their surface area, pore structure, and functional groups, thereby improving their adsorption capabilities. Typically, this process involves treating carbonaceous materials, such as activated carbon, biochar, and carbon fibers, with strong acids such as H_3PO_4 , H_2SO_4 , and $HNO₃$. The acid activation process not only creates new pores but also widens the existing pores, leading to a well-developed pore structure, which is essential for various adsorption applications.

 H_3PO_4 has emerged as the predominant acid activator in the production of activated carbon for carbon capture applications, undergoing transformation into polyphosphate during the activation process.¹¹⁷

$$
2P_2O_5 + 5C \rightarrow P_4 + 5CO_2 \tag{19}
$$

$$
2H_2P_2O_7^{2-} \to P_4 + 6O_2 + 2H_2O
$$
 (20)

Activation with H_3PO_4 not only stimulates the development of pores but also introduces novel functional groups into the structure of carbon. For instance, the immersion of Lantana camara biomass in 85 wt% H_3PO_4 solution led to a significant increase in the specific surface area of the activated biochar from 117 to 1177 m^2 g⁻¹ and a 20-fold increase in pore volume compared to the untreated biochar.¹¹⁸ Furthermore, the oxygen

content of the biochar increased, with the XPS analysis con firming the presence of surface groups such as $C-PO₃$, $C-P₂O₆$, and $C-P_2O_5$. Similarly, Bonga *et al.* observed surface groups such as $C-O-PO₃$ and $C-PO₃$ in biochar produced through phosphoric acid activation in the temperature range of 400– 600 °C.¹¹⁹ However, it is notable that the acidic P_xO_y groups resulting from H_3PO_4 activation have been found to hinder CO_2 chemisorption in carbon materials. In this case, fine-tuning of the surface chemistry and porous structures of carbon materials can be achieved by adjusting the concentration of H_3PO_4 and the treatment temperature.¹²⁰ Although a porous structure plays a crucial role in $CO₂$ capture at lower temperatures, surface chemistry, including the presence of functional groups, becomes increasingly influential at higher temperatures during adsorption. Thus, meticulous attention is warranted in both the preparation and application of H_3PO_4 -activated carbon. Review Worker on the binden increased, with the XPs are
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Sulfuric acid is an extremely reactive substance, which can interact with organic compounds, such as carbohydrates, to extract water and break down organic precursors into carbon, according to the following reaction:¹²¹

$$
C_nH_{2n}O_n + H_2SO_4 \to nC + H_2SO_4 \cdot nH_2O \qquad (21)
$$

Additionally, sulfuric acid reacts with the mineral content in lignocellulosic materials, serving as an effective cleaning and de-ashing agent in the preparation of activated carbon. This makes H_2SO_4 a cost-effective option for carbonization processes, offering both economic and functional advantages.¹²¹ Furthermore, the activation process using sulfuric acid is known to enhance the porous structure of the resulting activated carbon. Notably, the porous carbon treated with $H₂SO₄$ exhibited a commendable $CO₂$ adsorption capacity of 3.60 mmol \rm{g}^{-1} , marking a 39.5% improvement compared to the untreated sample.¹²² The acid penetrates the material, creating a surface with a combination of large and medium-sized pores, which is beneficial for a variety of applications. $121,123$

Additionally, nitric acid activation is primarily used to introduce a variety of oxygen-containing functional groups, such as carboxyl, hydroxyl, and carbonyl groups, on the surface of carbon. This treatment is usually conducted at room temperature to 100 °C. Nitric acid oxidizes the surface of carbon, creating defects and functional groups, which enhance the adsorption capacity for polar molecules, including heavy metals and organic pollutants. The reaction mechanism is as follows:¹²⁴

$$
HNO3 \rightarrow NO2 + O2 + H2O
$$
 (22)

$$
2NO2 + C \rightarrow C(O) + 2NO
$$
 (23)

$$
C(O) + H_2O \rightarrow COOH \tag{24}
$$

$$
C + HNO3 \rightarrow C-OH + 2NO2
$$
 (25)

$$
C + O_2 \rightarrow C = O \tag{26}
$$

The oxidation process increases the surface acidity and introduces active sites, which improve the interaction of carbon

materials with the adsorbate, enhancing their performance in various applications.

4.1.2.3 Salt activating method. Salt activation is a crucial method for tailoring the properties of carbon materials, offering a versatile approach to enhance their surface area, pore structure, and adsorption capabilities. This technique employs various salts, such as potassium carbonate (K_2CO_3) , sodium carbonate (Na₂CO₃), zinc chloride (ZnCl₂), and ferric chloride $(FeCl₃)$, to activate the carbonaceous precursors through thermal treatment. Unlike traditional alkaline activators such as KOH and NaOH, which are strong bases, salts such as K_2CO_3 and $Na₂CO₃$ also contribute to activation by providing different chemical environments. These salts activate the carbonaceous precursors through thermal treatment, promoting the development of porous structures *via* specific reaction mechanisms.¹²⁵

 K_2CO_3 is a common activating agent, which upon heating typically between 700 °C and 900 °C, decomposes to release $CO₂$ and K_2O . The CO_2 generated in this reaction acts as a gasifying agent, creating pores within the carbon matrix. Additionally, $K₂O$ can further react with the carbon structure to form metallic potassium, which intercalates into the carbon lattice, enhancing its porosity, as follows:

$$
K_2CO_3 \rightarrow K_2O + CO_2 \tag{27}
$$

$$
K_2O + C \rightarrow 2K + CO \tag{28}
$$

This process leads to a significant increase in surface area and pore volume, making K_2CO_3 -activated carbons highly effective for adsorption applications.¹²⁶

 $Na₂CO₃$ follows a similar activation pathway to $K₂CO₃$. During thermal treatment, $Na₂CO₃$ decomposes into sodium oxide (Na₂O) and (CO₂). CO₂ aids in the development of a porous structure by gasifying the carbon material. $Na₂O$ further reacts with carbon to produce elemental sodium, which intercalates into the carbon structure, enhancing the formation of pores, as follows:¹²⁷

$$
Na_2CO_3 \rightarrow Na_2O + CO_2 \tag{29}
$$

$$
Na2O + C \rightarrow 2Na + CO
$$
 (30)

Carbon materials activated by Na_2CO_3 exhibit a high surface area and improved adsorption properties, making them suitable for various environmental applications, including pollutant capture and energy storage.

 $ZnCl₂$ is widely recognized as an effective activation agent, particularly due to its strong dehydrating properties. The activation process with $ZnCl₂$ typically involves three stages during pyrolysis. Initially, biomass decomposes into small particles and volatiles as the temperature increases. In the subsequent stage, $ZnCl₂$ interacts with these particles, producing char, tar, and volatiles through the following reaction:¹²⁸

$$
ZnCl_2 + C \rightarrow Zn + Cl_2 + CO \tag{31}
$$

In the final stage, the char undergoes further reaction with ZnCl₂, leading to significant pore development. Carbon

materials activated with $ZnCl₂$ are noted for their increased prevalence of mesopores and macropores, which is attributed to the deep penetration of molten $ZnCl₂$ into the carbon lattice, which expands the pore structure. Co-activation with $CO₂$ further enhances the specific surface area, total pore volume, and even generates ultra-fine pores.^{129,130} Conversely, activation with ferric chloride $(FeCl₃)$ is more effective in promoting micropore formation, as demonstrated by Elewa et al., who achieved a micropore volume of 0.468 $\rm cm^3~g^{-1}$ using FeCl₃.¹³¹ However, the environmental implications of $ZnCl₂$ activation warrant attention. During this process, some $ZnCl₂$ can be converted to ZnO, which may adsorb on the activated carbon, posing the risk of soil and groundwater contamination. Considering these concerns, the EU proposed restrictions on high concentrations of ZnO since 2017, with similar measures initiated by China in 2018.¹³²

 $FeCl₃$ is another effective activating agent, particularly favoring micropore formation. In the activation process, $FeCl₃$ acts as a catalyst, promoting the removal of volatile compounds and facilitating the development of a highly porous structure, as follows:

$$
2\text{FeCl}_3 + 3\text{C} \rightarrow 2\text{Fe} + 3\text{Cl}_2 + 3\text{CO} \tag{32}
$$

This reaction results in carbon materials with a high micropore volume, which is beneficial for applications such as gas storage and separation.¹³³

4.1.2.4 Metal doping. The impregnation of biomass-derived carbon materials with metals such as sodium, calcium, magnesium, aluminum, nickel, and iron has attracted significant attention as a strategy to enhance their $CO₂$ capture capabilities. Biomass-derived carbon materials, including biochar and activated carbon, are attractive due to their sustainable and eco-friendly nature, offering a renewable alternative by recycling agricultural and forestry waste. These materials naturally possess high surface areas and porous structures, making them highly effective for gas adsorption. Furthermore, the introduction of metal species into these carbon matrices can enhance their $CO₂$ adsorption capacities and selectivity, making them more efficient and versatile for carbon capture applications.¹³⁴

Sodium and calcium, both alkaline metals, have been shown to significantly enhance $CO₂$ capture through chemisorption mechanisms. When impregnated into biomass-derived carbon materials, these metals form stable carbonates, such as Na_2CO_3 and CaCO₃, respectively, which effectively bind $CO₂$ molecules. This chemisorptive process not only increases the efficiency of $CO₂$ capture but also reduces the likelihood of desorption, making these materials more reliable for long-term $CO₂$ storage.¹³⁵ The stability of these carbonates ensures that the captured $CO₂$ remains securely bound, reducing the risk of its release back into the atmosphere. Magnesium, another alkaline earth metal, also enhances the capture of $CO₂$ when incorporated into biomass-derived carbon materials. The formation of $MgCO₃$ within the carbon matrix, combined with the high surface area of the material, creates an effective platform for $CO₂$ adsorption. The presence of MgCO₃ not only increases the

affinity of the material for $CO₂$ but also contributes to the overall stability of the adsorbed $CO₂$.¹³⁶ Aluminum impregnation introduces Lewis acid sites on the surface of carbon through the formation of alumina $(Al₂O₃)$. These acid sites interact strongly with $CO₂$ molecules, significantly enhancing both the adsorption capacity and selectivity of the material.¹³ The strong interaction between $CO₂$ and the aluminum-impregnated carbon material increases the efficiency of $CO₂$ capture, making it particularly suitable for selective $CO₂$ adsorption in mixed-gas streams.

