

Review on Biochar Upgrading Methods for Its Application in Thermochemical Conversion Processes and Critical Materials Recovery

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Review

Review on Biochar Upgrading Methods for Its Application in Thermochemical Conversion Processes and Critical Materials Recovery

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Abstract

With the rapid increase in solid waste generated worldwide, sustainable approaches for the recovery of resources considering environmental protection are required. As one of the emerging strategies in recent years, biochar has shown great potential due to its high carbon stabilization, adjustable porosity and tunability. This review focuses on the critical assessment of the available technologies for biochar upgrading, with a specific objective of biochar physicochemical functionality improvement and critical materials recovery in line with circular economy targets. We systematically review physicochemical activation methodologies, functionalizations and leaching approaches, accounting for their effects on surface area, porosity and functional group chemistry. Particular attention is paid to the dual functionality of upgraded biochar (i) as a catalyst support for thermochemical processes and (ii) as a medium for the recycling of essential nutrients (e.g., phosphorus, potassium, magnesium, calcium). It is evidenced that customized activation can further improve its adsorption and catalytic efficiency as well as promote near-total nutrition extraction. This review positions advanced biochar as an enabling multipurpose technology across sustainable material production, nutrient cycling and waste valorization in the circular bioeconomy.

Keywords: biochar; upgrading methods; materials recovery; chemical leaching



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1. Introduction

The exponential increase in waste production in all sectors has become a major global burden resulting in serious ecological pollution and threats to environmental and public health as it attracts disease organisms [1,2]. Conventional biowaste management options, including landfilling, are less desirable because of their environmental, social, and economic impacts, such as the taking up of valuable land, production of greenhouse gases (GHGs) like methane, and the potential for soil and groundwater contamination. Worldwide production of solid waste is expected to rise from 2.01 billion tons per year to a staggering 3.40 billion tons by 2050, hence the necessity for creative and sustainable waste management solutions [3,4]. In 2023, the global solid waste sector generated an estimated 173.2 million tonnes of CO₂e with around 70% released from open dumping, a practice that is still in use in many low- and middle-income countries. These projections, illustrated in Figure 1, suggest that emissions under a business-as-usual scenario could reach 203.4 Mt

CO₂eq in 2030 and 289.5 Mt CO₂eq in 2050 if there are no changes to existing practices. This underscores the need to develop more sustainable and systemic solutions. The circular economy (CE) provides an alternative path by stressing the prevention of waste, recovery of materials, and closed-loop resource streams flows [5]. CE deployment in the waste sector could bring us an order of magnitude closer to reaching net-zero emissions by 2050. Switching systems from linear to circular waste is not only key to reaching GHG reduction targets, it is also an important improvement in resource efficiency, preventing environmentally degrading discharges of hazardous and other waste streams, with potential benefits in support of the circular and bioeconomy [6].

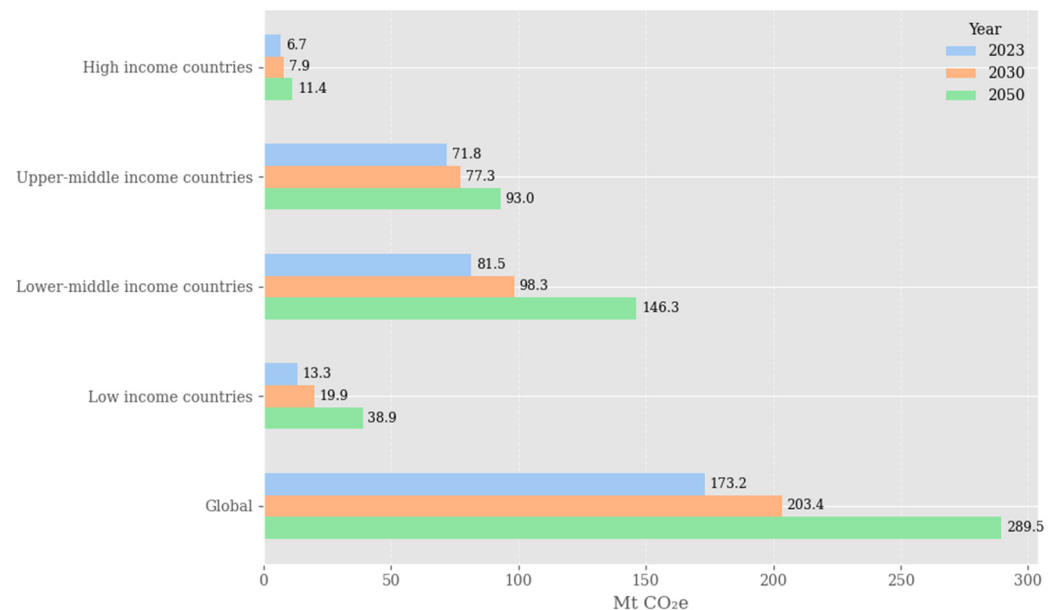


Figure 1. Estimated global GHG emissions from solid waste management across income groups for the years 2023, 2030, and 2050. Elaborated from: [7].

One promising approach is converting biowaste into biochar through carbonization. This method not only reduces waste, but also recovers raw materials, making it a smart step toward circular- and resource-efficient systems [8]. Resulting biochar can be modified to have high surface area, tunable porosity and rich surface chemistry, which justifies its use in applications including carbon sequestration, CO₂ capture, heavy metal immobilization and removal of organic micropollutants from wastewater [9]. Crucially, during biochar upgrading operations it is possible to recover materials, such as phosphorus, potassium, magnesium and calcium, which are considered critical raw materials by the European Union because of their scarcity and increasing demand [10].

Biochar is typically produced through slow pyrolysis of lignocellulosic biomass at temperatures ranging from 400 to 750 °C and in oxygen-limited environments. The derived biochar is a carbon rich solid which contains variable amounts of hydrogen, oxygen, nitrogen, phosphorus, and ash minerals. The processing temperature, residence time, reactor type, and pre-treatment steps can affect the physicochemical characteristics of the produced biochar. These process variables essentially govern the carbonization degree and the development of surface features that characterize biochar's functioning in diverse applications. Steps like drying, grinding, and activation can further enhance biochar's performance and functionality [11]. The ash content in biochar is made up of minerals like phosphorus, calcium, aluminum, iron, potassium, and silicon which vary depending on the feedstock sources [12]. Its potential for long-term carbon storage is due to the high proportion (often over 87%) of the stable and resilient carbon it contains [13].

Biochar also exhibits promising performance as a multifaceted catalyst in many reactions, including, among others, methane cracking, biodiesel production, and syngas upgrading [14,15]. Because of its properties, like high porosity and carbon content, and its potentially lower costs, using biochar as an alternative to traditional solid metal-based catalysts shows an interesting potential [16,17]. Two key physicochemical features, including surface functional groups and the dispersion of metal nanoparticles on biochar surfaces, play a crucial role in enhancing its catalytic performance. Moreover, properties like Specific Surface Area (SSA), pores, and its hydrophobic nature also impact the effectiveness in its application as a catalyst [18].

On the basis of the well-established multifunctionality of biochar in the environmental, agricultural, and industrial contexts, the combined thrust of biochar application via upgrading methods with the aim of material recovery as fertilizers and applications of activated biochar as catalyst in methane cracking, biodiesel production, syngas upgrading and even cleaning up environmental contaminants constitutes a derivative approach that simultaneously addresses resource recovery and waste valorization. This represents a win-win strategy which meets the needs of circular bioeconomy approaches. Although a vast amount of literature is available related to the production, properties, and applications of biochar, the original contribution of this work is formed through a deep look at the single stage and integrated chemical–physical activations, the subsequent development of pathways toward upgrading the properties of biochar, catalytic and adsorptive wise (enhanced surface area, porosity and designed functional groups) and also the recovery of valuable nutrients (e.g., phosphorus, potassium, magnesium) from waste derived feedstocks. This dual application serves to contextualize the biochar process in the context of the circular bioeconomy of waste valorization and resource provision. Furthermore, this paper integrates developments, including environmental trade-offs and establishing biochar as a versatile platform to connect sustainable material production with nutrient recovery and circular economy ambitions.

2. Biochar Production Methods

There are several main processes (see Figure 2) for producing char, that differ in feedstock flexibility and operational parameters as well as in the resulting quality and yield of the products. Among the prevalent techniques, slow and intermediate pyrolysis stand out, with slow pyrolysis proving to be the most efficient approach for generating substantial quality of biochar considering the carbon retention and longest residence time [19]. It is worth mentioning two additional pre-treatment methods for transforming carbon from biomass: torrefaction and hydrothermal carbonization (HTC), whose solid products have different structures in comparison to biochar.

