

RESEARCH ARTICLE

Sustainable ultra-strong thermally conductive wood-based antibacterial structural materials with anti-corrosion and ultraviolet shielding

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Abstract

In light of the uprising global development on sustainability, an innovative and environmental friendly wood-based material derived from natural pinewood has been developed as a high-performance alternative to petrochemical-based materials. The wood-based functional material, named as BC-CaCl₂, is synthesized through the coordination of carboxyl groups (–COOH) present in pinewood with calcium ions (Ca²⁺), which facilitates the formation of a high-density cross-linking structure through the combined action of intermolecular hydrogen bonds. The as-prepared BC-CaCl₂ exhibits excellent tensile strength (470.5 MPa) and flexural strength (539.5 MPa), establishing a robust structural basis for the materials. Meanwhile, BC-CaCl₂ shows good water resistance, thermal conductivity, thermal stability, UV resistance, corrosion resistance, and antibacterial properties. BC-CaCl₂ represents a viable alternative to petrochemical-based materials. Its potential application areas include waterproof enclosure structure of buildings, indoor underfloor heating, outdoor UV resistant protective cover, and anti-corrosion materials for installation engineering, and so forth.

KEYWORDS

coordination crosslink, excellent mechanical performance, ultra-strong, wood-based composite

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1 | INTRODUCTION

Petrochemical energy serves as a vital component for industrial operations, playing a crucial role in human production and daily life.^{1–3} Its products, derived from petroleum processing, includes gasoline, kerosene, diesel and various raw materials for polymer materials such as plastics, synthetic fibers, and synthetic rubber and so on.^{4,5} These petroleum products have been widely used in various packaging and construction industries as a structural functional material.⁶ However, the ever-increasing exploitation of oil as a non-renewable energy source will inevitably lead to energy depletion. Additionally, the diminishing availability of raw materials will necessitate a reduction in the production of petroleum-based materials.⁷ Furthermore, the stable chemical properties of such materials make them difficult to degrade effectively.⁸ Consequently, various waste materials derived from the petroleum-based products accumulate in the environment exerting great pressure on the environment.^{9,10} Today, green development is a global consensus for sustainable growth. After the pandemic, many countries have adopted national policies with the primary goal of achieving “green recovery”.¹¹ In this context, the development of bio-based materials utilizing biomass as raw materials has gained significant momentum,^{12–15} positioning it as one of the most promising industries at present. Moreover, the bio-based materials are poised to become a leading industry, driving scientific and technological innovation as well as economic development on the global scale.¹⁶ Therefore, it is crucial to develop an all-green sustainable high-performance bio-based structural functional materials that rely solely on 100% biomass.^{17,18}

Biomass is considered to be the main sustainable source of organic carbon on the earth,^{19–24} a perfect equivalent to oil, and a sustainable green raw material

for producing fuel and fine chemicals with net zero carbon emissions.^{25–27} Meanwhile, bio-based composite materials are a new type of material that has emerged in recent years,^{28–30} which refers to a new type of material that uses renewable biomass (such as crop straw fiber, bamboo and wood fiber, etc.) or materials made from biology as raw materials and is then compounded with other materials through biological, chemical, and physical methods.^{31–34} Since the production of new building materials and building components is an indispensable link in the development of the modern construction industry, this marks a major change in the production method and represents a future development trend in the furniture and building materials industry.^{35–37} Meanwhile, in the “Guiding Opinions of the General Office of the State Council on Promoting the Steady Growth of the Building Materials Industry” issued by the State Council of China, it is clearly proposed to “support the development of biomass building materials by using crop straw, bamboo fiber,³⁸ wood chips, and so forth, and develop biomass fiber reinforced wood plastic building materials and other products”. Hence, the new biomass structural functional materials developed are the most original ecological synthetic environment-friendly materials, which conform to the direction of novel environment-friendly synthetic materials.^{38–40} However, the biomass structural functional materials constructed so far either have poor mechanical properties or rely on the synthesis of petrochemical adhesives (phenol-formaldehyde resin or urea-formaldehyde resin).⁴¹ These drawbacks will not only cause accelerated consumption of petrochemical energy, but also lead to serious environmental pollution and ecological damage. Therefore, it is of great significance to develop an advanced method to prepare new sustainable bio-based structural functional materials.^{42,43}