Transition metals such as iron and nickel play crucial roles in improving $CO₂$ capture. Iron, forms iron oxides, such as $Fe₂O₃$ and $Fe₃O₄$, on the surface of carbon. These oxides facilitate both physisorption and chemisorption mechanisms, significantly enhancing the overall $CO₂$ capture efficiency.¹³⁷ Alternatively, Ni doping is a promising strategy to enhance their properties for various applications, particularly in $CO₂$ adsorption and electrochemical energy storage. Nickel doping involves impregnating carbon materials derived from biomass with a nickel precursor such as nickel nitrate $(Ni(NO₃)₂)$ or nickel chloride ($NiCl₂$). This process begins with the pyrolysis of biomass to produce a porous carbon framework. Subsequently, this carbon material is impregnated with a nickel salt solution, allowing nickel ions to diffuse and become uniformly distributed within the carbon matrix. After impregnation, the material undergoes thermal activation, typically at temperatures in the range of 300 °C to 900 °C in an inert atmosphere. This thermal treatment reduces the nickel salts to metallic nickel, which is embedded within the carbon structure, forming nickel oxides and carbonates, serving as active sites for $CO₂$ interaction.¹³⁸ PSC Advances are entired on 18 202 are noted for their increased arthitly of the national CO₂, Numation incredicts can be a created on 18 202 and the stational interest are the stational interest are the stational incre

The incorporation of nickel into biomass-derived carbons significantly enhances their $CO₂$ adsorption capacity and selectivity. Nickel carbonates and oxides provide additional active sites, which promote stronger interactions with $CO₂$ molecules, thereby increasing the amount of $CO₂$ that can be adsorbed. This is particularly advantageous for applications requiring selective adsorption, such as carbon capture and storage.¹⁰² The presence of nickel also improves the selectivity of materials for $CO₂$ over other gases, making it more efficient in separating $CO₂$ from gas mixtures. This enhanced selectivity is due to the specific interactions between $CO₂$ molecules and the nickel-based active sites, which are less favorable for other gases. In addition to gas adsorption, nickel doping signicantly improves the electrochemical properties of biomass-derived carbons. In applications such as fuel cells, batteries, and supercapacitors, nickel acts as a catalyst, which enhances the electrocatalytic activity and stability of the carbon material. Also, the presence of nickel increases the electrical conductivity and facilitates efficient charge transfer, which are crucial for high-performance energy storage and conversion devices.¹³⁹

4.1.2.5 Heteroatom doping. Heteroatom doping provides a promising pathway for tailoring the characteristics of carbon materials by deliberately introducing elements such as nitrogen, sulfur, and boron into the carbon framework. This approach alters the electronic structure and surface chemistry of carbon, thereby improving its conductivity, catalytic activity, and ability to adsorb various substances. Nitrogen doping has

attracted considerable interest due to its capacity to create active sites and enhance the electrochemical performance of carbon-based materials. Similarly, sulfur and boron doping have been explored for their potential to augment the adsorption capacity and stability of carbon materials across diverse applications. Although micropores play a pivotal role in carbon dioxide capture, surface functional groups also signicantly contribute to the adsorption process, accounting for about 38%. Surface alkalinity plays a crucial role in $CO₂$ adsorption, and heteroatom doping has been demonstrated to enhance the surface alkalinity, thereby improving the $CO₂$ uptake.¹⁴⁰ Nitrogen doping, being the most extensively researched method, not only regulates the pore structure of carbon materials but also enriches them with various forms of N-containing groups such as pyridone, cyanide, pyrrole, pyridine, amine, quaternary, and pyridine-N-oxide. Compounds such as PEI, TEPA, and DETA have been shown to be effective as nitrogen dopants for $CO₂$ capture.¹²⁹ However, the additional cost associated with nitrogen doping underscores the importance of identifying nitrogen-rich biomass sources for the synthesis of carbon materials. However, although biomass with a high nitrogen content tends to yield nitrogenous biochar, not all nitrogen in biomass can be converted to N in carbon materials, given that some may be transformed into $NH₃$ or other gases during the process. Beyond single nitrogen doping, other elements such as sulfur and co-doping have been proven to be effective. For instance, researchers synthesized sulfur-doped porous carbon with a $CO₂$ adsorption capacity of 4.5 mmol $\rm g^{-1}$, surpassing similar materials doped with nitrogen, which is possibly due to the higher binding energy of $CO₂$ –S compared to $CO₂–N$, resulting in more stable adsorption and higher capacity.¹⁴¹ Guo et al. utilized poplar sawdust as a raw material and sulfur-containing wastewater as a modifier to co-load nitrogen, sulfur, and oxygen on activated carbon, yielding porous carbon with a CO₂ adsorption capacity of 5.61 mmol g^{-1} at 0 °C.¹⁴² Open Access Article. Published on 18 2024. Downloaded on 15-11-2024 20:10:22. This article is licensed under a [Creative Commons Attribution-NonCommercial 3.0 Unported Licence.](http://creativecommons.org/licenses/by-nc/3.0/) **[View Article Online](https://doi.org/10.1039/d4ra04537h)**

4.1.2.6 Other modification methods. Besides the primary modification methods discussed previously, supplementary approaches such as ball milling, microwave irradiation, ultrasound treatment, and plasma modification show potential for further improving carbon materials. Ball-milling modification utilizes the kinetic energy produced during the movement of balls in machinery to diminish the particle size of materials or disrupt their chemical bonds, thereby modifying their pore structures. An examination of crayfish shell biochar pre- and post-ball milling unveiled notable enhancements in its specific surface area and micropore volume, escalating from 128 to 290 $\mathrm{m^2\,g^{-1}}$ and 0.028 to 0.061 $\mathrm{cm^3\,g^{-1}}$, respectively. 143 Additionally, the comparison among pristine biochar, ball-milled biochar, and ball-milled-N-doped biochar demonstrated superior pore structures in the modified biochar, coupled with increased $CO₂$ adsorption capacities.¹⁴⁴ Microwave radiation, an emerging heating method, presents advantages such as swift heating, convenient regulation, and uniform heat dispersion. Given their elevated microwave absorption capacity, carbon materials can be transformed under microwave heating, generating new carbons with tailored characteristics.¹⁴⁵ The rapid heating facilitated by microwave radiation may prompt the formation of new pores and modify the pore structures and surface functional groups in carbon materials. Studies suggested an increase in the number of functional groups on the molecular bonds of graphene materials after microwave treatment, providing additional adsorption sites for the adsorption process.¹⁴⁶

Ultrasonic treatment has emerged as a novel method for altering carbon materials. By exploiting cavitation and microjet effects, ultrasonic treatment effectively eradicates the lumps formed during the pyrolysis of biomass, thus preventing pore blockage and facilitating the creation of additional microporous structures.¹³⁴ Typically, ultrasonic treatment is paired with chemical modification techniques to optimize its effectiveness. For example, in the study by Cao et al., biochar was synthesized through ultrasonic treatment followed by amination at room temperature. Preceding amination, ultrasonic treatment of biochar induced the detachment of the graphite layer, resulting in the formation of new pores.¹⁴⁷ The resultant exfoliated biochar exhibited an abundance of functional groups such as – $COOH$, -OH, and -CH (O) CH-, which facilitated amine grafting. As a result, the biochar treated with ultrasound and amines exhibited a nine-times greater $CO₂$ adsorption capacity than that of the untreated biochar. Plasma, consisting of cations, particles, and free electrons, serves as a neutral substance for modifying the surface characteristics of materials, which is typically generated through corona, glow, or microwave discharge methods. During the modification process, interactions between plasma and the material surface facilitate the formation of new groups. In the study by Khan et al., they utilized atmospheric pressure plasma to modify activated carbon, observing improvements in its surface area and pore volume together with the enrichment of surface oxygencontaining functional groups.¹⁴⁸

4.1.3 Impact of activation temperature. The impact of activation temperature on the properties and efficacy of activated carbon cannot be overstated. This comprehensive review explores the intricate interplay between activation temperature and the resultant characteristics of activated carbon. A multitude of factors, including pore volume, pore size distribution, microporosity development, and surface area, are signicantly influenced by the temperature regime employed during the carbonization and activation processes. Higher temperatures facilitate the removal of moisture and volatile matter from the precursor material, fostering the creation of pores, which are essential as adsorption sites. Consequently, an increase in temperature often results in the generation of a more extensive network of pores, enhancing the adsorption capacity of materials. However, this comes at the cost of the carbon content in the activated carbon, impacting its purity and suitability for specific applications.

Furthermore, the choice of chemical treatment is pivotal in determining the optimal activation temperature. Various activating agents interact differently with the carbonaceous components in biomass, leading to diverse outcomes in terms of pore structure and surface characteristics. Studies conducted by numerous researchers have underscored the significance of selecting the appropriate chemical treatment based on the reactivity with the precursor material and its specific chemical properties.¹⁴⁹–¹⁵¹ Factors such as boiling and melting points, as well as the ability to mobilize metallic ions, further influence the effectiveness of the activation process. Understanding these intricate relationships among chemical treatment, activation temperature, and the resultant properties of activated carbon is essential for tailoring materials to meet the diverse needs of industrial, environmental, and technological applications.

4.1.3.1 Optimum KOH activation temperature. Determining the optimal activation temperature for KOH is paramount for the production of activated carbon, given that it profoundly influences the properties and performance of materials. This review summarizes a vast body of literature dedicated to pinpointing the ideal activation temperature using KOH as the activating agent. The activation temperature serves as a critical parameter in modulating the pore structure, surface area, and adsorption properties of the resulting activated carbon. By collating insights from various studies, this review endeavors to elucidate the multifaceted factors shaping the selection of the optimum activation temperature for KOH, considering variables such as precursor material, activation duration, and intended applications. Dewi asserted that the most effective activation temperature for KOH treatment is in the range of 700 °C to 800 °C.¹⁵² Correspondingly, Williams et al. observed a significant enhancement in surface area and micropore volume with an increase in activation temperature from 700 °C to 800 °C.¹⁵³ The surface area experienced a remarkable 33% enhancement, increasing from 1955 $\mathrm{m^{2}\,g^{-1}}$ to 2600 $\mathrm{m^{2}\,g^{-1}}$, in contrast to the less than 10% improvement witnessed when the temperature was increased from 600 °C to 700 °C. This trend was corroborated by a group of researchers, further demonstrating that elevating the activation temperature from 700 °C to 800 °C led to a substantial increase in surface area, reaching up to 3228 $\mathrm{m^2\,g^{-1}}$, albeit at the expense of carbon yield and surface area when the temperature exceeded 850 $^{\circ}$ C.¹⁵⁴⁻¹⁵⁶ Additionally, Ma et al. observed a progressive widening of the average pore width from 0.520 nm at 500 °C to 0.573 nm at 800 °C.¹⁵⁶ This exponential enhancement can be attributed to the formation and mobilization of metallic potassium at around 700 °C, facilitating pore development. Furthermore, the interaction among potassium, carbon and water yields potassium carbonate, which mitigates precursor burn-off, contributing to enhanced pore formation. Understanding these intricate temperature-dependent mechanisms is indispensable for optimizing KOH activation processes and tailoring activated carbon for diverse applications. PSC Advances are the interest in more article in the studies, this resistance is the competition of the best competition of the section common access Article is likely are the section of the section of the common common c

4.1.3.2 Optimum H_3PO_4 activation temperature. Determining the most effective activation temperature for H_3PO_4 is crucial in producing activated carbon, given its significant influence on the characteristics and performance of materials. This review presents an in-depth exploration of the extensive research dedicated to identifying the optimal activation temperature using phosphoric acid as the activating agent. The activation temperature plays a pivotal role in shaping the pore structure, surface area, and adsorption capabilities of the resulting activated carbon. By summarizing the insights from various

studies, this review aims to highlight the factors influencing the selection of the best temperature for H_3PO_4 activation, considering variables such as precursor material, activation duration, and intended applications. Notably, empirical studies suggest that the most suitable activation temperature for H_3PO_4 -treated precursors is typically 500 °C or lower, regardless of the impregnation ratio or precursor source.^{152,157} For instance, researchers treated Paulownia wood with H_3PO_4 at a ratio of 4 and subjected it to carbonization in the temperature range of 300 °C to 600 °C.¹⁵⁸ The results indicated that higher temperatures were associated with a decrease in the activated carbon yield. Additionally, a notable trend emerged, wherein the micropore volume and BET surface area exhibited trend for activated carbon carbonized in the range of 300 °C to 400 °C, followed by a decline as the carbonization temperature increased from 400 $\mathrm{^{\circ}C}$ to 600 $\mathrm{^{\circ}C}.^{152,157}$ This phenomenon was attributed to the contraction of the carbon structure on Paulownia wood at elevated temperatures, resulting in a diminished void space. Moreover, the activated carbon produced in these experiments demonstrated a dual-porosity nature, comprising micropores and mesopores, which is consistent with ndings from prior studies utilizing alternative precursor materials such as date pits.