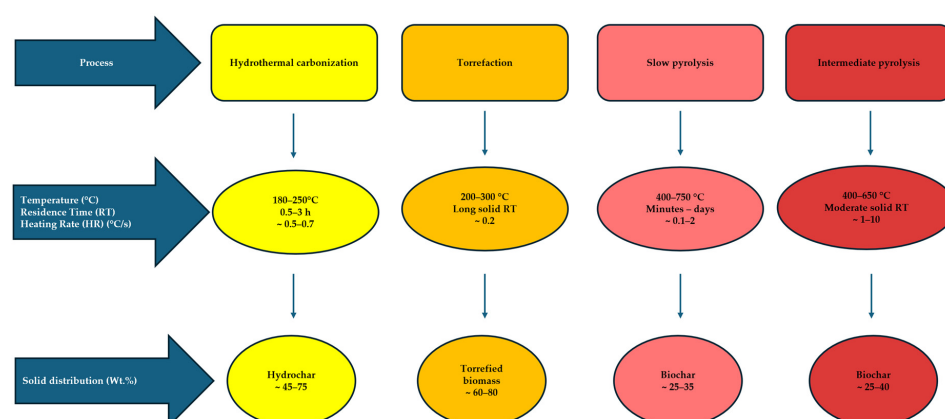


Figure 2. Operating parameters and typical solid distribution in different processes [20–22].

2.1. Hydrothermal Carbonization (HTC)

Hydrothermal carbonization is a thermochemical process that transforms wet biomass (75–90% water content, without requiring a drying pre-treatment step [23]) into a carbon-rich product called hydrochar. In this process, wet biomass is heated to 180–280 °C in water under pressure (10–40 bar). The reaction produces three main outputs: hydrochar (a solid fuel material), process water (rich in dissolved organics), and a small amount of gas (mainly nitrogen and carbon dioxide, comprising approximately 1–2% of the input), the yield of which can vary based on the feed composition [24]. Hydrochar production generally requires less energy consumption than biochar production. It holds potential as a coal substitute due to its comparatively high Higher Heating Value (HHV) [25]. In comparison to biochar, hydrochar has lower carbon, ash content and also a smaller specific surface area and pore volume.

2.2. Torrefaction

Torrefaction is a moderate temperature pre-treatment process in which biomass is heated slowly to about 200–300 °C for a short duration prior to pyrolysis. This process produces a light, porous solid rich in carbon with reduced water content, which in terms of energy density and grindability, is more conducive to handling, storage and transportation of the material [26]. The total carbon yield is influenced by the treatment temperature, residence time, feedstock and furnace atmosphere. For example, lignin from beech starts to decompose at approximately 200 °C, most hemicellulose at about 230 °C and cellulose at temperatures well above 270 °C [27]. Under pilot-scale operations with hardwood and switchgrass pellets, solid yields greater than 77 wt% have been reported [28]. Limited surface area, lower porosity and unstability of torrefied biomass make it useful as a fuel, but not for environmental and catalytic processes [29].

2.3. Slow Pyrolysis

Slow pyrolysis is recognized as a suitable method to convert biomass into a stable solid product enriched with carbon and valuable elements (phosphorus, etc.), depending on the feedstock [30]. This thermochemical process is well known, and it offers a sustainable way to produce pyrochar from biomass. In slow pyrolysis, approximately 50% of the feedstock carbon content can be permanently converted into a stable carbon structure [17].

Together with biochar, other condensable (also known as bio-oil) and incondensable fractions (pyrogases) are produced. These fractions can be either valorized and/or used to supply the process energy demand. In the context of the slow pyrolysis process with energy recovery (Figure 3), the biochar yield and quality are highly dependent on the process parameters. Lower temperatures, slower heating rates, longer residence times and larger particle sizes tend to increase char yield because they promote secondary reactions that fix carbon in the solid phase [31]. The composition of feedstock is also important, since biomass with a high lignin content generally generates more biochar due to its higher thermal stability when compared to cellulose and hemicellulose [32]. While the optimal operating conditions depend on feedstock properties, slow pyrolysis generally follows a tendency where more moderate pyrolysis conditions favor carbon recovery in biochar; but higher temperatures and faster heating rates lead to a production distribution shift to bio-oil and pyrogas [33].

In general, since intermediate and fast pyrolysis processes use higher heating rates, the yield of biochar is typically lower compared to the slow pyrolysis [34]. According to the reported studies, at a temperature of 400 °C, by increasing the heating rate from 5 °C/min to 30 °C/min, the biochar yield decreases from 34.2 to 29.7 percent [35]. These considerations

allow us to remark that the process conditions and the feedstock quality significantly affect both the final product yield as well as its chemical–physical characteristics.

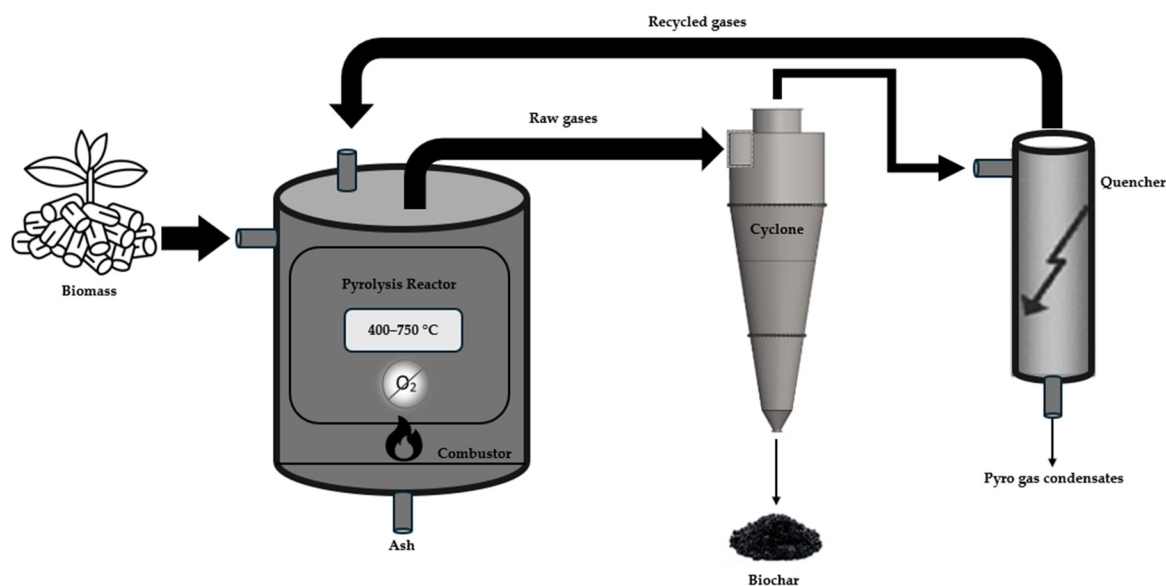


Figure 3. Basic conceptual scheme of slow pyrolysis for biochar production [36].

The biochar produced by slow pyrolysis (pyrochar) with a low heating rate and high residence time is generally more structurally stable, has a higher carbon content and greater porosity suitable for high temperature conversion processes as a medium, and can even help with the cleanup of pollutants. To put it more simply, pyrochar is more stable in the long term as a conditioner and amendment for soil and environmental use [37]. The production of biochar involves the breakdown of organic materials and the removal of volatile compounds, resulting in a stable matrix with enhanced aromatic carbon structures. These structural features are crucial for biochar’s catalytic functions, as they contribute to its large surface area, increased porosity and abundant active sites, all of which facilitate chemical interactions and make biochar an effective catalyst [38].

Consequently, torrefied biomass obtained at low temperatures has its fuel properties significantly upgraded, but it still does not have the aromaticity and surface area required for more advanced applications [39]. HTC hydrochar also contains high volatiles and higher H/C ratios (>1.4), and it is more suitable as a soil additive and nutrient reuse than catalytic applications [40]. On the contrary, pyrolysis biochar has lower H/C and O/C ratios (<0.7), high aromaticity and structural stability, as well as well-tuned porosity. Moreover, the inherent alkaline nature and rich mineral element content of pyrolysis-derived biochar with the abundant surface functional groups (phenolic and carboxylic acids) also contribute to its catalytic properties [41]. The combined properties described above render slow pyrolysis biochar a potential candidate as multifunctional material for adsorption, environmental applications, and catalyst support [14].