As a traditional building material, wood has been widely used in ancient and modern architecture.^{44–46}

As a kind of coniferous plant, pinewood is widely used in structural building materials and furniture products because of its low price, natural texture and long service life.⁴⁷ However, pinewood also has many defects. On one hand, it is soft and easy to crack and deform due to its high moisture content. On the other hand, the large amount of rosin in the pinewood will block its tracheid. It is often subject to degreasing treatment during use, which makes its processing complex.⁴⁸ Therefore, the high-value utilization rate of pinewood is not high. To combat this, high-temperature steaming or chemical treatment is usually used to reduce the resin content of wood. However, these methods can damage the structure of the pine itself. Hence, the extraction of pine resin by extraction has become a common treatment method for the high-value utilization of pinewood.

The performance of bio-based structural functional materials can be enhanced through the design of micro-nano structures.⁴⁹ Thereinto, Ca^{2+} , as a common metal ion, is usually cross-linked through coordination with $-\text{COOH}$, combining with high-density hydrogen bonds between cellulose microfibrils and hemicellulose, to form a special high-density cross-linked network inside the wood. Under this treatment condition, the strength and dimensional stability of bio-based structural and functional materials are greatly enhanced, and they are endowed with multiple functions. Hence, aiming to make full use of pinewood to prepare high-performance and sustainable wood-based structural functional materials, a robust and effective method is reported in this study. The degreased pinewood is first chemically pretreated, then coordinated with $-\text{COOH}$ in the pinewood under the action of Ca^{2+} , and finally thermally molded into high-strength wood-based composite. The material shows remarkable mechanical properties, good dimensional stability, excellent thermal conductivity and outstanding UV resistance.⁵⁰ The excellent mechanical properties of BC- CaCl_2 provide a solid foundation for its application as a structural material in the building materials industry, such as anti-seismic buildings. In particular, pinewood is a fast-growing wood. The prepared high-performance wood-based composites address the inefficient use of pinewood and achieve the high-value utilization of natural wood. Therefore, this study complies with the ultimate strategy of global sustainable development. Meanwhile, the production process of this material is green and pollution-free, which meets the policy requirements of prioritizing ecological environment in socioeconomic development.⁵¹ In addition, the material meets the requirement of excellent properties for a renewable and low-cost alternative to traditional petrochemical-based functional materials.⁵² The potential application areas for BC- CaCl_2 include waterproof enclosure structure of

buildings, indoor underfloor heating, outdoor UV resistant protective cover and anti-corrosion materials for installation engineering, and so forth.

2 | MATERIALS AND METHODS

2.1 | Raw materials and preparation of wood-based composites

The pinewood was purchased from Suqian Timber Wholesale Market in Jiangsu Province. After careful selection (from the same pinewood and free from cracks and growth defects), it was cut into wood chips of the same specification (length 50 mm, width 50 mm, thickness 5 mm). NaOH (96%), Na_2SO_3 (98%), and CaCl_2 (99%) were purchased from Shanghai Macklin Biochemical Co., Ltd., China.

Preparation of wood-based composites: First, the pine resin was fully removed from the cut pinewood chips by Soxhlet extraction (BC-E). Second, the extracted pine chips were immersed in the mixed aqueous solution of 2 M NaOH and 0.4 M Na_2SO_3 (solid-to-liquid ratio = 1:10), stirred and heated at 96°C for 4 h, rinsed with deionized water, and freeze-dried (BC-P). Next, the pretreated pinewood chips were immersed in CaCl_2 solution for 24 h in vacuum, rinsed with deionized water, and freeze-dried (BP- CaCl_2). Finally, the extracted, pretreated and impregnated pinewood chips were placed into the customized molds respectively (the dimensions of the slot: length 50 mm, width 50 mm), and the molds were pressed in a hot press machine (Dongguan Zhenggong Electromechanical Equipment Technology Co., Ltd., China) at a pressure of 45 MPa and a temperature of 180°C. After hot pressing for 1 h, the mold was cooled to room temperature and three kinds of the wood-based composites were taken out, named as: BC-E, BC-P, and BC- CaCl_2 .