The observed trends in activation behavior can be elucidated by considering the chemical properties and organic chemistry of phosphoric acid. In the temperature range of 300 °C to 400 ° C, phosphoric acid not only acts as a heat facilitator but also as a dehydrating agent, capturing minerals to inhibit ash formation, and thereby promoting pore development. Additionally, H_3PO_4 forms a protective layer on carbon in liquid materials and various chemical structures such as acetic acid, methanol, and polyphosphates, which reinforce the pore stability against temperature-induced collapse. However, at higher temperatures, these supporting structures, particularly acetic acid, encounter resistance limits. Beyond 440 °C, acetic acid decomposes into a mixture of carbon dioxide and methane or water and ketene, resulting in compromised pore stability and subsequent surface area reduction. Thus, understanding these intricate temperature-dependent mechanisms is vital for optimizing the H_3PO_4 activation processes and tailoring activated carbon for diverse applications in industries, environmental remediation, and emerging technologies.

4.1.3.3 Optimum ZnCl₂ activation temperature. Determining the optimal activation temperature for $ZnCl₂$ is a critical factor in the production of activated carbon, given its significant impact on the properties and performance of materials. This review aims to delve into the extensive research dedicated to identifying the ideal activation temperature using zinc chloride as the activating agent. The activation temperature plays a pivotal role in molding the pore structure, surface area, and adsorption properties of the resulting activated carbon. By summarizing the insights from various studies, this review endeavors to elucidate the factors influencing the selection of the optimum temperature for $ZnCl₂$ activation, considering variables such as precursor material, activation duration, and intended applications. According to the study conducted by a research group, the optimal activation temperature for $ZnCl₂$

treatment was suggested to be 500 °C, which is attributed to the higher surface area and micropore volume reported for activated carbon precursor cherry stones.¹⁵⁹ This finding was corroborated by other researchers, where an increase in temperature from 500 °C to 800 °C resulted in a decrease in surface area.¹⁵¹ Additionally, Nurdin et al. reported the highest surface area of activated carbon precursor oil palm shell at 1429.71 m² g⁻¹ when activated at 500 °C, with a high micropore volume also observed.¹⁴⁹ However, an increase in activation temperature from 500 \degree C to 800 \degree C led to a decrease in specific surface area and micropore volume, coupled with an increase in average pore size distribution, as supported by similar research utilizing different precursors.¹⁵² Furthermore, in the adsorption-based study by Njeh et al., the highest iodine and methylene blue adsorption capacities were recorded for sawdust activated at 500 °C.¹⁶⁰ It was noted that $ZnCl₂$ can vaporize at temperatures above 400 °C, potentially leading to rapid carbon burnout due to the loss of protective carbon elements, according to Wei et al.¹⁶¹ However, Huang disagreed these claims, suggesting that the optimum activation temperature for $ZnCl₂$ treatment is around 700 °C.¹⁶² Though scarce, this assertion is reinforced by similar findings, where the highest micropore formation upon activation up to 800 °C is possibly linked to the physical properties of $ZnCl₂$, with a boiling point of 732 °C.^{151,163} In summary, higher temperatures generally result in increased pore formation, but each activating agent exhibits a specific range of optimum temperatures, depending on their chemical and physical properties such as boiling point. According to the literature, the suggested activation temperatures for precursors incorporating KOH, H_3PO_4 , and ZnCl₂ are 700–800 °C, 500 °C, and 500 °C, respectively. Review Wextwert Excellential Contents Articles. The Signal Common Contents Article and Theorem and Herbert Common Contents Article 20:10:22. This are also the common Common Common Common Common Common Common Common Common

4.1.4 Impact of impregnation ratio. The impregnation ratio stands as a pivotal factor shaping the quality and attributes of activated carbon manufacture, delineating the proportion of activating agent to precursor material employed throughout the process. This ratio has a profound influence on the ultimate pore structure, surface area, and adsorption capacity of the resulting activated carbon. Fine-tuning the impregnation ratio is paramount in customizing activated carbon for specific applications, demanding equilibrium among parameters such as porosity, yield, and cost-effectiveness. Additionally, another crucial element that greatly affects the characteristics of the final activated carbon is the impregnation ratio during chemical activation processes. This signifies the ratio between the activating agent weight and the raw material impregnated. Fundamentally, a higher impregnation ratio tends to induce more swelling, thereby facilitating the more vigorous release of volatile matter contents, consequently widening the pores. Conversely, lower concentrations of activating agent foster the steadier elimination of volatile matter content, while curbing tar deposition, resulting in the formation of more micropores.

In the case of biomass-derived materials, the impregnation ratio is particularly important, influencing their structural and functional characteristics. Several studies conducted by researchers underscore the significance of the impregnation ratio in shaping the pore structures.151,164,165 The typical impregnation ratios for biomass are in the range of 1 : 1 to 3 : 1, depending on the type of biomass and the specific requirements of the end product.¹⁶⁶ Studies have highlighted the effect of the impregnation ratio on the structural properties of activated carbons. For example, increasing the impregnation ratio has been shown to improve the porosity and surface functionality of activated carbon from coconut shells, enhancing its adsorption capabilities.¹⁶⁷ Additionally, optimizing the impregnation ratio is crucial for achieving a well-balanced pore structure and thermal stability, as seen in biochar produced from rice husks. Research also indicates that an impregnation ratio of 1 : 1 can result in activated carbons with higher surface areas and larger pore volumes compared to lower ratios.¹⁶⁸ Increasing the ratio of phosphoric acid from $1:1$ to $2:1$ was found to significantly increase the micropore volume in activated carbons.¹⁶⁹ However, it is important to note that excessively high impregnation ratios can lead to the overactivation of carbon materials, compromising their structural integrity. Therefore, it is essential to carefully optimize the impregnation ratio to achieve the desired balance among surface area, pore structure, and mechanical stability, tailoring the properties of activated carbon to meet specific application needs.

In essence, the impregnation ratio plays a critical role in the production of activated carbon, shaping its pore structure and surface area, and consequently affecting its adsorption capacity and performance. The optimal adjustment of this ratio is indispensable for tailoring activated carbon to specific applications, harmonizing diverse factors such as porosity, yield, and cost-effectiveness. A profound comprehension of the intricate interplay between the impregnation ratio and properties of activated carbon is pivotal for refining production processes and devising activated carbon materials endowed with enhanced performance attributes.

4.1.5 Impact of inert gases on carbonization process. The application of inert gases during the carbonization process plays a crucial role in determining the structural characteristics of the resulting activated carbons. Inert gases create an oxygenfree environment, which prevents oxidation and undesirable side reactions that can compromise the quality of the carbon material. Carbonization involves the thermal breakdown of organic substances in this inert atmosphere, producing a carbon-rich material with unique textural properties.¹⁷⁰ The choice of inert gas, together with the specific conditions under which it is applied, has a significant impact on the development of the porosity, surface area, and adsorption capabilities of materials.

Research indicates that the temperature at which carbonization occurs and the flow rate of the inert gas, typically nitrogen are critical factors in determining the porous architecture of carbon materials. Studies have shown that the optimal conditions for achieving the maximum specific surface area, pore volume, and micropore content are carbonization temperature in the range of 800–850 $\mathrm{^{\circ}C}$ with a nitrogen flow rate of 7.5-15 dm^3 min⁻¹. These parameters help in effectively removing volatile components from the precursor material, promoting the formation of a well-developed porous structure and increasing the surface area.¹⁷¹

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Inert gases such as nitrogen play multiple roles during the carbonization process. Primarily, they prevent the oxidation of the carbon material by displacing oxygen and other reactive gases, thus maintaining the integrity of the carbon structure. Additionally, the flow rate of nitrogen influences how quickly volatile compounds are expelled from the carbon matrix. A higher nitrogen flow rate facilitates the faster removal of volatiles, enhancing the development of a more extensive pore network and higher surface areas. Alternatively, a lower flow rate can lead to a less porous structure due to the slower release of volatiles and potential re-deposition of carbonaceous residues within the pores.^{96,171}

Furthermore, the interaction between the carbonization temperature and inert gas flow rate is vital in achieving the desired textural properties. Higher temperatures enhance the decomposition of the precursor material, and when combined with a controlled nitrogen flow, can result in a uniform and highly porous structure. However, if the temperature is too high or the nitrogen flow is inadequate, the pore structure may collapse or unwanted tar formation may occur, clogging the pores and reducing the overall effectiveness of the material.¹⁷²

Overall, the use of inert gases such as nitrogen during carbonization is critical for regulating the textural characteristics of activated carbon. By optimizing the carbonization temperature and gas flow rate, it is possible to fine-tune the pore structure and surface area, thereby enhancing the adsorption capacity and overall performance of the material.

4.2 Utilizing waste-derived activated carbon for $CO₂$ capture

The widespread application of activated carbon spans multiple sectors, encompassing gas and water purification, gold processing, wastewater treatment, metal extraction, decaffeination,

and pharmaceuticals. The rapid expansion of the AC market is underscored by the significant rise in its annual demand. According to Market Data Forecast, the global AC market is projected to grow from \$9.17 billion in 2023 to \$11.46 billion by 2029, indicating robust growth.⁹⁶ Thus, to meet escalating demands and address cost considerations, researchers are actively exploring alternative pathways for the production of AC, with a specific emphasis on low-cost or waste materials. The utilization of waste materials not only addresses environmental and waste management challenges but also unlocks lucrative opportunities for the business sector. In parallel, biomass can be broadly classified into two main categories, conventional waste and non-conventional waste. Conventional waste is comprised of biomasses sourced from the agricultural and wood industries, such as coconut shells, rice straws, sawdust, and wood, while non-conventional wastes include materials such as plastic, sugarcane bagasse, citrus peel, and tires, originating from municipal and industrial sources. Coconut shells, in particular, have attracted significant attention as a frequently studied precursor due to their elevated carbon content and robust mechanical properties, as evidenced by the compositional analysis in Fig. 9. Comparative studies conducted by several researchers have focused on determining the adsorption capacity and practical feasibility of coconut shell-derived AC precursors.¹⁷³–¹⁷⁶ The literature underscores the importance of understanding the carbon content and compositions of potential raw materials, given that these factors profoundly influence the characteristics of the resulting AC. Crucially, parameters such as fixed carbon and volatile matter levels play pivotal roles in evaluating the effects of carbonization and pore formation processes, with carbonization notably enhancing pore development by augmenting the mobility and reducing the volatile matter content. **PESC Advances** Vew Rets Articles. The main of the significant interior in the significant interior of the significant interior in the significant interior in the significant interior in the significant interior in the sig

Table 4 Variations in compositional elements linked to diverse precursors

The analysis presented in Table 4 highlights that nutshell, coconut shell, algae, and sea mango possess a carbon content exceeding the average of 44.95 wt%. Other studies suggest that utilizing a concentrated carbon precursor can potentially enhance the yield of AC by leveraging its higher carbon content, thereby facilitating the formation of microporous structures.²¹⁵ Moreover, gaining a thorough understanding of the structural intricacies of various waste materials, influenced by variations in lignocellulose compositions, is considered crucial.

Comparative studies indicate that precursors abundant in lignin tend to yield activated carbon with spherical porous structures, while that rich in cellulose tends to produce cylindrical pore structures. These observations emphasize the pivotal role of precursor type in determining the texture of AC.177,216 However, achieving the optimal activated carbon for $CO₂$ adsorption poses significant challenges due to the diverse composition properties of waste materials from different sources. Notably, it can be observed in Table 4 that algae, walnut shell, and waste palm shell exhibit elevated nitrogen levels compared to other waste samples. With nitrogen levels of 7.20 wt%, 3.20 wt%, and 3.00 wt%, respectively, surpassing the average nitrogen content of 1.67 wt% among the samples. Despite the significance of the carbon content in precursor selection, this indicates that a higher nitrogen concentration in the precursor enhances both the carbon dioxide adsorption performance and $CO₂$ selectivity, irrespective of the carbon composition.^{178,217} Treeweranuwat et al. emphasized that a low carbon content in the precursor leads to a reduction in the surface area of the resulting activated carbon. However, incorporating heteroatom-rich materials during the carbonization and activation of shrimp-derived AC not only increased its surface area but also boosted the carbon yield.²¹⁸ Furthermore, the addition of melamine, sodium thiosulfate, and KOH was shown to double the AC yield due to the enhanced thermal stability.