3. Biochar as Catalyst Support

Biochar can stabilize and make nanoparticles uniformly spread on its surface which makes it act as a very effective catalyst as it has active sites for different reactions. The mesoporous structure allows for good dispersion of metals such as manganese, copper, cobalt and iron and immobilization in the pores, preventing leaching into water systems [42]. Its unique features include heterogeneity (easy isolation from reaction mixtures), bifunctionality (promoting both transesterification and esterification reactions), recyclability, porosity, and non-graphitizability, which makes it resistant to the crystallization process at high tem-

peratures [43]. Compared to other catalysts, biochar is low-cost, degradable and recyclable, which as a result is useful in a wide range of applications in energy and environment, such as biodiesel production, methane and tar cracking, syngas and pollutant removal [44]. For example, the native inorganic elements, including Fe and K, in biochar increase tar cracking activity [45] or having biochar support as an eco-friendly and efficient catalyst for methane cracking to produce hydrogen [46]. However, raw biochar often suffers a disadvantage of low porosity, low surface area and poor catalytic performance, since biochar has a plenty of functional surface groups (such as -OH, C=O, etc.) and needs to be modified to improve its functions (including feedstock activation, synthesis control and surface functionalization) [47]. Surface activations and modifications for biochar exhibit marked enhancement in porosity and efficiency, making it widely used in the biorefinery and energy production [48].

3.1. Defined Key Properties of Biochar as Catalyst

In recent years, biochar has been considered as a catalyst for thermochemical processes thanks to a few key qualities: it can handle high heat, resists chemical poisons such as sulfur and contains helpful metals such as iron. These metals serve as active sites, making the reaction more efficient and stable [49]. To boost its performance, the structure of biochar should be optimized. A higher surface area and more pore volume help trap carbon better during the reaction. Thermal treatments can also create intentional flaws and structural defects that act as energetic spots where methane's strong C-H bonds can break more easily [50]. While temperature and gas composition matter, it is ultimately the catalyst's design that often determines how much methane actually converts [51].

In the summary of the discussion of biochar upgrading according to the research, the following parameters should be taken into consideration.

3.1.1. Surface Functionality

Biochar inherently consists of oxygenated functional groups including hydroxyl (-OH), carboxyl (-COOH) and carbonyl (-C=O) moieties which markedly improve its catalytic activity [52]. The improved performance is largely due to the presence of active surface sites that promote both the adsorption and redox processes, positioning biochar as a material with considerable potential across numerous applications [53]. Studies have shown the catalytic functions of biochar and focused on the contribution of oxygen-containing groups in pollutant removal and catalytic conversion [54]. Functionalization like sulfonation or metal deposition further contributes to catalytic activity. For example, sulfonated biochar is considered a high-efficiency solid acid catalyst in cellulose hydrolysis [55].

3.1.2. Inorganic Elements

Another important characteristic of biochar is the small presence but significant quantities of inorganic compounds like potassium, sodium, calcium, magnesium and iron. The source feedstock of biochar is an important factor controlling the amount and composition of these minerals. For example, biochar prepared from woody, herbaceous, or aquatic plants generally contains significantly less inorganic substances than that prepared from other sources [56]. To investigate the impact of different types of metallic compounds on the activity of carbon catalysts, two primary upgrading methods can be used [57,58]: the first approach involves removing metals from high-ash carbon materials which contain various metals through acid treatment; the second method introduces different metals onto low-ash carbon-based catalysts to evaluate their effect on catalytic performance.

3.1.3. Particle Size

In catalytic processes, the effect of particle size on mass transfer, as well as reaction rate, is important, because with larger catalyst particles, internal mass transfer resistance is greater, which reduces slower reaction kinetics [59]. It is proved by several studies that smaller catalyst particles give a greater amount of the catalytic surface area, which is good for mass transfer and the process reaction [60,61]. But smaller particles can also result in a greater pressure drop in the reactor, and their application in practice may be restricted [62].

In addition to biochar features, there are also exceptionally stable thermal and mechanical properties of biochar that can be employed well in high-temperature processes. Such inherent stability is believed not only to secure stable performance of biochar under environmentally adverse conditions but also to contribute to the reusability of biochar as a catalyst for multiple cycle [17].

4. Review on Biochar Upgrading Techniques

Biochar quality can be improved using a variety of post-production treatments, such as gas activation, reduction, acid and alkali exposure, oxidations and metal ions treatment [63]. Lately, there has been growing interest in improving biochar's ability to absorb substances by expanding its surface area, increasing its porosity or introducing functional groups [64]. The selection of a particular method relies upon its intended final field of application [65]. Chemical reagents employed in biochar upgrading can induce alterations in their physicochemical traits, consequently boosting their sorption capabilities. Upgrading processes significantly increases biochar effectiveness and broadens its potential applications, especially in the remediation of pesticide-contaminated environments [66] as can be seen in Table 1.

Table 1. Biochar upgrading methods [67,68].

Method	Functionality	Effect
Acid/Alkali	Remove impurities and change specific surface area and pore volumes	Improve adsorption capacity
Oxidation	The surface of biochar is oxidized by coating oxidant	
Reduction	The surface non-polarity of biochar is increased by reducing agent treatment	Increase the specific surface area and oxygen-containing functional groups of biochar
Steam	Promotes the porosity of carbonized materials	
Organic solvent	Organic modifiers can increase the pore structure of biochar	

4.1. Methods for Biochar Upgrading

Upgrading methods aim to improve biochar properties, widening its range of applications [69]. Biochar upgrading methods are generally divided into two main groups: physical and chemical activation. In physical activation, the material is treated with CO₂ gas, steam or a combination of both at temperatures exceeding 700 °C to enhance its porosity and surface area. During this process, highly reactive carbon atoms are eliminated [70,71]. Chemical activation involves using alkali substances such as NaOH, KOH and alkaline earth metal hydroxides (Ca(OH)₂) or acids (such as HCl, H₃PO₄, and HNO₃) either during carbonization or after the biochar has been produced to further modify its properties [72].

4.1.1. Physical Methods

Biochar porosity can be enhanced by physical activations like ball milling and also by using mediums like air, CO₂, or steam at high temperatures. This process generally occurs in two stages: first, the biomass undergoes a thermochemical transformation, and then it is activated to develop a porous structure [73]. In general, increasing the activation temperature leads to a greater release of volatile organic matter, which in turn generates more pores. Pores larger than 50 nm are classified as macropores, those between 2 and 50 nm are termed mesopores, and pores smaller than 2 nm are referred to as micropores. The formation of pores on biochar's surface is a key physical feature that helps dissolve toxic pollutants during adsorption and boosts its catalytic performance [74].

- Thermal-oxidation treatment

After the biochar is heated to the desired temperature (800–900 °C) for 1–2 h, it is exposed to hydrogen, air, or argon gases, and new functional groups are formed on its surface [75]. As illustrated in Figure 4, heating biochar in an air environment significantly changes its surface chemistry, which in turn influences its ability to absorb organic and inorganic compounds. Overall, thermal treatment enhances the adsorption of both polar and nonpolar substances. Thermal treatment can significantly increase the porosity and surface acidic groups of biochar and creates new adsorption sites on biochar for organic/inorganic molecules. The optimum thermal air oxidation temperature is in the range of 400–500 °C. Considering the potential variability related to the feedstock, biochar made from lignin-rich feedstocks is preferable for thermal air oxidation treatment [76–78].

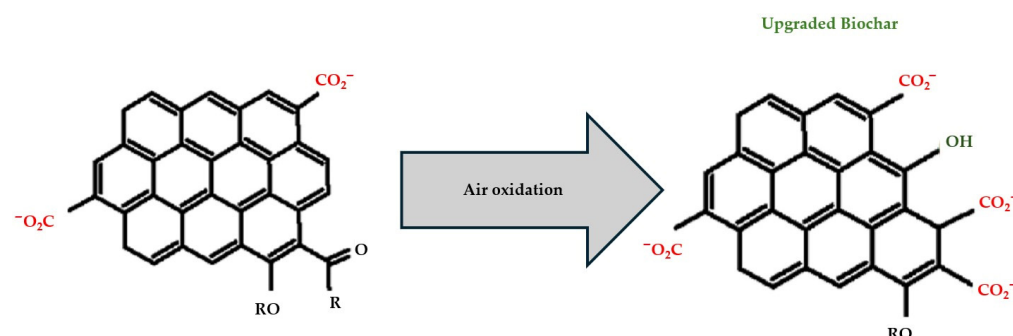


Figure 4. General thermal treatment mechanism of biochar through air oxidation (Colors on structure: red = oxygenated functional groups, green = new functionalities, black = carbon framework). Adapted from [76], *Environmental Functional Materials*, 1 (2022), 187–195. © 2022 The Author. Published by Elsevier B.V. on behalf of KeAi Communications Co., Ltd.