2.2 | Characterization

In each test, three samples were analyzed in parallel according to the requirements of GB/T 24511-2017. The density ρ of the prepared wood-based composites was calculated according to the formula $\rho = \text{mass}/\text{volume}$. The AGS-X universal mechanical testing machine (Shimadzu, Japan) was used to investigate the mechanical properties of the samples. Specifically, the test was performed on a sample of 50 mm × 6 mm × 3 mm at room temperature at a displacement rate of 1.5 mm/min. The lightness of the samples was measured with a colorimeter (SR60, Mitutoyo Precision Measuring Instrument Co., Ltd., Guangdong,

China). In the water resistance test, the prepared three samples were placed in water, and the changes of water absorption rate and water absorption thickness swelling rate were compared within 48 h. The DSA100S contact angle measuring instrument (KRUSS) was used to measure the contact angle of samples. 2 μ L of water was dropped onto the sample each time, and the contact angle was recorded within 10 s. Sulfuric acid (4 μ L, 78%) (Nanjing Chemical Reagent Co., Ltd, China) was dropped on three samples respectively, and their corrosion resistance was analyzed by observing their surface states.

Thermal conductivity of the three samples was characterized by a thermal conductivity tester (DRPL-2B, Xiangtan Instrument Co., Ltd., China). In the case of continuous heating by the laser heater for 3 min, the samples were photographed and characterized by an infrared thermal imager (FLIR E5xt, FLIR, USA). The thermal stability of the samples were analyzed using a thermogravimetric analyzer (TA Instruments, TGA55 USA) at a heating rate of 20°C/min from room temperature to 640°C in a nitrogen atmosphere.

In the flame-retardant test, three samples were burned under an external flame of the alcohol lamp for 10 s. The burning conditions and final combustion conditions of the samples in different time periods were observed and compared. At 210°C, the samples were heated with acrylonitrile butadiene styrene (ABS), polyvinyl chloride (PVC), polypropylene (PP), polycarbonate (PC) and polyamide (PA) purchased from Chang'an Wald Plastic Products Factory in Guangdong, China, and their morphological changes were compared.

The ultraviolet visible (UV-Vis) spectrophotometer (UV1900, Shandong Qingdao Jingcheng Instrument Co., Ltd.) was used to characterize the UV transmittance of the samples. Quanta 200 scanning electron microscope (SEM, FEI, China) was used to characterize the cross-section morphology and energy dispersive spectra (EDS) of the samples.

The antibacterial performance of *Escherichia coli* was evaluated against a sample plate with a length and width of 2 cm \times 2 cm.⁵³ The antibacterial properties of the samples were characterized according to the distribution range of *Escherichia coli*. Furthermore, this antibacterial experiment should be repeated 4 times.

2.3 | Chemical characterization

The National Renewable Energy Laboratory (NREL) method was used to determine the cellulose, hemicellulose, and lignin content in the samples. Functional groups of the samples were analyzed using VERTEX 80 V ATR Fourier infrared spectroscopy (FT-IR, Bruker

Technology Co., Ltd., Germany). X-ray diffraction (XRD, Beijing Puwei General Instrument Co., Ltd., China) was performed in the scanning range of 5 to 60° (2 θ) to analyze the diffraction patterns. X-ray photoelectron spectroscopy (XPS, Shimadzu Enterprise Management Co., Ltd., China) was used to investigate the elemental composition and molecular structure of the samples. The ¹³C-NMR spectra of the samples were tested by high-resolution solid-state nuclear magnetic resonance spectroscopy (NMR, Bruker 400 M, Germany).