An essential criterion for an effective precursor in the production of adsorbents is a minimal ash content.²¹⁹ Douglas fir stands out with its notably low recorded ash content of 0.21 wt% (as indicated in Table 4), which is signicantly lower than the average of 4.12 wt%. In contrast, algae and rice husk exhibit a higher ash content of 6.70 wt% and 16.70 wt%, respectively. The reduced ash content in Douglas fir suggests its potential as a superior adsorbent precursor. Although the ash content does not directly influence the development of porosity, it impacts the adsorption properties due to the formation of inert sites. Understanding the prevalent lignocellulose structure is crucial due to the diverse compositions of waste materials. Comparative analyses indicate that precursors rich in lignin tend to yield activated carbon with spherical porous structures, while that with a higher cellulose content tends to yield cylindrical pore structures.^{177,180} The varying composition properties of waste materials pose challenges in achieving the optimal activated carbon for $CO₂$ adsorption. Agricultural residues, such as palm kernel shells, rice husks, coconut shells, and bagasse, are extensively utilized for producing activated carbon due to their abundance, high carbon content, and relatively low ash levels. Conducting a comprehensive precursor analysis is essential to facilitate the conversion of AC into a suitable $CO₂$ adsorbent. Besides possessing a substantial surface area and high microporosity concentration, waste-derived synthesized AC materials must be economically feasible, possess basic functionalities, exhibit low heat absorption, high $CO₂$ selectivity, and be suitable for industrial-scale production. Also, the composition of waste products plays a crucial role in determining the quality of the resulting activated carbon.

4.2.1 Utilizing agricultural waste for the synthesis of activated carbon. Utilizing agricultural waste for the synthesis of activated carbon presents a signicant opportunity to address pressing environmental challenges, while simultaneously contributing to sustainable waste management practices and resource optimization. This comprehensive review critically examines a range of agricultural residues and elucidates their conversion methodologies, resulting properties, and myriad applications of the produced activated carbon. Through meticulous investigation, this review underscores the

transformative potential of agricultural waste as a versatile and eco-friendly alternative across diverse industrial and environmental sectors. Recent research efforts have increasingly focused on harnessing agricultural waste as a valuable source of activated carbon precursors tailored for $CO₂$ adsorption applications. For instance, in the study by Hidayu and Muda, palm kernel shells and coconut shells were employed as primary raw materials for the synthesis of activated carbon.²¹⁹ These agricultural residues were subjected to carbonization at 800 $\mathrm{^{\circ}C}$ after being ground to a uniform size, followed by physical and chemical activation using $ZnCl₂$ in a 1:1 ratio. This study revealed that coconut shells exhibited a higher yield of activated carbon precursors compared to palm kernel shells, positioning them as promising candidates for raw material utilization in carbon capture applications.

Similarly, another research group investigated the efficacy of utilizing bagasse and rice husk as precursor materials for the production of activated carbon.²⁰⁰ Employing $ZnCl₂$ as an activator at a temperature of 500 °C for 60 min, this study demonstrated that bagasse-derived activated carbon displayed superior adsorption capabilities over its rice husk-derived counterpart across various adsorption temperatures. These ndings underscore the potential of leveraging readily available agricultural waste streams for the scalable and economically viable production of activated carbon, thereby facilitating the transition towards a more sustainable waste management paradigm. However, despite the notable advancements in $CO₂$ adsorption performance observed in these studies, challenges persist in ensuring the thermal stability of the synthesized activated carbon for sustained $CO₂$ capture efficacy. Particularly, the decline in $CO₂$ adsorption performance at elevated temperatures underscores the need for targeted modifications to enhance the thermal stability of activated carbon materials tailored for $CO₂$ capture applications. According to the findings gleaned from extensive literature studies, coconut shell and palm kernel shells emerge as the most promising agricultural waste precursors for the production of activated carbon, owing to their high carbon content and low ash composition.^{176,219} This ranking underscores the critical importance of selecting appropriate precursor materials to optimize the performance and sustainability of activated carbon production processes. **PEC Advances**

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> 4.2.2 Utilizing industrial waste for the synthesis of activated carbon. The utilization of industrial waste for synthesizing activated carbon has emerged as a compelling strategy for promoting sustainable resource management and environmental remediation endeavors. This comprehensive review delves into the diverse array of industrial by-products and waste materials, assessing their potential as viable precursors for the synthesis of activated carbon, while elucidating the intricacies of various conversion techniques employed in their transformation. For instance, in a study, sewage sludge was investigated as a precursor for $CO₂$ adsorption, undergoing activation with KOH and NaOH at temperatures in the range of 600 °C to 800 °C.²²⁰ Despite yielding activated carbon with a relatively modest surface area of 179 m² g^{-1} , this material exhibited an enhanced $CO₂$ adsorption capacity, which was attributed to its high content of basic functional groups. Notably, optimal $CO₂$

adsorption was achieved when solid NaOH was utilized as the activating agent, facilitating enhanced interaction between hydroxide and the carbon elements within the sewage sludge matrix. Furthermore, Wang et al. explored alkaline activation (KOH) to develop activated carbon fibers, varying the KOH to fiber ratio and activating the samples at 700 °C.²²¹ The resulting activated carbon fibers demonstrated a high surface area and micropore concentration, with the incorporation of nitrogen contributing to expedited sorption kinetics, which is a desirable attribute for CO₂ adsorbents. Similarly, another research group investigated municipal solid waste compost as a precursor, activating the waste using H_2SO_4 in the temperature range of 400 °C to 800 °C.²²² Interestingly, the activated carbon impregnated with H₂SO₄ before pyrolysis exhibited superior properties such as enhanced surface area and micropore volume compared to the post-pyrolysis incorporation of the activating agent, with a notable CO₂ adsorption of 2.6 mmol $\rm g^{-1}.$ Through the repurposing of industrial waste streams, this approach not only tackles waste management challenges but also presents a cost-effective and environmentally friendly avenue for producing activated carbon with multifaceted applications in pollutant removal, energy storage, and beyond. Review Web articles. The control of the methods control of the methods are the control of the set of

4.2.3 Utilizing food waste for the synthesis of activated carbon. Exploring the potential of food waste for the synthesis of activated carbon represents a pivotal step towards sustainable waste management practices and resource optimization. This review critically examines various types of food waste as promising precursors for activated carbon production, elucidating their respective conversion methodologies and the resulting properties of the synthesized activated carbon. For instance, researchers utilized chitin aerogel as a raw material, activating it with potassium hydroxide at a ratio of 3 : 1 activating agent to carbon, and subjecting it to activation at 850 °C for 4 h.²²³ Similarly, Jia et al. investigated shrimp shells as activated carbon precursors, leveraging their high nitrogen content to enhance the formation of basic functional groups in the synthesized activated carbon.²¹⁷ In another study, coffee grounds were utilized as the raw material and activated with KOH at 400 °C, yielding activated carbon with an average $CO₂$ adsorption capacity of 0.27 cm³ g⁻¹ at 0 °C.²²⁴ Moreover, Wen et al. demonstrated the potential of beer waste as an activated carbon precursor, employing hydrothermal carbonization followed by activation at 800 °C for 2 h, resulting in activated carbon with high $CO₂$ selectivity compared to other studied precursors.²²⁵ Comprehending the structural and elemental makeup of waste materials is crucial for confirming their viability as precursors for activated carbon in $CO₂$ capture. Although agricultural residues are favored due to their abundance, high carbon content, and low ash content, food waste shows promise despite its lower average carbon content, which is attributed to its high nitrogen heteroatoms. Furthermore, industrial residues often possess elevated concentrations of functional groups, potentially enhancing the carbon capture capabilities. Consideration of factors such as volatile matter fraction, ash content, moisture content, and compositional complexity of biomass is crucial to ensure the uniform performance of the produced activated carbon, particularly for

commercial applications where a high variation in carbon dioxide adsorption capacity is intolerable. Agricultural wastes are favored due to their abundance, ease of management, and the ability to categorize the raw materials based on species, age, and origin. By repurposing food waste streams, this approach not only mitigates food waste disposal challenges but also offers an environmentally friendly solution for producing activated carbon with diverse applications in water treatment, air purification, and energy storage.

5. Application

5.1 $CO₂$ capture with carbon materials

Carbon materials are increasingly recognized for their potential in industrial applications, particularly due to their costeffectiveness, efficient regeneration, and exceptional ability to capture substantial amounts of $CO₂$. These materials play a crucial role in addressing the growing need for $CO₂$ capture from various emission sources, leveraging their high surface area and adaptable pore structures to serve as effective adsorbents for $CO₂$ molecules. The versatility of carbon materials extends beyond $CO₂$ capture, where they are also employed in pollutant adsorption, catalysis, and energy storage, making them vital in confronting environmental challenges and promoting sustainable technologies. Recent research has focused on the use of carbon materials for $CO₂$ capture, largely driven by the increasing release of carbon dioxide from industrial activities, which exacerbates the global warming crisis. In this case, biomass, rich in carbon content, is emerging as a valuable resource for producing biochar or hydrochar, contributing to efforts in reducing $CO₂$ emissions. However, pristine carbon materials face limitations in their textural structure and chemical composition, which restricts their effectiveness in $CO₂$ capture. Thus, to overcome these challenges, pre-capture modifications are necessary to enhance the properties of carbon materials and improve their $CO₂$ capture efficiency. This review critically examines the reported studies on biomass-derived carbon materials for $CO₂$ capture, exploring the underlying adsorption mechanisms and identifying promising avenues for future research and practical application. By understanding these mechanisms and advancing the development of modified carbon materials, it is possible to optimize the $CO₂$ capture processes, thereby contributing to the global effort to mitigate the impacts of climate change and move towards a more sustainable future.

5.1.1 The process of $CO₂$ capture. Capturing $CO₂$ using carbon materials is a process in which carbon dioxide molecules adhere to the surface of these materials, aided by their large surface area and specialized pore structures, which enhance adsorption.²²⁶ This process involves a variety of interactions, including van der Waals forces, π – π interactions, and electrostatic forces, which draw $CO₂$ molecules to the surface of carbon materials. The effectiveness of carbon materials in $CO₂$ capture stems from their high porosity, extensive surface area, and well-optimized pore size distribution, which enhance their interaction with $CO₂$. The $CO₂$ capture mechanism in biomassderived carbon materials occurs through two primary

processes, i.e., physical adsorption and chemical adsorption. Physical adsorption, or physisorption, is an exothermic process dependent on temperature and pressure, where $CO₂$ molecules adhere to the surface via van der Waals forces and other noncovalent interactions.²²⁷ This process is energy-efficient for regenerating the adsorbent, especially under higher pressure conditions.²²⁸ Alternatively, chemical adsorption, or chemisorption, involves the formation of covalent or ionic bonds between $CO₂$ molecules and the functional sites on the adsorbent, driven by acid–base neutralization reactions.²²⁹ Although chemisorption is energetically favorable during the adsorption phase, desorption and adsorbent regeneration require more energy. By incorporating basic organic groups or inorganic metal oxides, such as amines and alkali metals, respectively, the efficiency of chemisorption can be signicantly enhanced. For example, polyethyleneimine-functionalized mesoporous silica and nitrogen-doped activated carbon have shown increased CO₂ capture performances through these mechanisms.²³⁰ These ndings underscore the importance of both adsorption processes in optimizing the $CO₂$ capture efficiency, making carbon materials derived from biomass crucial for advancing sustainable carbon capture technologies. When captured, $CO₂$ molecules are temporarily held on the carbon surface, effectively extracting them from the gaseous phase. The continual release of anthropogenic $CO₂$ into the atmosphere has led to the significant and escalating issues of global warming and climate change.^{231,232} Between 1970 and 2004, the annual global $CO₂$ **PEC Advances** (*i.e.*, physical assorption and elembral assorption). San excellented on 2024. Downloaded on 2022. The creative of the creative is licensed under the creative Common Access Article is a reached on the distr

emissions substantially increased by 80%, contributing to the current $CO₂$ concentration in the air exceeding the maximum permissible limit for climate safety and forecasted to reach 700 ppm by the end of the century. As a result, a temperature rise of 1.8 °C to 4 °C is expected due to global warming, posing significant risks to the ecosystem and causing irreparable harm to the environment.²³³ This surpassed climate safety threshold underscores the urgent need for remedial measures to mitigate anthropogenic greenhouse gas emissions. $CO₂$ capture methods can be applied directly from the atmosphere or from industrial processes flue gas streams, employing suitable techniques.²³⁴ Subsequently, the captured $CO₂$ can be utilized in various strategies, including the production of clean energy fuels for the future or as a carbon source for the synthesis of industrially important chemicals.