- Steam activation

In this process, the raw material is first heated to a temperature between 300 and 700 °C for 1 to 2 h without any oxygen present. As the second step, the decomposed sample is subjected to a steam flow at a temperature of 800 to 900 °C for 30 min to 3 h, at a rate of 2.2 to 5 mL/min [79]. Steam improves the properties of biochar by removing elements resulting from incomplete combustion during thermal decomposition and oxidizing the carbon surface. The high-temperature steam reacts with the carbon, removing carbon atoms as CO or CO₂, thus developing a more porous structure [80].

- Activation with CO₂

CO₂ can be used as an oxidizing agent, operating between 200 and 900 °C, resulting in the formation of pores and the increase of the existing ones. The CO₂ is usually not very reactive, but when combined with steam at higher temperatures, it becomes active. This enhanced reactivity helps in producing biochar with similar dimensions of pore sizes [70].

The reactivity of CO₂ increases with temperature, facilitating its strong interaction with biochar via the reverse Boudouard reaction. The char (C) is activated by the following reaction: $C + CO_2 \leftrightarrow 2CO$ [81]. During the reaction, carbon dioxide (CO₂) is reduced and attaches to the carbon surface to yield a surface oxide (C(O)) and carbon monoxide (CO). The biochar structure post steam activation and CO₂ purification is shown in Figure 5. Surface oxide (C(O)) is then desorbed, which further refines the pore structure of the char [82]. In Table 2, the reactions involving biochar in different mediums are presented.

Table 2. Physical activation of carbon-based material with various mediums.

Medium	Reaction	Energy Balance
Air/O ₂ [83]	$C + O_2 \rightarrow CO_2$ (1)	Exothermic
Steam/H ₂ O [84]	$H_2O + C \rightarrow CO + H_2$ (2)	Endothermic
	$H_2 + 2C \rightarrow 2C(H)$ (3)	
CO ₂ [85]	$CO_2 + C \rightarrow C(O) + (CO)$ (4)	Endothermic
	$C(O) \rightarrow CO$ (5)	
	$CO_2 + C \rightarrow 2CO$ (6)	

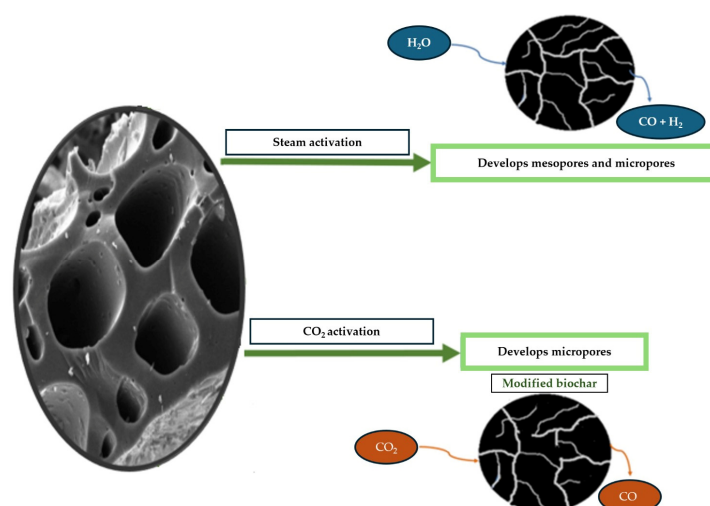


Figure 5. The microporous structure of biochar forms as a result of steam activation and CO₂ treatment. Adapted from [86], Biomass Conversion and Biorefinery, 14 (2020), 5759–5770. © 2020 Springer-Verlag GmbH Germany. Reproduced with permission.

- **Ball milling**

Another physical modification method is ball milling which represents a solvent free, environmental and convenient method. In this manner, the biochar is subjected to mechanical grinding, either by itself or with pulverized solid additives, in order to enhance its physicochemical properties. Ball milling has the ability to produce a smaller mean diameter, increase specific surface area and add functional groups, etc., leading to a high permeability rate of adsorptive species and catalytic efficiency or mixing dispersion in composites [87].

4.1.2. Chemical Methods

The activation process involves using various agents, including bases like KOH and NaOH, acids such as HNO₃ and H₃PO₄, oxidants like H₂O₂ and sulfonating agents (SO₃H), and ZnCl₂. The choice of activation method depends on how the biochar will be used. If it needs to attract negatively charged elements, it is treated with bases to create a positive charge on its surface, improving its ability to bind with the target substance. On the other

hand, acid activation can be used when biochar needs to capture positively charged elements [88]. Additionally, activation helps remove moisture and volatile compounds that could lead to tar formation, ultimately increasing its porosity and carbon yield [89]. Activated biochar is carefully washed to remove any leftover chemicals and expose functional groups that enhance its ability to absorb different substances.

- Alkali activation

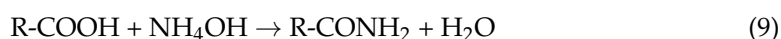
In this process, biochar is treated with reagents like potassium hydroxide (KOH), sodium hydroxide (NaOH) and ammonium hydroxide (NH₄OH) [90].

In a study by Ma [91], Biochar produced from sawdust through pyrolysis in temperatures between 300 and 700 °C was treated with a 2 mol/L potassium hydroxide (KOH) solution for 5 h. During this process, various chemical reactions occurred, as described below:



The addition of more functional groups containing nitrogen to the structure of biochar is made possible by using NH₄OH, which increases the elemental nitrogen in biochar [67].

At this stage, -NH₂ groups were introduced onto the surface of the biochar.



After being treated with NH₄OH, due to the significant removal of ash from the pores, the biochar surface exhibited a smoother post-treatment texture. The application of alkalis led to the thinning of micropores, increasing the biochar's SSA [68].

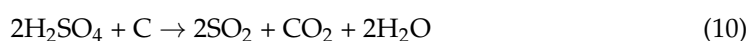
In another study by Haque et al. [92], activated carbon (AC) was obtained through traditional carbonization and KOH activation, while amorphous silica was derived from rice husk using alkali extraction followed by acid precipitation. The ideal temperature for AC production was found to be 700 °C. AC produced via alkali extraction showed greater activity than that obtained through KOH activation.

- Acid activation

Acid treatment can also modify the surface characteristics of biochar, such as its texture, porosity and surface area, and its effects are different with the types and the concentrations of the acids [93]. The application of acidified biochar has been acknowledged for its notable efficacy as catalyst support for thermochemical processes, for enhancing soil quality and for promoting agricultural productivity. Acidification of biochar is known to influence various physicochemical properties, fostering favorable conditions for nutrient retention, positive ion exchange capacity, and soil fertility [94]. Hydrochloric (HCl), sulfuric (H₂SO₄), phosphoric (H₃PO₄) and nitric (HNO₃) acids are typically used in biochar upgrading.

The experimental results [95–97] show that the modified biochar with the highest specific surface area was obtained through acid treatment followed by KOH activation. Similar to basic activation, acid upgrading increases the number of functional groups containing oxygen in biochar [98]. As a result, the surface charge and functional groups are altered, leading to a lower point of zero charge and improved surface purity. Below are several important reactions reported for the acidic upgrading of biochar using different acids.

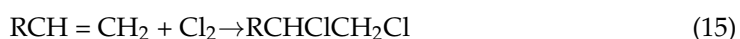
Sulfuric acid reacts with the carbon atoms in biochar through the following chemical process [99]:



Phosphoric acid interacts with the hydroxyl (-OH) groups on the biochar surface, forming ester bonds. This process helps integrate phosphorus atoms into the carbon structure, improving the material's chemical characteristics:



Hydrochloric acid undergoes nucleophilic substitution to the biochar [100]:



During the upgrading process with nitric acid (HNO_3), the aromatic structures within the biochar undergo chemical reactions with the acid. This results in the formation of organic structures containing nitro groups [72]:



As pointed out by Kasozi et al. [101], although the textural qualities of a biochar may not be improved through organic upgrading, its surface chemistry can be impacted quite considerably.

Organic acids, such as acetic acids, also impact the characteristics of biochar. They decrease the porosity of biochar while increasing the formation of new functional groups on the surface, likely due to esterification reactions. As stated by Xu et al. [102], acid treatment does not change the specific surface area (SSA) and the total pore volume (TPV) of biochar because of pore blockage by adsorbed organic acid molecules. This indicates the biochar upgraded with citric acid. Typically, an increase in the atomic O/C ratio signifies an improvement in biochar's surface chemistry. Carboxylic groups are among the identified surface functionalities on biochar modified with organic acids. Sheng et al. [103] demonstrated the upgrading of water hyacinth-derived biochar with citric acid; in the study, the biomass was pyrolyzed at 300 °C for 2 h in a nitrogen atmosphere with a 5 °C/min heating rate, followed by treatment with 0.7 mol/L citric acid for 2 h. The application of citric acid led to a reduction in the specific surface area of biochar, likely due to the dissolution and removal of lignin during the acid treatment.