3 | RESULTS AND DISCUSSION

3.1 | Preparation of wood-based composites

Figure 1 shows the preparation process of high-strength wood-based composites. After most of the pine resin inside the extracted pinewood chips is removed to form a loose and porous structure, we obtain the raw material denoted as BC-E (Figure 1A). After chemical pretreatment with NaOH and Na₂SO₃, the sulfonation reaction occurs that etches the pinewood chips to remove part of the lignin (Figure S1A), and the raw material of BC-P is obtained.⁵⁴ Under vacuum-assisted conditions, the pretreated pinewood chips are fully immersed in the CaCl₂ solution to obtain the BC-CaCl₂ raw material. Finally, the three raw materials are pressed into regular plates with stable morphology by one-step hot molding.⁵⁵

During the thermoforming process, part of the lignin seeps out of the plates in the molten state, and covers the surface of the material after cooling, forming a dense protective layer. The prepared high-performance wood-based structural functional material BC-CaCl₂.⁵⁶ The black surface of BC-CaCl₂ verifies that lignin covers the surface of the materials. The BC-CaCl₂ exhibits better density, mechanical properties, thermal conductivity, UV resistance, and water stability than BC-E and BC-P (Figure 1B). Especially, BC CaCl₂ shows excellent mechanical strength, indicating that under the action of Ca²⁺, the treated pinewood produces a stable cross-linked structure. Meanwhile, BC-CaCl₂ is green and pollution-free, which enables its usage in building materials to replace the traditional petrochemical-based structural materials.⁵⁷

3.2 | Chemical mechanism of preparation of BC-CaCl₂

Figure 2 reveals the strengthening mechanism of Ca²⁺ on the internal molecular structure of pretreated