Numerous adsorbents have been proposed for $CO₂$ capture, each possessing unique characteristics that contribute to their effectiveness in this application. As shown in Fig. 10, an optimal $CO₂$ adsorbent should exhibit high adsorptive capacity, selectivity for $CO₂$ in gaseous mixtures, rapid sorption kinetics, a microporous structure, and morphological and chemical stability, while also offering a low heat of adsorption, high regeneration rate, and cost-effectiveness.²³⁵ The following table presents a comprehensive overview of various AC and carbon materials derived from different precursors, including details on their synthesis and activation methods. It includes information on the precursor materials, synthesis techniques,

Fig. 10 Characteristic properties required for an ideal adsorbent for $CO₂$ capture.

Table 5 Properties of activated carbons from various precursors and activation methods

activation agents, and the ratio of precursor to activation agent used. Additionally, this table summarizes key textural properties such as surface area, pore size or volume, and pore size distribution (PSD), as well as the surface functionalities of each material. This comparative analysis provides valuable insights into the performance and characteristics of different carbon materials for applications such as $CO₂$ capture (Table 5).

Recent studies on $CO₂$ capture, both pre- and postcombustion, have investigated various techniques, including adsorption on functionalized porous materials, absorption in liquid amine solutions, membrane separation, and cryogenic separation at ultra-low temperatures.²⁶⁰ However, each method has its strengths and limitations, such as the use of harsh chemicals, equipment wear and tear, disposal challenges, performance inconsistencies, and high associated costs. In contrast, activated porous carbons derived from biomass have emerged as promising and economically viable options for addressing the elevated $CO₂$ levels in the atmosphere. These carbons produced via biomass pyrolysis offer a sustainable approach by converting unwanted biomass materials into efficient adsorbents suitable for large-scale $CO₂$ capture.²⁶¹ Porous carbon materials offer a triple advantage in $CO₂$ emission reduction, as follows: firstly, by mitigating $CO₂$ emissions from decomposing biomass; secondly, by amending soils with porous carbon to diminish $CO₂$ generation; and thirdly, by utilizing porous carbons for $CO₂$ capture in flue gas streams during both pre- and post-combustion processes. This section delves into the potential applications of biochar and activated porous carbons derived from various biomass precursors for $CO₂$ capture. Although pristine biochar may possess limited porosity and efficacy for $CO₂$ capture, activation procedures can enhance its surface area and porosity, making it suitable for this purpose. Researchers have explored diverse biomass sources

and experimental parameters to develop various types of activated porous carbons, aiming to optimize their $CO₂$ capture efficiency.

5.1.2 Optimizing carbon for $CO₂$ capture. Carbon materials are widely employed for $CO₂$ capture due to their expansive surface area and customizable pore structures, making them effective adsorbents for $CO₂$ molecules. These materials play a vital role in addressing the urgent challenge of reducing $CO₂$ emissions from diverse sources, spanning industrial processes to atmospheric sources. By harnessing their inherent attributes, such as high adsorption capacity and selectivity, carbon materials offer a promising avenue for capturing $CO₂$ and mitigating its atmospheric concentration. Various strategies have been explored to enhance the $CO₂$ capture capabilities of carbon materials. Table 6 offers an overview of carbon materials prepared through different modification methods, together with a comparison of their $CO₂$ capture efficiencies. Pristine carbon materials often exhibit limited pore structures and surface chemistry, constraining their $CO₂$ capture effectiveness. For example, the pristine biochar derived from coffee grounds displayed a modest CO₂ capture capacity of 0.14 mmol g^{-1} , whereas KOH activation (500 °C, 1 h) significantly increased it to 4.76 mmol g⁻¹.²⁶⁹ Thus, modifications such as acid or alkaline activation, heteroatom doping, and metal impregnation are indispensable for achieving heightened $CO₂$ capture capacities. The efficacy of these modifications relies heavily on the choice of modifier, treatment temperature, and duration. One research group explored the synthesis of porous carbon from tobacco stems via KOH activation at varying temperatures, observing that higher temperatures yielded an augmented surface area and micropore volume, initially enhancing the $CO₂$ capture capacity before reaching a saturation point.²⁶² This phenomenon was ascribed to the reduction in oxygen content in the **PEX Advances**
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carbon framework at elevated temperatures, typically enhancing the adsorption of $CO₂$. Additionally, activated carbon was produced from corn stover and activated using H_3PO_4 , CO_2 , and H_2O , with the H_3PO_4 -activated carbon exhibiting the highest CO $_2$ capture capacity of 10.9 mmol $\mathrm{g}^{-1},$ owing to its elevated specific surface area and microporous volume resulting from chemical activation.²⁶⁴ Moreover, urea-doped nanoporous carbon synthesized from walnut shells showed an increase in $CO₂$ capture capacity with higher urea doping levels.²⁶⁶ Furthermore, the $CO₂$ uptake of activated carbon was enhanced by modification with NiO and CuO, respectively, resulting in notable improvements in $CO₂$ capture capacity.²⁶⁷ Subsequent sections of this review will explore the application of various types of carbon materials in $CO₂$ capture, including pristine carbon, activated carbon, heteroatom-doped carbon, metal-loaded carbon, and carbon-based nanomaterials, examining their individual advantages and challenges in $CO₂$ capture.

5.1.2.1 Pristine carbon materials. Pristine carbon materials, in their natural state, serve as the fundamental building blocks for various applications, including the capture of $CO₂$. However, despite possessing inherent carbonaceous properties such as surface area and porosity, pristine carbon materials often have limited $CO₂$ adsorption capabilities due to their underdeveloped pore structures and surface chemistry. These materials are directly derived from the thermochemical conversion of biomass, such as biochar or hydrochar, well-known for their environmentally friendly characteristics and wide availability from diverse biomass sources. Despite their initial limitations, pristine carbon materials provide the foundation for subsequent modifications aimed at enhancing their $CO₂$ capture efficiency. Through the implementation of various activation methods and doping techniques, pristine carbon materials can be customized to meet specific requirements for $CO₂$ capture, making them a versatile platform for developing effective $CO₂$ adsorbents to counteract the rising atmospheric $CO₂$ levels. Numerous studies have highlighted the $CO₂$ adsorption potential of biochar, which is attributed to its polarity, high specific surface area, and functional groups. 270 For instance, composite biochar synthesized from sewage sludge and Leucaena wood in different proportions showed superior $CO₂$ uptake compared to pure sewage sludge biochar. Increasing the Leucaena wood biochar content resulted in a higher carbon content, and subsequently enhanced $CO₂$ adsorption capacity.²⁷¹ Similarly, date palm leaf biochar produced at varying temperatures exhibited a corresponding increase in $CO₂$ adsorption capacity with higher preparation temperatures, owing to the increased carbon content of the biochar.²⁷² These findings underscore the potential of pristine carbon materials, particularly biochar, as effective $CO₂$ adsorbents and emphasize the significance of optimizing preparation conditions to maximize their $CO₂$ capture efficiency.

5.1.2.2 Activated porous carbon. Activated porous carbon (APC) has emerged as a pivotal material for $CO₂$ capture due to its unique properties, including highly porous structure and large surface area, which enhance its adsorption capabilities. Traditionally, APC has been extensively utilized in various applications such as pollutant removal, industrial purification, and air filtration. Its versatility is attributed to its ability to be tailored through different activation methods, both physical and chemical, to achieve the desired porosity and surface characteristics. Biomass provides a distinctive opportunity for creating activated porous carbons with beneficial structural characteristics. This approach is considered more advantageous than using traditional fossil fuel-based precursors for the production of activated carbons due to low cost and abundant availability of biomass and the environmentally friendly nature of the pyrolysis process used. The process of converting biomass into APC involves pyrolysis, followed by activation using agents such as steam, $CO₂$, and chemical reagents such as KOH or ZnCl₂. This process results in a material with a high surface area and tunable pore size distribution, making it highly effective for capturing CO_2 ²⁷³

Physical activation, such as $CO₂$ or steam activation, plays a crucial role in enhancing the microporosity of the carbon structure, thereby increasing its specific surface area and improving $CO₂$ adsorption. The choice of biomass precursor and the conditions of the pyrolysis process, including temperature and residence time, significantly influence the final properties of the APC. For instance, $CO₂$ activation is known to produce microporous carbons with uniform pore structures, which are ideal for gas adsorption, while steam activation can introduce mesopores and macropores, potentially enhancing the diffusion of $CO₂$ molecules into the carbon matrix.²⁷⁴ Alternatively, chemical activation can further enhance the porosity and surface functionality of APCs, making them even more effective for $CO₂$ capture. Activating agents such as KOH and $ZnCl₂$ facilitate the development of a hierarchical pore structure and the introduction of functional groups that can interact with $CO₂$ molecules, improving the adsorption efficiency.^{226,275} The ability to engineer APCs with specific surface properties through controlled activation processes makes them highly promising materials for addressing the growing challenge of $CO₂$ emissions and climate change. Review Wext Scheme The article on 18 2022. Article on 10 20: Additional Access Article is articled on 18 2022. And the article is articled on 18 202. And the article of the article of the article of the article is article

> Fig. 11 illustrates the process of $CO₂$ capture using activated porous carbon derived from biomass, which was chemically activated with potassium hydroxide (KOH). The process begins with the raw biomass of Jujun grass, which undergoes KOH activation to create a highly porous carbon material. Subsequently, this activated porous carbon is used for $CO₂$ adsorption, effectively capturing $CO₂$ molecules at 25 °C as a function of pressure.²⁷⁶ This highlights the potential of KOH-activated porous carbon as a highly efficient material for $CO₂$ capture, which is crucial for mitigating the impact of greenhouse gases on the environment.

> 5.1.2.3 Non-functionalized activated carbon. Nonfunctionalized activated carbon derived from various precursor materials undergoes activation processes without additional chemical modifications to its surface, preserving its inherent physical and chemical attributes. In contrast to functionalized activated carbon, which incorporates specific functional groups to target adsorption, non-functionalized activated carbon relies solely on its porous structure and high surface area for adsorption. Generally, activated carbons demonstrate

superior CO₂ adsorption capacities compared to biochar due to their improved textural properties and surface chemistry resulting from activation treatments. $CO₂$ capture mechanisms involve both physical and chemical processes. Physical adsorption arises from van der Waals forces between $CO₂$ molecules and the carbon surface, with surface area and pore size playing critical roles in the adsorption efficiency. Conversely, chemical adsorption is attributed to the alkali functional groups on the carbon surface, enhancing the interaction and uptake of $CO₂$. Activation significantly enhances the textural properties of carbon materials, thereby boosting their $CO₂$ adsorption performance.