Acid treatment generally extracts metals such as iron, aluminum, titanium, calcium, sodium and sulfur from carbon-based catalysts and adds oxygen-containing groups, like carboxyl, to the biochar surface [61,104]. While this treatment increases the surface area and pore volume of biochar, enhancing its sorption properties, it also alters its composition and reactivity. Bases like NaOH and KOH can react with sulfur and important minerals found in coal to make compounds like silicate and aluminate. These compounds can then be dissolved by acids, chemicals like sulfuric acid and hydrochloric acid, along with other unused minerals [30].

4.1.3. Inorganic Methods

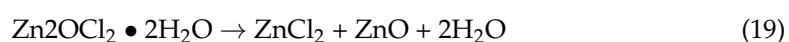
The use of salts, such as chlorides or phosphates, allows for influencing both the physical structure and surface chemistry of biochar. Li et al. [105] noted an alteration in the surface of biochar when it was treated with K_3PO_4 ; in their experiment, the particles were soaked in K_3PO_4 and stirred for 6 h. Afterward, the impregnated biomass was subjected to pyrolysis at 550 °C for 1 h in a nitrogen atmosphere, with a heating rate of 10 °C/min [106].

This pre-treatment procedure resulted in an enrichment of $P_2O_7^{-4}$ on the surface of the biochar formed from the breakdown of HPO_4^{-2} .



The generated components significantly contribute to the adsorption of heavy metals onto biochar. Treating biochar with K_3PO_4 enhanced its textural properties and surface chemistry. This modifier expedited the decomposition of biomass polymers, leading to the formation of numerous oxygen-based groups on the surface of biochar [107].

In an experiment, biochar was treated by soaking it with a solution containing 30% $ZnCl_2$ [98]. During this upgrading process, the following chemical transformations took place within the biochar:



4.1.4. H_2O_2 Upgrading

Biochar can also be enhanced using hydrogen peroxide (H_2O_2), which is often considered a more cost-effective and environmentally friendly alternative to strong acids, bases and salts. In Tan et al. [108], biochar was produced from rape stalk through pyrolysis at 600 °C in a nitrogen atmosphere, with a heating rate of 10 °C/min and a processing time of approximately four hours. To enhance its properties, the biochar was treated with 30% H_2O_2 for 24 h. During this treatment, H_2O_2 reacted with the biochar, resulting in gas formation.



The changes in the biochar properties observed across various studies following different upgrading treatments are presented in Table 3.

Activation techniques result in quite different performances of the biochar. The KOH-activated char had the highest surface area, reached the SSA of 2183.8 m²/g, and developed the sufficient amount of micropores which is suitable for the removal of heavy metal ions and pollutants. Activations by $ZnCl_2$ and H_3PO_4 , which obtained SSA values (>1000 m²/g) and pore volumes as high as 0.6 cm³/g, the resulting hybrid microporous–mesoporous materials are promising for catalysis and tar removal. Physical activation using either CO_2 or O_2 yielded moderate improvements (SSA 200–850 m²/g) with potentially active surfaces useful for syngas upgrading or methane cracking, while N_2 activation was found the least effective (SSA < 122 m²/g) which could be ascribed to both non-reactive atmosphere (inert environment) and few amounts of functional groups being formed. The chemical–physical combined treatments enable synergistic promotion, with a SSA value of up to 1419 m²/g and the PVs of up to 0.83 cm³/g, having potential application in diverse catalysis.

Table 3. Physical–chemical methods used in different sources.

Upgrading Method	Specific Surface Area (SSA) (m ² /g)	Pore Volume (cm ³ /g)	Effects
Chemical activation with KOH at 800 °C on <i>Undaria pinnatifida</i> derived biochar [109]	69.7–1156.2	0.04–0.67	Activated biochar (2.32 nm pore size) with mesoporous structure, more accessible adsorption sites, oxygen-containing functional groups, chemical reactivity
Physical activation with direct CO ₂ at 750 °C on hybrid willow [110]	738.74	0.37	Activated biochar with 2 nm pore size with increased surface area and chemical reactivity
Chemical-Physical activation on rice husk biochar with (KOH, H ₃ PO ₄ , ZnCl ₂) and N ₂ at 800 °C [111]	89.93-KOH: 390.59 H ₃ PO ₄ : 346.3 ZnCl ₂ : 394.5	0.079–0.224 0.189 0.221	Increased surface area provides more active sites for chemical reactions, enhancing catalytic performance, best adsorption capacity for the tar removal
CO ₂ physical activation on the biochar at 900 °C [112]	215–345	0.14–0.19	Enhanced porosity development with the pore size of 2.51 nm
Chemical activation of Pyrolyzed cellulose with 5% H ₃ PO ₄ at 70–80 °C for 2 h [113]	199.5–558	0.02–0.23	The H ₃ PO ₄ upgrading enhanced SSA and pore size
Chemical activation on swine manure derived biochar immersed in H ₃ PO ₄ solution (24 h at 25 °C) [114]	227.6–373.4	0.07–0.14	Enhances surface area due to the generated micropores
Chemical activation on pinecone biochar with ZnCl ₂ : biochar = 2:1 (800 °C) [115]	0.583–1067.9	0.02–0.512	One-step ZnCl ₂ activation increased the specific surface area more efficiently
Physico–chemical activation on rice husk-derived biochar with (CO ₂ , KOH, H ₃ PO ₄) at 600 °C [116]	183.16-CO ₂ : 511.07 KOH: 1124.31 H ₃ PO ₄ : 973.11	0.059–0.246 0.604 0.502	High surface area and pore volume: creates microporous–mesoporous hybrid structures
Physical activation of larch wood biochar with nitrogen at 700 °C [117]	8–122	0.01–0.85	Biochar with pore size of 4.62 nm with developed microporosity limited functional group formation due to inert atmosphere
Chemical activation with KOH on corn straw biochar [95]	2183.79	well-developed micropores	Strong adsorbent for heavy metals and PAHs cleanup from water
Chemical—physical activation of mixed woody sawdust with KOH and N ₂ at 900 °C [118]	327–1419	0.15–0.83	Promising catalyst with increased porous structure for use in syngas production
Chemical–physical activation of rice husk char with (KOH, H ₃ PO ₄ , ZnCl ₂) and N ₂ at 800 °C	89.93–390.59 346.30 394.50	0.079–KOH: 0.224 H ₃ PO ₄ : 0.189 ZnCl ₂ : 0.221	Best adsorption capacity for the tar removal

5. Comparative Discussion on Upgrading Methods

Having reviewed the pros and cons of different activation methods (Table 4) and gone through various studies, it was found that activation of biochar through acids like HNO_3 , HCl , H_2SO_4 and H_3PO_4 has been reported to significantly modify biochar surface functional groups toward increased oxygen content, amplified polarity and lessened ash percent. Different acids provide unique surface functionalization: e.g., HNO_3 treatment that triggers nitration and oxidation makes it possible for $-\text{NO}_2$ and $-\text{COOH}$ groups to be attached [72]. However, H_3PO_4 activation forms phosphorus-containing structures ($\text{P}=\text{O}$ and $\text{P}-\text{OOH}$), which can build stable complexes with pollutants [119]. During HCl treatment, the oxygenated functional groups are enhanced and mainly their carbonyl structures such as phenols, ethers and lactones are reduced. Such functional groups, largely present on the biochar edges and outer surfaces, facilitate hydrophilicity and polarity, whereas demineralization from acid activation tends to reduce ash content and surface charge which favors electrostatic attraction for cationic pollutants [120]. For acids, activation with HNO_3 does not significantly affect surface area and porosity in comparison with HCl , H_2SO_4 and H_3PO_4 . HNO_3 treatment at initial stages may cause micropore development and solubilize the inorganic impurities resulting in improved surface area and pore volume. But the long-time activation could cause the coagulation of crystallite and the narrowing of pores, reducing the change in the overall porosity [121]. It is thus claimed that H_3PO_4 is a milder acid, better preserving the carbon skeleton and more efficient in developing the micropores, and thus particularly useful for porosity engineering. On the other hand, alkaline activation (typically NaOH or KOH) enhances biochar's alkali groups and nonpolar surface sites, thereby promoting the adsorption of anionic pollutants. These processes mostly enhance the surface area and the pore volume, and KOH -activated biochar has a higher surface area and higher adsorption capacity with respect to steam activation [122]. Nonetheless, an over-alkaline treatment or higher temperature during processing might destruct pore structure because of a vigorous gasification reaction, which in turn results in decreased surface area [123]. From a chemical standpoint, it is well reported that KOH should enhance the surface functional groups (i.e., carboxylic, alcoholic and phenolic), but oxygen groups decompose upon heating, causing reducing overall functionality and raising carbon content [124]. In contrast, NaOH activation has received much less attention, although it is believed to be less expensive, less hazardous and more eco-friendly, and Na^+ can enter the biochar structure more effectively than K^+ because of its minor ionic radius [125].