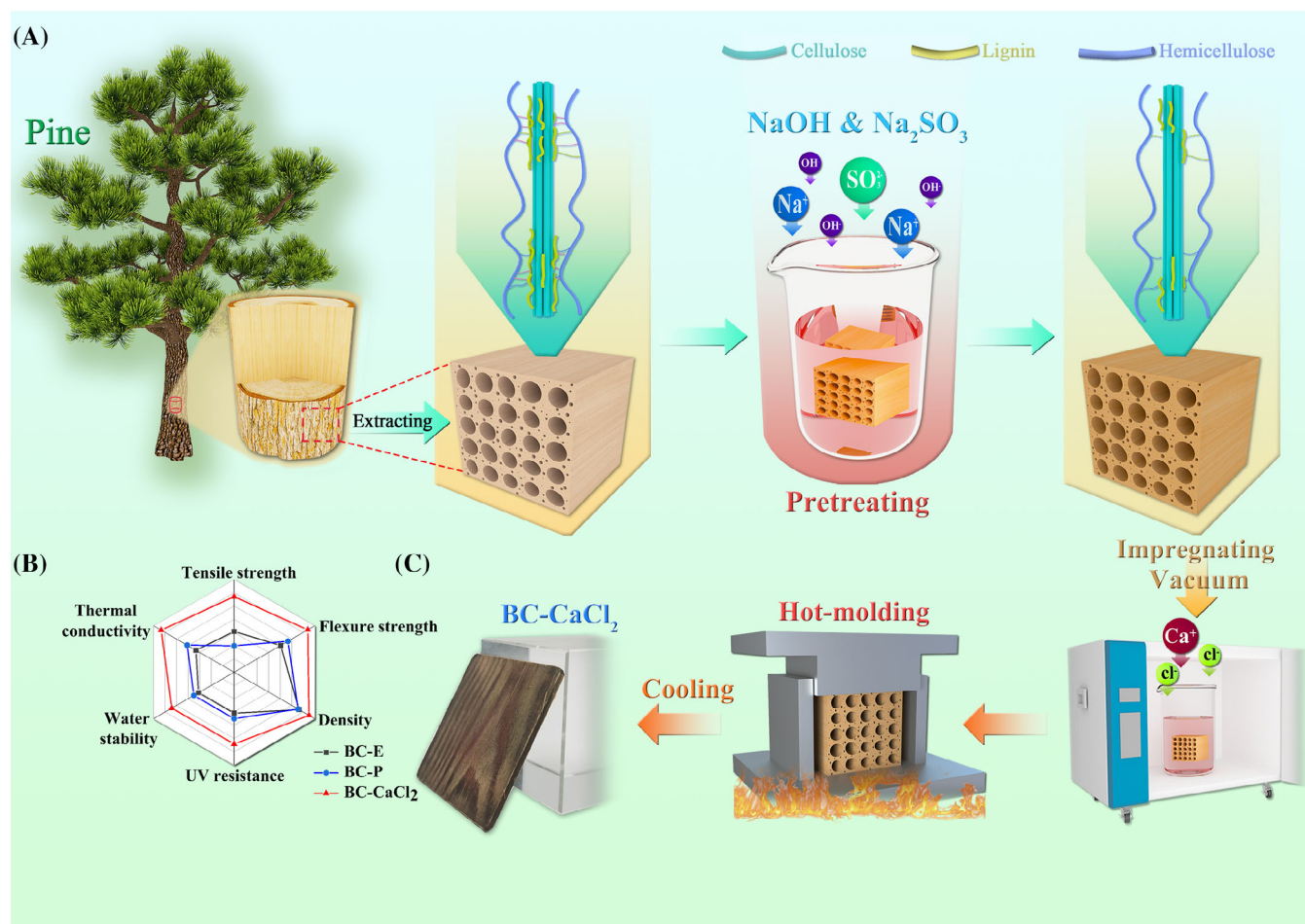


FIGURE 1 (A) Schematic diagram of the preparation process of high-strength wood-based composites. (B) The density, mechanical property, thermal conductivity, UV resistance and water stability of BC-E, BC-P, and BC-CaCl₂.

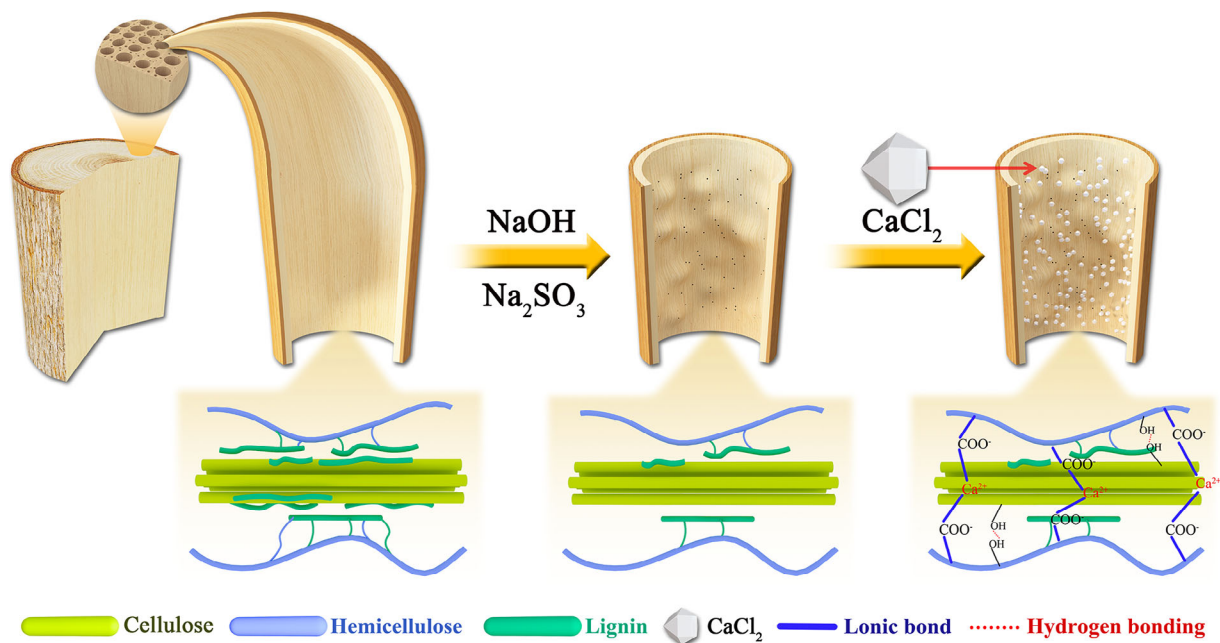


FIGURE 2 Schematic diagram of chemical mechanism of BC-CaCl₂.