Highly microporous KOH-activated carbons derived from Jujun grass (JG) and Camellia japonica (CJ) biomass exhibited significant $CO₂$ capture capabilities in both pre- and postcombustion scenarios.²⁷⁷ The activated carbons produced at 700 °C with a KOH impregnation ratio of 2 demonstrated the highest CO $_2$ adsorption at 1 bar (4.9–5.0 mmol $\rm g^{-1})$ under room temperature conditions. Conversely, that prepared at 800 °C with a higher KOH impregnation ratio of 4 showed the highest $CO₂$ adsorption at 20 bar (21.1 mmol $g⁻¹$). Mesquite wood, when transformed into carbon-rich material through singlestep KOH activation at 800 °C, achieved a high surface area of 3167 m² g⁻¹ and CO₂ adsorption of 26.0 mmol g⁻¹ at 23 °C and 30 bar.²⁶¹ Similarly, microporous activated carbons synthesized from Arundo donax at 600 °C displayed an excellent $CO₂$ adsorption capacity of 15.4 mmol g^{-1} at 30 bar, with a surface

area of 1122 $m^2 g^{-1}$. ZnCl₂ activation of Arundo donax resulted in an even higher surface area of 3298 m² g⁻¹ and CO₂ adsorption of 30.2 mmol g^{-1} at 30 bar.²⁷⁸ These results emphasize the critical role of surface area and porosity in enhancing the $CO₂$ adsorption by activated porous carbons.

The microporous content of activated carbon significantly influences $CO₂$ adsorption, with a higher micropore content correlated with increased $CO₂$ adsorption.²⁷⁹ Research on KOHactivated rice husk char showed that lower activation temperatures (640–710 °C), reduced KOH impregnation ratios (1:1), and elevated nitrogen content resulted in the high $CO₂$ uptake of 2.11 mmol g^{-1} at 0.1 bar. A narrow pore size distribution and micropores less than 0.7 nm were identified as crucial for the enhanced $CO₂$ adsorption at sub atmospheric pressures. These activated carbons also exhibited high selectivity (19.9) for $CO₂$ over N_2 . Another study highlighted the importance of ultramicropores in $CO₂$ capture using KOH-activated bamboo porous carbons, reporting a notable $CO₂$ adsorption value of 7.0 mmol g^{-1} at 0 °C and 1 bar, which was attributed to the micropores of 0.55 nm.²⁸⁰ However, wider pores resulting from increased KOH impregnation amounts led to reduced $CO₂$ adsorption due to the limited pore diffusion.

Likewise, Patel et al. highlighted the importance of narrow micropores (≤ 1 nm) in CO₂ capture by activated porous carbons produced through the KOH activation of sawdust.²⁸¹ Mild KOH treatment resulted in narrow micropores with lower surface areas compared to severe treatment, yielding high $CO₂$ uptake

 $(4.8 \text{ mmol g}^{-1})$ at room temperature. The presence of a microporous structure, hydroxyl groups, and heteroatoms such as nitrogen and oxygen collectively regulated the $CO₂$ adsorption on activated porous carbons derived from KOH activation of gelatin and starch biomasses, resulting in substantial $CO₂$ adsorption (7.49 mmol g^{-1}) at 1 bar and 0 °C, with favorable selectivity for CO_2 over N_2 (52–98%).²⁸² The isosteric heats of adsorption values (21.7–62.9 kJ mol $^{-1})$ suggest the involvement of both chemical and physical adsorption mechanisms. Additionally, wood ash-based activated porous carbon has been proposed for $CO₂$ capture, primarily via chemical sorption.²⁸³ Analysis pre- and post- $CO₂$ adsorption revealed the participation of alkali and alkaline earth metals in wood ash, facilitating chemical reactions with $CO₂$ and $H₂O$ and resulting in the formation of diverse carbonate products.

5.1.2.4 Heteroatom-doped activated porous carbon. Doping activated porous carbon with heteroatoms such as nitrogen, oxygen, and sulfur during activation modifies its electronic properties and surface chemistry, enhancing its adsorption capabilities and selectivity, particularly for $CO₂$. Although micropores are crucial for $CO₂$ capture, they also promote the adsorption of other gases such as CH_4 and N_2 , leading to reduced CO₂ adsorption capacity due to competitive adsorption. However, heteroatom doping addresses this issue by introducing functional groups that exclusively interact with $CO₂$, boosting the $CO₂$ selectivity. Incorporating basic or electronrich heteroatoms such as nitrogen into the carbon framework further enhances the $CO₂$ absorption capacity, given that the abundant basic sites on the surface act as anchors for capturing the weakly acidic $CO₂$ molecules. For example, Nazir et al. added nitrogen-containing functional groups to activated porous carbons for $CO₂$ extraction during carbonization using an NH₃ solution.²⁶⁵ This three-step process resulted in a final carbon material with a high nitrogen content (7.21%) incorporated as functional groups, exhibiting a higher surface area (2511 m^2) g^{-1}) compared to the carbon prepared using KOH activation alone (2064 $\mathrm{m^2\,g^{-1}}$). The enhanced surface area and basic sites facilitated significant CO₂ adsorption at both 0 \degree C (7.19 mmol $\rm g^{-1})$ and 25 °C (5.5 mmol $\rm g^{-1})$, with high ultra-microporosity aiding in physical and chemical $CO₂$ adsorption.²⁸⁴ Additionally, the sample demonstrated high selectivity for CO_2 over N_2 (30.75%), rapid kinetics, and efficient regeneration, presenting a promising approach for $CO₂$ capture using biomass-based materials.

In a study conducted by a research group, activated carbon was produced from husk using carbonization and KOH activation, with chitosan introduced during its activation.²⁰⁰ The resultant doped activated carbon displayed an enhanced $CO₂$ adsorption capacity, reaching 5.83 mmol g^{-1} at 0 °C. The XPS analysis detected pyridine-N and pyrrole-N on the surface of the doped activated carbon, enhancing its surface alkalinity and facilitating $CO₂$ uptake. Similarly, another research team enhanced the $CO₂$ capture performance of activated carbon by modifying it with ammonium sulfate.²⁸⁵ The presence of sulfur functional groups enabled interaction with $CO₂$ molecules, thereby improving their adsorption. Nitrogen doping plays a vital role in heteroatom doping to boost the $CO₂$ capture

efficiency. Nazir and Wang et al. employed various amines (TEPA, MEA, DEA, PEI, and DETA) to modify waste wood ashes for $CO₂$ capture. Among them, the TEPA-sorbent exhibited a superior $CO₂$ capture performance, with a capacity of 1.76 mmol g^{-1} and an impressive regeneration rate of 92.6% (Fig. 12).265,286 Despite the DEA-sorbents having the highest actual nitrogen loading, their adsorption capacity was lower than that of TEPA, highlighting the multifaceted nature of $CO₂$ adsorption, where the nitrogen content alone does not determine the capacity. The significance of micropores in $CO₂$ adsorption was demonstrated by the optimized loading of TEPA (45 wt%), which yielded a superior $CO₂$ capture performance (2.02 mmol g−¹). Furthermore, the type of nitrogen-containing functional groups significantly impacts the $CO₂$ capture efficiency. Wang et al. investigated the roles of various nitrogen functional groups in the $CO₂$ adsorption process, with pyridine-N showing the highest CO₂ adsorption energy $(-21.4 \text{ kJ mol}^{-1})$ due to its robust dipole–dipole interactions with $CO₂$, similar to H⁺ and O^{2-269,287} Thus, understanding these mechanisms is crucial for tailoring adsorbents for the optimal $CO₂$ capture performance. Review Wext Common temperature. The presence of a microscopic article, North and DEVA DOWN COM DOWN COMPOSITE and the composition-non-temperature is likely and the property and on the composition-non-temperature is likely

5.1.2.5 Metal/metal oxide-embedded activated porous carbon. Metal or metal oxide-embedded activated porous carbons stand out as a versatile and highly effective class of carbon-based materials, which are characterized by the integration of metal or metal oxide nanoparticles into their porous structure. These materials are synthesized using a variety of methods, such as impregnation, co-precipitation, and in situ growth, resulting in enhanced adsorption properties and catalytic activity owing to the incorporation of metal or metal oxide nanoparticles into the carbon matrix. Another strategy to boost the $CO₂$ adsorption capacity involves functionalizing activated porous carbons with metal atoms to create additional surface-active sites. Numerous studies have explored the impact of metal functionalization on $CO₂$ uptake, with alkaline metals proving particularly effective in adsorbing acidic carbon dioxide. In the adsorption process, $CO₂$ serves as an electron acceptor, capable of receiving electrons from metal oxides possessing basic characteristics.

The adsorption mechanism of $CO₂$ onto metal-impregnated AC is depicted in the Fig. 13. Metal oxides are typically incorporated in the surface of AC using the wet impregnation method. This involves mixing the carbon precursor with metal oxide solutions before subjecting it to a high-temperature carbonization process, or alternatively, mixing the already prepared AC with metal oxide solutions followed by calcination at elevated temperatures.¹³ For instance, Botomé et al. produced AC from wood treated with chromated copper arsenate (CCA), which was pyrolyzed at 700 °C, and the resulting material demonstrated an adsorption capacity of 1.88 mmol g^{-1} for CO₂.²⁸⁸ The metal oxide impregnation on AC significantly impacts both the structural characteristics and $CO₂$ adsorption capacity, depending on the metal loading. A low metal loading may not enhance the adsorption capacity, while an excessively high metal loading can cause pore clogging, which reduces the surface area and pore volume, thereby diminishing the adsorption performance.

Fig. 12 Heteroatom-doped carbons for $CO₂$ capture and $CO₂$ adsorption performance of wood ash (WA-1) activated with various polyamines, showing (a) CO₂ adsorption breakthrough profiles, and (b) adsorption capacity along with amine efficiency. Regeneration performance of WA-1 activated with different polyamines: (c) CO₂ desorption curves, and (d) adsorbent regeneration potential and conversion efficiency. Reprinted with the permission.^{265,286}

Recent studies have focused on enhancing $CO₂$ capture using carbon composites derived from walnut shells and doped with metals such as Mg and Mg–Al. Although metal loading led to reduced specific surface areas due to pore clogging, the $CO₂$ capture performance of these composites improved significantly.²⁸⁹ Notably, the composites with 10% Mg and 5% Al achieved a CO₂ uptake of 4.5 mmol g^{-1} . Scanning electron microscopy (SEM) and energy-dispersive X-ray spectroscopy (EDX) confirmed the effective distribution of Mg and Al on the carbon surface, while Fourier-transform infrared spectroscopy (FTIR) and X-ray diffraction (XRD) revealed weakened metal oxide peaks after $CO₂$ adsorption, indicating enhanced chemisorption. This improved performance is attributed to the combined effects of physical and chemical adsorption mechanisms in the metal-doped carbon materials.²⁹⁰

In parallel, biochar modified with $AICI_3$ demonstrated superior $CO₂$ uptake compared to unmodified biochar. This was due to the primary role of surface adsorption facilitated by the

metallic oxides present. Additionally, activated porous carbons derived from pine cones and treated with KOH, incorporating nitrogen and metals, showed high $CO₂$ adsorption capacities.²⁹¹ For instance, the samples prepared at 700 $\mathrm{^{\circ}C}$ with a KOH ratio of two, containing 0.5% nitrogen and calcium, achieved a $CO₂$ uptake of 4.7 mmol g^{-1} .²⁹² These materials also exhibited an excellent performance in the oxygen reduction reaction (ORR), suggesting their potential in various applications beyond $CO₂$ capture. Table 7 provides a comparative overview of various types of sorbents, highlighting their physical and chemical properties together with their $CO₂$ adsorption capacities.