Physical activation is further developed as a green approach for activating biochar, and steam, CO_2 and air/oxygen are the most used agents. The biochar developed by steam activation have excellent microporosity and adsorption capacity; however, the energy demand to produce steam for steam activation is questionable with regard to efficiency and sustainability [126]. On the contrary, CO_2 activation prefers the formation of mesoporosity and is efficient in accommodating large molecules, but its slow activation rates and the need for long processing times add up to energy requirements and a carbon footprint [94]. Air or oxygen activation is the simplest and the most economical way to obtain activated carbon, but it has the least control, and often results in overburning, less yield and low stability in biochar [127]. Physical activation, although avoiding toxic residues from chemical reagents such as KOH or H_3PO_4 and therefore being more environmentally friendly, has high thermal energy needs which remain significant environmental concerns [128]. Generally, in comparison with traditional activated carbon, physically activated biochars consume less in terms of ecosystem service as they are derived from renewable sources with less chemical inputs, especially in the field of wastewater treatment [129]. In addition, the environmental consequences are not limited to carbon costs; for example, soil quality improvements have

been recorded for steam-activated biochars, while other activation methods are associated with potential ecotoxic effects [130].

Table 4. Effects, risks and advantages of activation methods.

Method	Functional Effects	Advantages	Risks
Acid activation (HNO ₃ , HCl, H ₂ SO ₄ , H ₃ PO ₄)	- introduces oxygen-containing groups (-NO₂, -COOH, carbonyls). - Enhances polarity and hydrophilicity. - Demineralization reduces ash content. - H₃PO₄ forms phosphorus complexes.	- Improves adsorption of cationic pollutants. - H₃PO₄ preserves carbon skeleton and develops micropores. - HNO₃ increases surface area and pore volume (initially).	- Long HNO₃ treatment narrows pores and reduces porosity. - Risk of secondary contamination from strong acids. - Possible reduced stability with prolonged activation.
Alkali activation (NaOH, KOH)	- Enhances alkali groups and nonpolar surface sites. - Increases surface area and porosity. - KOH develops carboxylic, alcoholic, phenolic groups. - NaOH: cheaper, less hazardous, eco-friendlier.	- Strong adsorption of anionic pollutants. - KOH: Higher surface area. - NaOH integrates Na⁺ effectively in structure.	- Over activation or high temperature can collapse pores. - Heating decomposes oxygen groups, reducing functionality.
Physical activation (Steam, CO ₂ , Air/O ₂)	- Steam develops microporosity. - CO₂ favors mesoporosity (large molecules). - Air/O₂ oxidizes surface.	- Eco-friendly, no toxic residues. - Derived from renewable sources. - Steam produces excellent adsorption properties.	- Steam requires high energy, questionable efficiency. - CO₂ activation is slow and energy-intensive. - Air/O₂ can overburn, lowering yield and stability.

6. Potential Uses of Upgraded Biochar as a Carbon-Based Catalyst in Different Applications

Upgraded biochar holds promise far beyond its traditional uses, finding roles as catalyst support in methane cracking, biodiesel production, syngas upgrading and even cleaning up environmental contaminants. What makes it especially versatile is how its properties can be tailored, especially for catalysis [58,131]. Much of this capability stems from its natural mineral richness; elements like iron and potassium actively support catalytic processes [45].

6.1. Catalytic Methane Cracking

In the concept of methane cracking, research on the pore structure of a catalyst indicates that carbon catalysts with higher meso and macropore content achieve higher hydrogen production compared to those dominated by micropores [132]. This is caused by the diffusion of methane, which is hindered more severely in the presence of micropores, decreasing hydrogen production [133]. Figure 6, adapted from Harun et al. [134], illustrates a study on hydrogen production via catalytic methane decomposition with activated carbon, ruthenium-activated carbon and activated biochar catalysts at 800 °C. It is important to mention that the efficacy of these catalysts is influenced by their surface area, as a larger area enhances the adsorption of methane onto the catalyst surface.

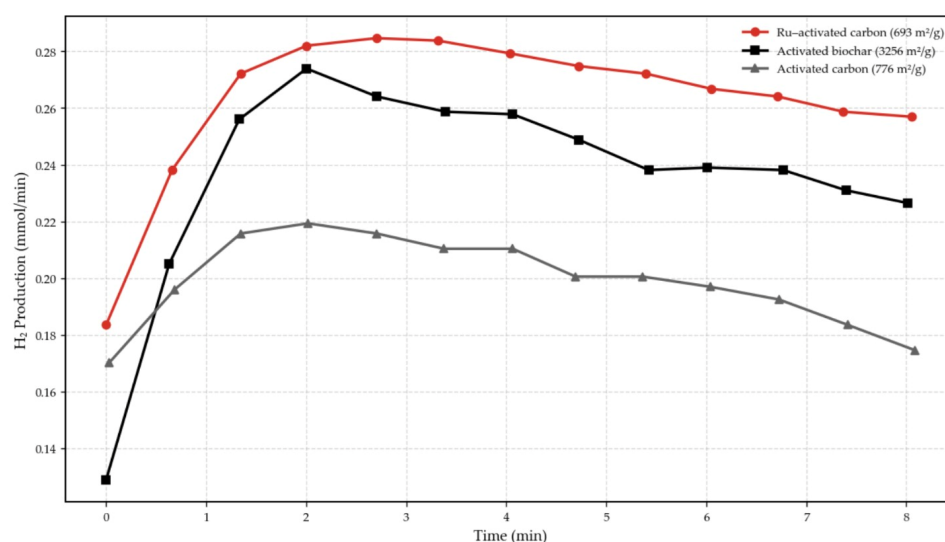


Figure 6. Hydrogen production from catalytic methane decomposition using Different Activated Carbons. Elaborated data from: [134].

Also, Abbas et al. [135] explored how varying the size of particles affects the methane conversion and decomposition rate using a fixed-bed reactor at a high temperature (850 °C) and a specific methane flow. Their findings suggested that changing the particle size within a fairly broad range (from relatively small to relatively large, specifically 135 to 1095 μm) had little noticeable influence on how much methane was converted, how quickly it decomposed or how long the catalyst remained active. Another study tested both biochar and activated char in methane decomposition, and their results showed that higher temperatures and lower methane concentrations improved conversion. At 900 °C with just 10% methane in a nitrogen stream, they achieved 72.3% conversion with activated char and 66.2% with biochar [49]. During acid treatments of biochar and extraction, some metals like Fe, Ni, Al and Ca, carbon-loaded catalysts showed improved methane conversion efficiency in the methane cracking processes [136].

Biochar's promise extends into dry methane reforming as well [137]. One study used a tungsten carbide-infused biochar, which maintained performance over 500 h. As CH_4/CO_2 ratios and temperatures increased, methane conversion decreased, but CO_2 transformation and hydrogen production grew steadily [138].

6.2. Support Biodiesel Production

Biochar has been found to be a promising catalyst and support for esterification and transesterification reactions in the processes of biodiesel production [139]. Its low cost, flexibility and mineral-abundant source of biowaste represent an appealing alternative to the typical homogeneous and heterogeneous catalysts [140]. Functionalization is one of the main factors that reinforces biochar catalytic activity. For example, a sulfonated biochar as a solid acid catalyst yielded approximately 97% biodiesel yield from FFAs (free fatty acids) and also produced 88% of productivity via simultaneous esterification and transesterification of vegetable oils [141]. However, biochar with rich content of inorganic minerals (such as calcium and potassium) can play a good role in solid base catalysts. For instance, palm kernel shell biochar, which was rich in CaO after calcination, exhibited 99% FAME (fatty acid methyl ester) content after the metanalysis of sunflower oil [142,143]. In another study, the KOH-activated pomelo peel biochar greatly achieved as high as 98% biodiesel yield because of its high basicity and enhanced surface area/porosity [144].