pinewood. First, most of the pine resin has been removed from the extracted pinewood, making it loose and porous, showing a well-defined cross-linked structure of cellulose, hemicellulose and lignin. Then, the internal structure of the pinewood is etched under the treatment of an alkaline solution, removing part of the lignin to expose the internal cellulose microfiber and hemicellulose structure. Finally, under the action of Ca^{2+} , a special high-density cross-linked structure can be formed with pinewood, including high-density hydrogen bonding between cellulose microfiber and hemicellulose, and coordination crosslinks between Ca^{2+} and $-\text{COOH}$. This high-density cross-linked structure can enhance the structural stability and strength of BC- CaCl_2 , and improve its hydrolysis resistance and durability.^{50,58}

Meanwhile, the presence of Ca^{2+} endows BC- CaCl_2 with certain antibacterial properties.

3.3 | Chemical properties of BC- CaCl_2

Under vacuum impregnation of CaCl_2 solution, the white crystalline building blocks in BC- CaCl_2 are composed of the cross-linked structure formed by $-\text{COOH}$ in hemicellulose and Ca^{2+} (Figure 3A).⁵⁹ Figure 3B,C shows the general distribution of C and Ca elements. Meanwhile, Figure 3D shows the mass percentage distribution of Ca element. Therefore, EDS composition analysis shows that BC- CaCl_2 contains a high concentration of Ca atoms, which further proves the strong binding between