Furthermore, the incorporation of metal oxides such as MgO, BaO, and $Fe₂O₃$ into activated carbon has been shown to enhance its $CO₂$ adsorption capacity. Basic oxides, in particular, are more effective than acidic or neutral oxides. For example, MgO significantly improved the $CO₂$ capture at higher temperatures, and BaO provided the greatest enhancement due to its high electronegativity.²⁹⁴ These findings underscore the

Fig. 13 Mechanism of impregnating activated carbon with metal oxides.

potential of metal-functionalized activated carbons in optimizing $CO₂$ capture and other environmental applications.

5.1.3 Processes involved in $CO₂$ capture. Here, the potential mechanisms for $CO₂$ uptake by carbon materials are depicted, distinguishing between physical adsorption and chemical adsorption based on the surface interactions between CO2 molecules and carbon materials (Fig. 14).269,295 Physical adsorption primarily relies on van der Waals forces and pore filling, with the adsorption temperature exerting a significant influence. In contrast, chemical adsorption involves the formation of chemical bonds between $CO₂$ and the adsorbents, predominantly facilitated by Lewis acid–Lewis base interactions and hydrogen bonding. Micropores, oxygen functional groups, N functional groups, and aromaticity emerge as critical factors influencing $CO₂$ capture.

Table 7. Overview of sorbent types, properties, and $CO₂$ adsorption capacities

Physisorption, a reversible process driven by van der Waals forces and electrostatic forces, is primarily governed by the textual properties of carbon materials, including pore structure and surface area. Although the large specific surface area and porosity of carbon materials enhance $CO₂$ capture, their pore size is more crucial than specific surface area and total pore volume for CO₂ adsorption. Micropores significantly contribute to $CO₂$ adsorption, particularly at low adsorption pressures. Notably, studies have identified narrow micropores with sizes in the range of 0.5 to 0.7 nm as critical for enhancing the $CO₂$ adsorption capacity.²⁹⁵ Under higher adsorption pressures, adsorption occurs through surface coverage, where the specific surface area plays a pivotal role. Adsorption temperature also influences physical adsorption, with higher temperatures promoting $CO₂$ diffusion and weakening $CO₂$ –carbon material interactions. Chemisorption, an irreversible chemical reaction process, involves the formation of new chemical bonds between adsorbents and adsorbates, facilitated by surface functional groups and alkali ions. Hydrogen bonds and Lewis acid–base interactions play crucial roles in chemisorption. Studies have highlighted the significance of hydroxyl groups in enhancing $CO₂$ capture, given that $CO₂$ molecules can form hydrogen bonds with $-OH$, promoting $CO₂$ adsorption. Nitrogencontaining polymers have been employed to modify carbon materials, enhancing their alkalinity and electron density to strengthen the Lewis acid–base interaction. Moreover, the covalent coordination bonds formed between $CO₂$ and the adsorbent surface contribute to chemisorption. Researchers have developed novel N, P co-doped porous carbon materials with high specific surface areas and rich mesopores, micro pores, and N, P, resulting in enhanced $CO₂$ uptake capacity.²⁹⁶ The presence of N groups in porous carbon derived from lotus stalks has been linked to excellent $CO₂$ adsorption capacity, with the N groups playing a dominant role in $CO₂$ adsorption at higher temperatures and lower pressures. However, the excessive loading of modifying agents such as melamine can clog the pores and reduce the specific surface area of carbon materials, thus impacting their $CO₂$ capture performance. Thus, achieving a balance between loading amount and porosity changes is essential to realize the optimal $CO₂$ capture efficiency. Studies have demonstrated an enhancement in surface area through heat treatment after melamine impregnation, highlighting the Provider and the spin interest article. Common article. Common article is the spin interest are and the spin interest are and the spin interest are a creative Commons are the spin interest are the spin interest at the spin

Fig. 14 Processes involved in $CO₂$ capture by carbon materials. Reprinted with permission.^{269,295}

importance of carefully managing the loading amounts to maximize the capture performance. These findings underscore the multifaceted mechanisms underlying $CO₂$ capture by carbon materials and emphasize the need for tailored material design to optimize the efficiency in various applications.

5.1.4 Traditional and alternative $CO₂$ adsorbents. Activated carbons, zeolites, and molecular sieves are among the traditional $CO₂$ adsorbents that have received extensive scrutiny for $CO₂$ capture applications due to their well-established traits and commercial accessibility. These materials frequently demonstrate impressive adsorption capacities and selectivity, but they may face challenges such as limited thermal stability and difficult regeneration. Table 8 provides an overview of the advantages and disadvantages associated with both

conventional and non-conventional $CO₂$ adsorbents. Identifying the optimal $CO₂$ adsorbent is intricate due to the complex interplay among the synthesis parameters for activated carbons and adsorption conditions. Discrepancies in research data hinder the unbiased assessment of the potential of waste products as $CO₂$ adsorbents given that their effectiveness cannot be solely gauged by comparing the $CO₂$ uptake under standardized conditions. The characteristics of activated carbons are heavily influenced by the synthesis process, which varies across different studies. Furthermore, many investigations fail to present a comprehensive view of the adsorption system, often overlooking potential competitive adsorbates such as CO, N_2 , and CH₄, while focusing solely on CO₂ removal efficiency. Besides cost considerations, factors such as operational intricacy, regeneration feasibility, source availability, and environmental impact must be carefully weighed when selecting appropriate adsorbents.

5.1.5 Mechanism of $CO₂$ adsorption on acrylic carbon. The application of acrylic carbon for $CO₂$ adsorption is increasingly recognized due to its promising potential with unique structural properties and customizable surface chemistry. Derived from the polymerization of acrylonitrile or its copolymers, acrylic carbon undergoes carbonization to form a material with a highly tunable porous structure and substantial surface area, both of which are essential for effective $CO₂$ capture. The adsorption process primarily involves physisorption and chemisorption mechanisms. In physisorption, $CO₂$ molecules are adsorbed on the acrylic carbon surface via van der Waals forces, allowing reversible adsorption without altering the chemical structure of $CO₂$. This characteristic is particularly beneficial for cyclic adsorption–desorption processes, given that it enables low-energy regeneration, making it an energyefficient option for repeated use.³¹⁸ The porous structure of materials is instrumental in enhancing $CO₂$ capture by providing numerous adsorption sites and promoting efficient gas diffusion. Moreover, the performance of acrylic carbon can be significantly enhanced by modifying its surface chemistry to introduce functional groups that facilitate chemisorption. Chemisorption involves the formation of strong chemical bonds between $CO₂$ molecules and surface groups such as amine, hydroxyl, and carboxyl, which can be introduced either during the synthesis of the polymer precursor or through postsynthesis modifications. For example, incorporating nitrogencontaining groups via ammonia treatment or co-polymerizing with nitrogen-rich monomers increases the $CO₂$ uptake due to the formation of carbamate species during chemisorption. This dual mechanism approach, leveraging both physisorption and chemisorption, positions acrylic carbon as a highly effective material for $CO₂$ capture applications.³¹⁹

Activated carbon fibers (ACFs) were developed from mechanically recycled acrylic (PAN), cotton (CO), and their blends (PAN/CO) to enhance CO_2 capture.³²⁰ The production process involved stabilization, carbonization, and chemical activation. However, although PAN is effective for ACF production, it is costly and has significant environmental impacts due to its energy-intensive production. Thus, to address these issues, recycled or renewable materials were utilized to reduce

both costs and environmental footprint. PAN fibers were stabilized using oxidation and phosphoric acid treatment, followed by carbonization at a temperature of 800 °C or 900 °C under nitrogen, and activation with various KOH ratios at 700 ° C. The stabilization process preserved the fiber structure and improved the heat resistance.³²¹ Generally, the PAN fibers maintained their integrity after carbonization and activation, although some samples showed rough surfaces due to the incomplete KOH removal.

The $CO₂$ adsorption capacity of ACFs made from recycled bers was notably high, with higher KOH ratios enhancing their surface area and pore volume, which in turn improved their $CO₂$ capture performance. These recycled ACFs showed comparable or superior $CO₂$ adsorption capacities to commercial ACFs, highlighting their potential for sustainable $CO₂$ capture. Additionally, this study evaluated the mass yield and elemental composition of PAN and PAN/cellulose (PAN/cell) nanofibers in producing activated carbon nanofibers (ACnFs). Oxidative stabilization reduced the mass of PAN/cell fibers more than PAN fibers. Carbonization and activation caused significant mass loss in the PAN/cell nanofibers due to their lower carbon content and thermal stability, although the PAN/cell fibers had a higher nitrogen content. The SEM and FTIR analyses confirmed the changes in the fiber diameter and chemical structure. Fig. 15a shows SEM images of biomass fibers before and after grafting with poly(acrylic acid) (PAAc), highlighting the increase in surface roughness.³²² Fig. 15b presents the SEM images of the PAN fibers at various stages, showing changes in surface texture and porosity with treatment. Fig. 15c shows a comparison of the untreated PAN fibers with that carbonized at different temperatures, revealing signicant structural transformations. The PAN-based ACnFs carbonized at 800 °C exhibited higher $CO₂$ adsorption compared to that processed at 900 °C, emphasizing the impact of the production conditions on CO_2 capture efficiency.³²⁴ Recent evaluations of the CO_2 adsorption capacities for various the carbon-based adsorbents at 1 bar and 25 °C highlight their effectiveness in $CO₂$ capture. This study shows that recycled ACFs derived from acrylic carbon show a comparable performance or surpass that of many commercial carbon adsorbents.³²³ Review WextRom Concerning Co. adsarbents. Identi- back comparison and environmental functions are the proposite and all proposite and all proposite and and and the proposite and and the proposite and and the proposite and

> Furthermore, doping acrylic carbon with metal nanoparticles, such as alkali and alkaline earth metals, has shown to boost the $CO₂$ adsorption significantly. Metals such as sodium, calcium, and magnesium interact synergistically with $CO₂$, combining the benefits of both physisorption and chemisorption. Metal-doped acrylic carbon adsorbents exhibit higher $CO₂$ capture capacities and improved selectivity, given that the metal sites provide additional reactive centers for $CO₂$ binding.²⁶⁵ These modifications not only enhance the $CO₂$ capture performance but also improve the stability and recyclability of the adsorbent, making them viable for long-term industrial applications.