6.3. Syngas Upgrading (Tar Reforming)

Biochar plays an important role in the prevention of tar generation during biomass gasification and pyrolysis and improving the production of the syngas (a mixture of CO and H₂) [145]. The highest H₂/CO ratio (1.5) reported was attained by CO₂ physically activated biochar (900 °C) with both great SSA and hierarchical porosity, which had an excellent performance with 78.2 wt% of syngas yield and 26 mmol H₂/g-biomass of hydrogen production with a gas energy conversion efficiency of 73% [118]. Still, biochar performance can fall short due to its limited porosity and surface area. To overcome this, researchers are refining the material, activating the feedstock, adjusting synthesis settings, adding surface treatments and combining it with other materials. Its structure, high surface area, fine pores and active minerals behave like conventional catalysts (dolomite, olivine, nickel and alkali metals) and as an effective catalyst in tar reforming [146]. For example, Ni/Ca/Fe-loaded biochar-nanocatalysts achieved 87% tar conversion and 42.46 mmol H₂/g-biomass which demonstrated the excellent catalytic activity toward metal modifications [147]. In the case of pyrogas, Ren et al. found that using biochar as a catalyst during biomass pyrolysis increased pyrogas yield dramatically from 15% to 46% at 480 °C [148]. Shen et al. noted that the same approach raised hydrogen content in syngas by up to 27% [75].

6.4. Environmental Cleanup

Activated biochar is widely recognized as a cost-effective, environmentally friendly and highly efficient adsorbent of contaminants from water and soil. It does this in two primary ways: by adsorbing pollutants onto its porous surface and by functioning as a catalyst support to accelerate reactions that break down toxic compounds [149]. Biochar can help clean up the environment by immobilizing pollutants, passing on electrons to break them down or enabling oxidants, like peroxides, to generate powerful cleaning agents [150]. In advanced oxidation applications, physically activated biochar (e.g., peanut shell char pyrolyzed at 900 °C) exhibited 98.3% sulfamethotyl removal in 120 min because of its high graphitic content, COOH groups and enhanced specific surface area [151]. Chemically oxidized biochar also performed well, with over 97% removal of the antibiotic sulfamethoxazole in 30 min [152].

7. Waste-Derived Biochar Chemical Leaching and Its Potential for Critical Materials Recovery

The evolving circular economy favors the recovery of critical materials, where biochar can assist. A shift is needed from conventional linear economic systems toward one that focuses on resource reuse and waste reduction. The supply of critical raw materials components of many industries faces potential vulnerabilities. Recovering valuable materials from waste biochar has the advantageous potential for helping to secure the sustainable supply of critical raw materials, which reduces reliance on environmentally harmful extraction processes [153].

Biochar's characteristics can be influenced by the quality of raw materials and processing conditions, requiring some additional refinement. While biochar has promise, it is often derived from biowaste which is rich in certain inorganic compounds, such as Fe, Si, K and Al, that dilute porosity and may inhibit catalytic processes [154]. Through chemical leaching, valuable materials can be retrieved from waste-derived biochar promoting circularity. Through this form of leaching, helpful nutrients such as phosphorus, potassium, magnesium and calcium can be extracted and used as fertilizers while heavy metals are removed. Leaching enhances biochar's porosity and adsorption capacity by reducing ash content, thus improving its application in environmental remediation and energy recovery. The process also helps in lowering alkaline earth metals (mostly Ca²⁺ and Mg²⁺), prevent-

ing carbonate formation that could otherwise diminish biochar's efficacy [30,155]. For this reason, chemical leaching can be considered as an upgrading process for the biochar.

As has been reviewed, alkali treatments using NaOH, KOH and $\text{Ca}(\text{OH})_2$ are particularly effective in extracting silica, alumina and clay minerals from coal, forming hydrated alkali bearing silicate and aluminate. These treatments are also suitable for extracting inorganic and organic sulfur [156]. In a study by Qiu et al. [157], potassium bicarbonate (KHCO_3) showed promise as an activating agent in pyrolysis for creating hierarchically porous biochar adsorbents (14.4 mg/g.min) from sewage sludge, supporting material recovery from water leaching which can be used for plant growth (Figure 7).

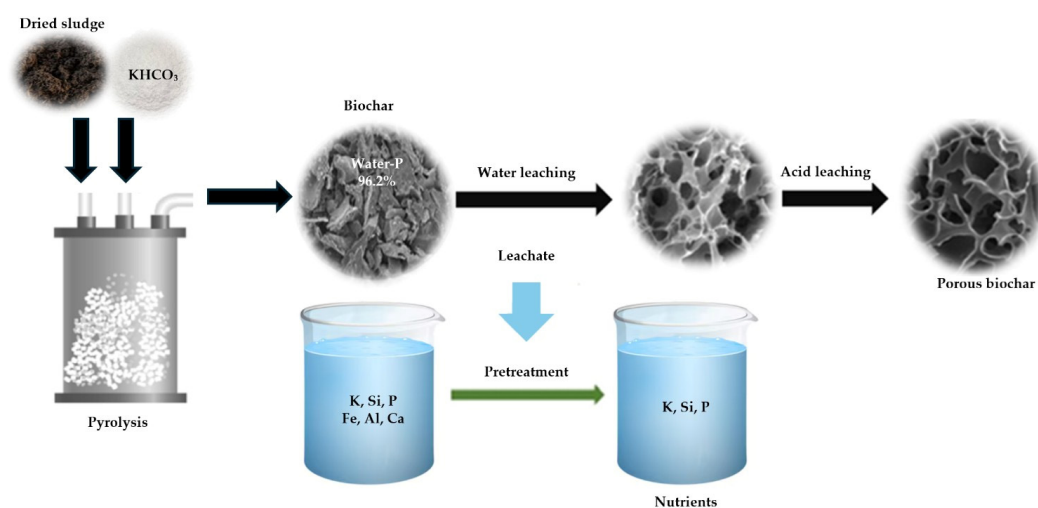


Figure 7. Biochar after both water and acid leaching treatments. Adapted from [157], Resources, Conservation and Recycling, 176 (2022), 105953. © 2022 Elsevier B.V. Reproduced with permission.

Conversely, acid leaching, which employs acids such as H_2SO_4 , HNO_3 and HCl , is more effective in removing carbonates, phosphates, iron (III) oxide and sulfides from low-grade coal. Sequential application of alkali and acid leaching is often beneficial, as alkalis react with sulfur and primary coal minerals to form compounds that acids can later dissolve [158]. Acid leaching is also employed for phosphorus recovery from sewage sludge and incinerated sludge ashes with HCl or H_2SO_4 , as they are particularly effective in dissolving phosphorus bound to aluminum and calcium [159].

In a different investigation carried out by Salimbeni et al., sewage sludge char produced through slow pyrolysis at 450°C was further treated using HNO_3 , HCl and KOH through chemical leaching under specific conditions ($60\text{--}80^\circ\text{C}$, 2 h contact time). The main phosphorus and calcium extraction results indicate that leaching could achieve almost complete removal (100% removal) and ash content reduction (61.4% removal). As a result, the biocoal obtained demonstrated substantially higher carbon content (rising from 14.4% to 27.4%) as well as greater surface area (SSA) of $70\text{ m}^2/\text{g}$. This greatly improved its potential valorization. While silica remained, and Al/P separation was challenging, single-stage HNO_3 leaching was deemed promising for producing P-rich fertilizer precursors and upgraded carbon material [30]. Also, Salimbeni et al. showed how chemical leaching can affect the char derived from industrial sludge and recover the inorganic material for the fertilizers, as can be seen in Figure 8. They started with slow pyrolysis of the dried sludge, continued by acid leaching of the resulting char / yield biocoal and a nutrient rich leachate, and subsequently physically activated the biocoal using CO_2 at $700\text{--}800^\circ\text{C}$. Chemical precipitation was employed for phosphorus recovery from the leachate. The results showed that leaching effectively removed around 88% of ash content in char, with close to 100% of P, Mg, Ca and Fe, and around 90% of Al. The activated carbon derived

exhibited a SSA of ($417 \text{ m}^2/\text{g}$), with potential as an adsorbent for emission or wastewater treatment, and suggested other potential applications like metallurgy [160].