> The application of acrylic carbon for $CO₂$ adsorption holds promise across various sectors, including flue gas treatment, air purification, and carbon capture and storage technologies. The versatility in its synthesis and the ability to finely tune its structural and chemical properties make acrylic carbon a highly

Fig. 15 Surface morphology of biomass and polyacrylonitrile (PAN)-based nanofibers for enhanced CO₂ capture: (a) pristine vs. PAAc-grafted biomass, (b) SEM comparison of untreated PAN, C800, and PANC/cell composites, and (c) PAN nanofiber morphology under varying treatments and magnifications. Reproduced from ref. 322 and 323.

adaptable material for $CO₂$ capture. Recent advancements focus on optimizing the synthesis routes to improve its pore structure and surface functionality, thereby maximizing its adsorption efficiency under different operational conditions.³²⁵ Continued research is essential to fully understand the adsorption mechanisms at the molecular level, which will aid in designing more efficient and cost-effective acrylic carbon-based adsorbents. Moreover, scaling up the production of these materials, while maintaining their performance and stability is crucial for practical applications. As global efforts to mitigate climate change intensify, the development of advanced materials such as acrylic carbon for $CO₂$ capture is becoming increasingly critical. The integration of these materials into existing $CO₂$ capture frameworks can significantly enhance their efficiency and contribute to reducing atmospheric $CO₂$ levels.³²⁶

5.2 Prospects for commercialization

The potential for commercializing a product or technology relies on diverse factors such as market demand, technological feasibility, scalability, regulatory compliance, and economic viability. In the domain of $CO₂$ adsorbents, the prospects for their commercialization depend on their capacity to meet or exceed industry-defined performance standards, while remaining cost-effective and environmentally sustainable. Additionally, considerations such as the availability of raw materials, manufacturing simplicity, and compatibility with existing

infrastructure play pivotal roles in determining the feasibility of their large-scale production and deployment. With various synthesis strategies for biomass-derived porous carbon materials, the transition from laboratory-scale or pilot plant development to industrial-scale production capacity becomes crucial. Achieving this transition requires integrating chemistry and chemical engineering technologies. Nonetheless, several challenges need to be addressed. Firstly, scaling up synthesis protocols from gram to kilo batch sizes poses a signicant hurdle, compounded by uncertainties in biomass supply and associated logistical costs. Establishing local biomass resources, enhancing annual yields, and devising new logistics routes can offer sustainable supply chains from field to processing plant. Subsequently, for improved qualities, micro and mesoporous carbon powder materials must be transformed into granules, pellets, beads, or extrudates with or without binders. Granulation techniques sometimes result in the loss of specific surface area, despite the fact that granular materials offer higher mechanical strength, decreased attrition, and lower resistance in the adsorbent bed. However, studies suggest that incorporating biomass-based pore-forming materials can enhance the sorption properties.³²⁷ Once the characteristics of materials are understood, finalizing the design of $CO₂$ capture plants becomes essential. Strategies may involve temperature swing adsorption (TSA), pressure swing adsorption (PSA), vacuum swing adsorption (VSA), or combinations thereof.

These next-generation carbon materials outperform traditional sorbents such as zeolites due to their inherent hydrophobic properties, low isosteric heat of adsorption, high $CO₂$ adsorption capacities, and long-term stability across multiple cycles. Fig. 16 schematically illustrates the transition from porous carbon powder to plant-level deployment. Further material and design considerations should be made based on the specific $CO₂$ removal application, whether for pre- or post-combustion scenarios requiring high $CO₂$ concentration removal or for confined environments such as cockpits and underground spaces with lower $CO₂$ concentrations. In these cases, highperformance materials such as carbon with strong interactions at low surface coverage, requiring minimal heat for regeneration, are essential. Additionally, anticipating the adsorption of $CO₂$ under humid conditions underscores the need for porous carbons capable of efficient operation in these circumstances.

5.3 Cost assessment framework

Cost assessment frameworks for producing activated porous carbons, especially for $CO₂$ capture, play a pivotal role in determining the economic feasibility of transitioning from laboratory-scale experiments to full-scale industrial production. Producing activated carbons from biomass encompasses various cost components, which can be broadly categorized into fixed and variable costs. Typically, fixed costs involve capital investments in infrastructure, equipment, and facilities necessary for the initial setup of the production plant. Alternatively, variable costs include raw materials, energy consumption, labor, maintenance, and other operational expenses. The interplay among these costs significantly impacts the overall

financial outlook of producing activated carbon on an industrial scale.

Establishing a production plant to meet industrial demand involves substantial capital investments, which are proportional to the plant size and the scale of equipment required for different production stages, from raw material preparation to the final collection of activated carbon. For instance, producing around 450 tonnes of physically activated carbon from biomass annually necessitates an initial capital investment of approximately \$2.12 million, coupled with annual operational costs of about \$1.22 million. In contrast, using a chemical activation method, such as with H_3PO_4 , for similar production volumes requires a total cost investment of approximately \$4.25 million, with market prices ranging from \$2.7 kg $^{-1}$ to \$2.9 kg $^{-1}$. These figures underscore the significant financial commitment required for large-scale production, emphasizing the importance of precise cost assessment frameworks.³²⁸ The cost dynamics of producing activated carbon from biomass are complex and multifaceted, as shown by recent studies. Skoczko et al. estimated that physical activation processes generally cost between \$1.5 and \$2.0 per kilogram, with variations influenced by factors such as the scale of operation and the type of biomass feedstock used.³²⁹ Chemical activation, particularly with H_3PO_4 , incurs significantly higher costs, with Raja et al. reporting costs ranging from \$3.0 to \$3.5 per kilogram.³³⁰ This difference underscores the importance of selecting the appropriate activation method, given that it plays a crucial role in the overall production costs.

The feedstock choice is another critical factor affecting the cost structure. Agricultural residues and forestry by-products are often more affordable options, but chemical activation methods tend to drive up costs due to the expensive reagents required. For instance, Ng et al. found that producing activated carbon from pecan shells costs between \$2.60 and \$2.76 per kilogram, depending on the activation method.³³¹ Similarly, Lima et al. achieved a competitive cost of \$1.38 per kilogram for activated carbon derived from organic waste using physical activation, assuming no raw material expenses.³³² In contrast, Stavropoulos et al. provided a comparative analysis of the production costs for activated carbon from various materials, revealing significantly higher costs for non-biomass sources such as used tires (\$10.88 kg^{-1}), wood (\$6.09 kg^{-1}), and lignite $($5.13 \text{ kg}^{-1}$).$ ³³³ Serafin *et al.* further illustrated the variability in production costs by estimating that producing 50 000 kg of activated carbon from fern leaves will cost approximately \$2.72 per kilogram based on an annual production cost of \$135.690.^{334,335} Additionally, energy consumption remains a key cost component due to the energy-intensive nature of the carbonization process, with market prices for activated carbon ranging from \$2.7 to \$3.3 per kilogram, depending on its quality and application.³³⁶

According to the comparison of these market prices with the estimated production costs, biomass-derived activated carbons show a competitive edge, particularly when considering the environmental benefits of using renewable resources and minimizing waste. Looking at the commercial landscape, the global production of activated carbon reached USD 7.2 billion in 2023 and is projected to grow by 12.42%, reaching USD 14.50 billion by 2031. This surge in demand for activated carbon can be attributed to various factors, including population growth, extensive consumption across industries, and stringent environmental regulations governing air and water purification systems. Currently, the commercial-scale production of activated carbon predominantly relies on coconut shell and coalbased precursors. However, these raw materials alone may not be sufficient to meet the escalating demand across diverse sectors. Therefore, exploring alternative precursors, particularly biomass, can offer a cost-effective solution to meet the projected demands. Globally, approximately 105 billion metric tonnes of biomass, sourced from both land and oceans, is available annually.³³⁷

Refining the cost assessment frameworks involves a comprehensive analysis of the production costs, market trends, and operational efficiencies. As technological advancements and process optimizations continue to lower production costs, the economic feasibility of large-scale biomass-based activated carbon production is improving, making it an increasingly attractive option for applications such as $CO₂$ capture. Ongoing research and development will be crucial in further reducing costs and enhancing the viability of this sustainable production pathway.

6. Research gap

Despite the considerable volume of research conducted on the potential of waste products as precursors for AC, a signicant research gap exists concerning the cost-effectiveness and scalability of the process. Many studies lack detailed procedures

during the synthesis stage, resulting in inconsistent characteristics and adsorption properties of the manufactured product. To commercialize this type of activated carbon, there is a need for a thorough investigation into the continuity of waste supply for precursors, given that abundant resources are essential. Although the average biochar yield is around 20 wt% compared to the raw materials, detailed statistics on precursor availability are necessary to ensure long-term sustainability. Additionally, more research effort is required to explore regeneration methods for activated carbon, given that current technologies often necessitate high heat, which may not be feasible in the long run. Furthermore, there is a lack of studies on the synthesis of hybrid activated carbons incorporating heteroatoms and metal oxides. These hybrid materials have the potential to exhibit enhanced adsorption properties and stability. Moreover, there is a dearth of research focusing on the detailed understanding of $CO₂$ adsorption at the quantum level. These insights are crucial for predicting the behavior and performance of activated carbons under various operating conditions accurately. Comprehensive studies in this area will significantly contribute to advancing the understanding of the $CO₂$ capture mechanisms and optimizing the performance of activated carbon-based adsorbents for practical applications. **PEX Advances**
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7. Summary and prospects for future investigations

This review article provided a comprehensive examination of the synthesis and modification techniques for biomass-derived carbon materials intended for $CO₂$ capture. It delved into the fundamental methodologies for converting biomass into carbon materials and the critical factors influencing their properties. The importance of selecting appropriate biomass sources and processing parameters to tailor the structure and chemical composition of carbon materials was emphasized. Various modification approaches, both physical and chemical, were outlined, highlighting their role in modifying pore structures or introducing functional groups or metals to enhance the $CO₂$ capture efficiency. However, despite the advancements, challenges and research opportunities persist, particularly in understanding the relationship between pore structure and factors such as biomass characteristics, pyrolytic conditions, and activation methods. Optimizing the pore structure of activated carbons, especially that activated with agents such as KOH and $ZnCl₂$, is crucial. Additionally, synthesizing hybrid activated carbons using multiple activating agents holds promise for improving the $CO₂$ capture performance through hybrid porous features.

Moreover, although renewable biomass serves as a costeffective precursor for carbon capture materials, pristine carbon materials have inherent limitations in terms of pore structure and surface chemistry, constraining their adsorption capacities. Although various modification methods have emerged to address these limitations, some processes are complex and costly, hindering their scalability for industrial applications. Template carbonization offers a viable alternative,

providing control of the material pore diameter and structure without additional modification treatments. However, much of the research on $CO₂$ adsorption by carbon materials remains confined to laboratory settings, overlooking real-world challenges such as moisture and corrosion. Future research should focus on enhancing the mechanical properties of carbon materials to withstand harsh environments and assessing their suitability for industrial applications. Furthermore, it is necessary to explore competitive adsorption mechanisms among different gases and simulate the carbon material adsorption performance in complex gas compositions. Additionally, research efforts should extend beyond $CO₂$ capture to encompass $CO₂$ conversion, leveraging the catalytic properties of biomass-based carbon materials. The in situ conversion of adsorbed $CO₂$ presents an intriguing avenue for sustainable $CO₂$ recovery. However, challenges related to regenerating $CO₂$ saturated carbon materials and mitigating the effects of contaminants require further investigation. Understanding the impact of contaminants on carbon materials and developing strategies to enhance their resistance can enhance their longevity and performance in $CO₂$ capture applications. **Review Excess Article is the continuum access Article is licensed on 18 2024.** The set is determined the creation and continuous article is licensed under the set is license and continuous article is licensed under a crea

Transitioning from laboratory-scale studies to pilot-scale investigations is crucial to evaluate material performance under realistic industrial conditions. Furthermore, gaining deeper insights into the underlying mechanisms for the formation of activated porous carbons from biomass and their regeneration processes can facilitate the design of advanced carbon materials for various applications. Addressing these research gaps and challenges is essential to harness the full potential of biomass-derived activated porous carbons for $CO₂$ capture, water decontamination, and energy storage, thereby advancing carbon capture technologies and promoting environmental sustainability.

Data availability

The data supporting the findings of this review can be obtained from the corresponding author upon request. Due to privacy concerns and other restrictions, the data are not publicly accessible.

Author contributions

Shreyase Kundu, Tasmina Khandaker & Md Al-Amin Mia Anik: writing - original draft, review and editing; Md. Kamrul Hasan, Palash Kumar Dhar, Sagar Kumar Dutta & M. Abdul Latif: proof reading and editing; Muhammad Sarwar Hossain: formal analysis, methodology, resources, validation, writing and editing, supervision and project administration.

Conflicts of interest

The authors declare that they have no known financial or personal conflicts of interest that could have influenced the work reported in this paper.

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