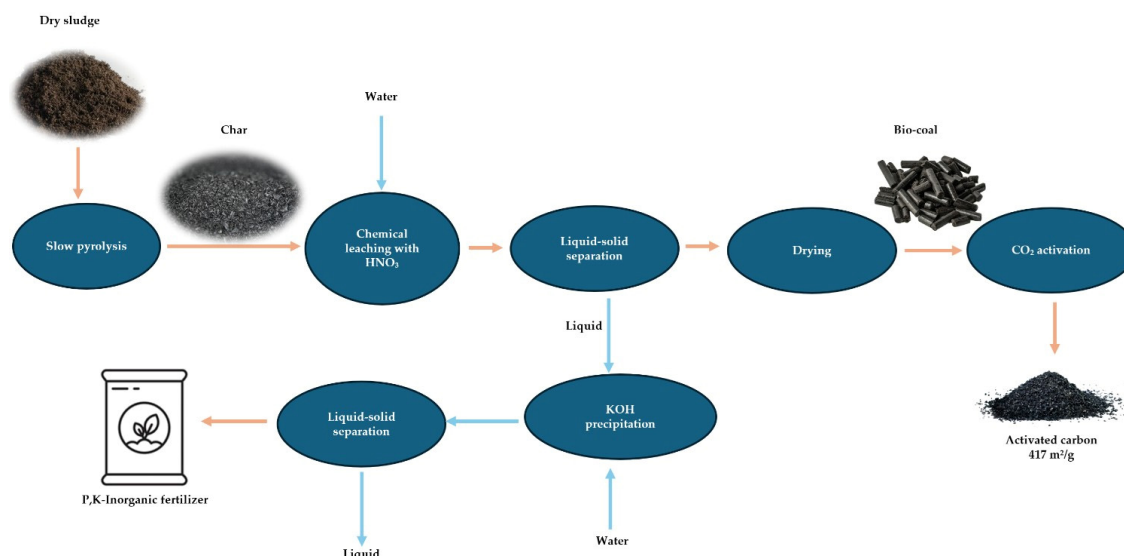


Figure 8. Elaborated methodology for the thermochemical treatment of derived-sludge char and material recovery [160].

8. From Production to Upgrading: Techno-Economic Concerns and Prospects

The future of biochar upgrading faces opportunities and critical pathways. Many of these are interconnected, and multiple aspects must be approached in order to optimize production processes, improve performance for specific applications and scalably deploy over time.

8.1. Biomass Compatibility

The yield and quality of biochar were determined by the type of biomass. When used as an amendment in soil remediation, carbon sequestration or water treatment technologies, biochar performance is often impeded by contamination and impurities that are present in the biomass. Unsuitable size or shape of feedstock often require expensive pre-treatment operations. It will require long-term commitments to stimulate energy crop production, promote sustainable land management practices and exploit the agro-forestry by-products [161].

8.2. Scale-Up Limitations

Biochar production can suffer from the variability in pyrolysis conditions and operating parameters so that quality is inconsistent and performance is uncertain. Scaling up is difficult due to high costs, inadequate infrastructure and the lack of standard setting in industrial production. Solving these issues will involve advancing reactor designs and energy-efficient processes, as well as common quality standards backed by policy incentives and certification schemes that can promote competition and confidence in the market to foster widespread use [162].

8.3. Balancing Upgrading Efficiency with Environmental Risks

Although the main objective of increasing biochar physicochemical performance is to render it more efficient in different applications, its sustainability may be compromised by the upgrading methods. For example, modifications with metal salt increase the presence of oxygen-containing functional groups, which directly increases inorganic contaminant adsorption. These treatments also alter the leaching potential of metal ions and pose a risk

to cationic secondary contamination [163]. Regarding physical activation when it comes to high-energy ball milling, surface area, particle size and functional groups are said to be more important, as they are a solvent free, reproducible and cost-effective green technology. On the other hand, they present issues such as high energy consumption, equipment durability, and dust emission [164]. Also, the polycyclic aromatic hydrocarbons (PAHs) that are formed during pyrolysis need to undergo accurate risk assessments, as PAH-contaminated soil and crops. Therefore, actions must be taken in balancing the efficiency–sustainability trade-off in biochar upgrading using innovation or life-cycle assessments and risk management.

8.4. Economic Obstacles

Economic constraints continue to be a major hurdle for widespread biochar upgrading. Today, the cost of biochar production is still too high and would need to decrease by two-thirds, or carbon subsidies would need to rise between two- and ninefold above current EU emissions trading scheme (ETS) prices (≈ 88 EUR/t CO₂eq) to become economically competitive according to studies [165]. The market itself is still immature and lacks universal quality standards, which constrains investor confidence and adoption. On the agricultural side, where biochar is often treated as a soil amendment, entry capital costs may be high. For instance, when 40 t/ha of biochar was used at a cost of ~ 190 /t in a Mediterranean study, the break-even time under normal condition was about 5.5 years in an amortization period [166]. Relative to other forms of carbon dioxide removal (CDR), biochar has received small levels of support, while existing subsidies are too weak to ensure widespread uptake. Despite all of the tunable properties of activated biochar, some difficulties such as high fabrication cost, poor long-term stability, industrial scale-up and lack of understanding of their catalysis mechanism still exist. These issues must be addressed in order to realize the full potential of biochar in a sustainability circular bioeconomy with respect to industrial scale. In the future, biochar upgrading will go beyond agriculture and forestry by offering new research opportunities in environmental remediation, horticulture, construction and materials science. These choices necessitate pathways, low in cost, to upgrade them, such as the production of ‘green’ biochar from a variety of residues, which also may contribute to reductions in costly feedstock dependence and to circular resource use. It is also crucial to establish the lowest effective application levels of advanced biochar at which they can yield top performance while minimizing costs and risks of collateral pollution. Ultimately, the translation from bench to application will depend on the success of extended field trials and weathering studies to guarantee novel modification approaches yield highly performant biochar that is both durable and safe at large-scale production. Collectively, they envision the opportunities and difficulties that form the basis for the next phase of biochar upgrading technologies.

9. Conclusions

This review has discussed the comparative properties of different thermochemical conversion processes, highlighting that slow pyrolysis continues to be the most efficient pathway for producing well-tuned pore structure and stable biochar, making it suitable to valorize waste streams and supports through the short-term mitigation of greenhouse gases by sequestering carbon. The following upgrading steps substantially enhance biochar performance with the specific surface area up to 2100 m²/g by KOH activation, and hybrid microporous–mesoporous structure is obtained with a pore volume of 0.83 cm³/g through a combined (KOH-N₂) treatment while acid activation provides near-total removal of ash (removal factors of up to 88–100%) enriched with the surface oxygen functionalities that are necessary for suitable adsorption and catalytic activity. At the catalytic level, upgraded biochar significantly outcompetes common catalysts. In the case of methane

cracking, activated char gave 72.3% CH₄ conversion at 900 °C and 66.2% in the case of raw biochar. As the catalyst support for biodiesel production, the sulfonated biochar was capable of achieving a 97% biodiesel yield from the free fatty acids. For syngas utilization, CO₂-activated biochar co-gasified performed a high 78.2 wt% vol syngas yield at 26 mmol H₂/g biomass and produced energy conversion efficiencies of up to 73%. Environmental remediation performance was similarly strong, with oxidized biochars removing >97% of antibiotics, and thermally activated chars removing 98.3% of sulfamethoxazole. Regarding materials recovery, sequential acid–alkali leaching of waste-derived biochar enabled a nearly 100% recovery of P, Mg, Ca and Fe, as well as the substantial decrease of ash content by 61–88%. This not only allowed the recovery of EU-listed critical raw materials but also increased carbon content (from 14.4% to 27.4%) and specific surface area (up to 417 m²/g), at the same time improving the biochar’s catalytic and adsorptive capacity.

Despite all of the progress in biochar upgrading, there are several issues which need to be considered, such as production cost, scaling up the upgrading process and disposal of solid residues. Overcoming these barriers will require technological advancements and innovative policy frameworks. Crucially, biochar upgrading provides real-world opportunities in developed economies. It is consistent with circular economy objectives and carbon neutrality commitments; decentralized schemes can displace open burning, enhance soil fertility and decrease dependence on imported fertilizers. Moving forward it will be crucial to interface the emerging technologies with sound economic projections and enabling regulations if biochar is to realize its potential as a sustainable climate change mitigation and resource recovery.

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Conflicts of Interest: The authors declare no conflict of interest.

Abbreviations

The following abbreviations are used in this manuscript:

SSA	Specific Surface Area
TPV	Total Pore Volume
PV	Pore Volume
HRT	Highest Reactor Temperature
HR	Heating Rate

HHV	Higher Heating Value
RT	Residence Time
PAH	Polycyclic Aromatic Hydrocarbons
HTC	Hydrothermal Carbonization
OCFGs	Oxygen-Containing Functional Groups
AC	Activated Carbon
CE	Circular Economy
GHGs	Greenhouse Gases
ETS	Emissions Trading Scheme

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