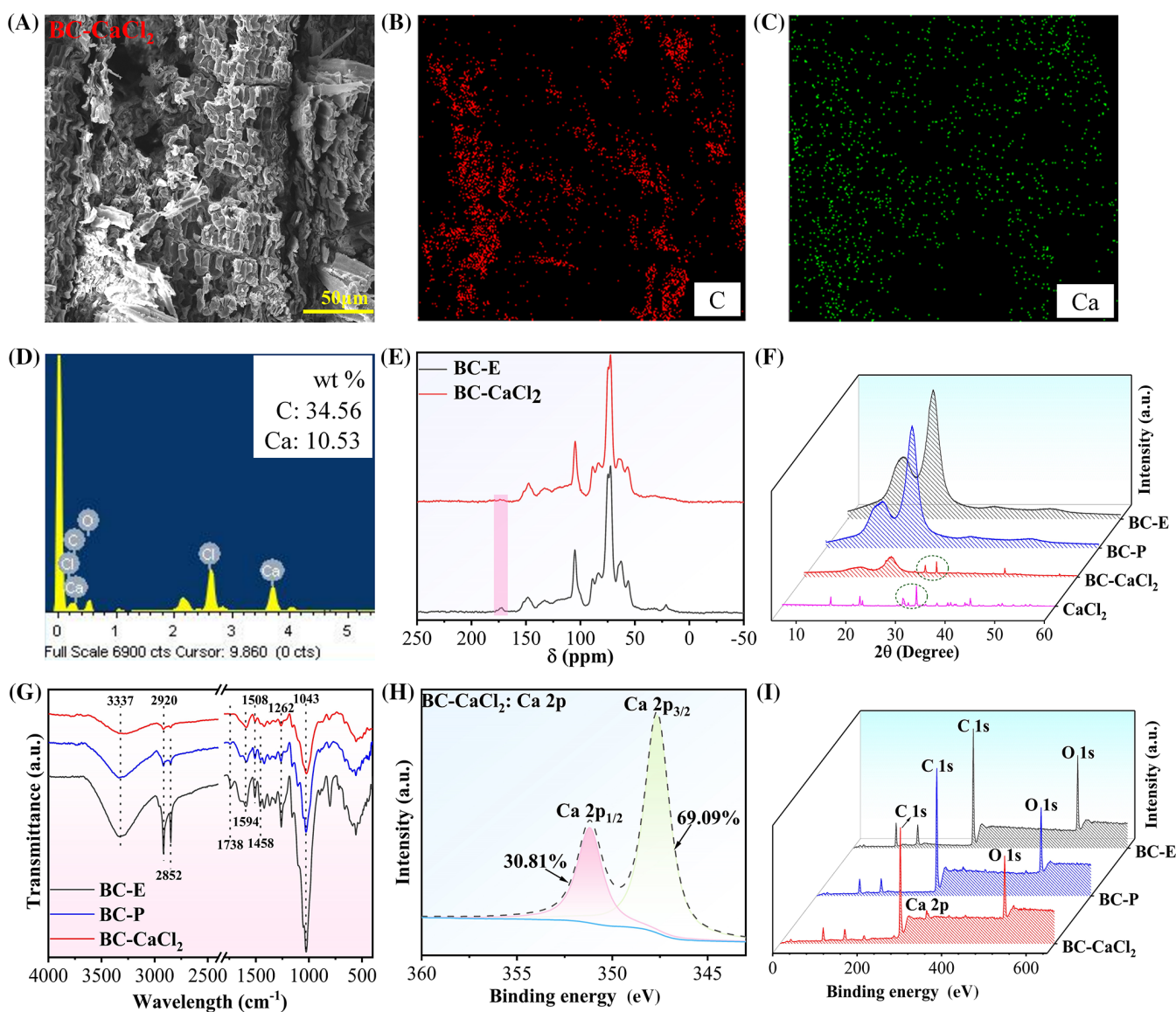


FIGURE 3 (A) Section SEM image of the fractured BC- CaCl_2 . The mapping of elemental (B) C and (C) Ca. (D) EDS spectrum the element content in BC- CaCl_2 . (E) Comparison of ^{13}C -NMR of BC-E and BC- CaCl_2 in the range of -50 – 250 ppm. (F) XRD patterns of BC-E, BC-P, BC- CaCl_2 , and CaCl_2 . (G) FT-IR and (I) XPS total spectra of BC-E, BC-P, and BC- CaCl_2 . (H) Ca 2p XPS spectra of BC- CaCl_2 .

–COOH in hemicellulose and Ca^{2+} .^{60,61} Through ^{13}C -NMR spectrum analysis, the peak value of BC- CaCl_2 in hemicellulose (170–175 ppm) disappears almost completely, which is due to the coordination crosslinks between –COOH in hemicellulose and Ca^{2+} , forming a stable wood structure (Figure 3E).⁶² As shown in Figure 3G, the peak of BC- CaCl_2 at 3337 cm^{-1} (O–H bending vibration) is significantly reduced compared to BC-E and BC-P, which indicates the reduction of free hydroxyl. The absorption peaks of lignin at 1594, 1508, and 1458 cm^{-1} (benzene ring skeleton vibration) of BC- CaCl_2 are consistent with those of BC-P and much lower than those of BC-E.^{63,64} This is because the lignin in pinewood is partially removed under the pretreatment of alkaline solution. Meanwhile, the peak of BC- CaCl_2 at 1738 cm^{-1} almost completely disappeared, indicating the removal of the characteristic absorption peak (C=O stretching vibration) of hemicellulose formed by the acetyl and carboxyl groups of hemicellulose, which is consistent with ^{13}C -NMR results.⁶⁵

In addition, the characteristic peaks of Raman spectrum in the range of $1535\text{--}1650\text{ cm}^{-1}$ are mainly derived from lignin (Figure S1B). The image shows that the peaks of BC- CaCl_2 at 1537 cm^{-1} (CH_3 bending in O– CH_3) and 1643 cm^{-1} (C=C telescopic vibration) are significantly weaker than those of BC-E, which indicates a substantial removal of lignin and is consistent with the above results.⁶⁶

X-ray diffraction patterns show that only BC- CaCl_2 has a diffraction peak consistent with CaCl_2 in the range of diffraction angle from 30 to 40° (Figure 3F). This proves that CaCl_2 exists in the interior of the pretreated pinewood after vacuum impregnation, thus providing coordination conditions for the internal crosslinks of BC- CaCl_2 . In addition, the weakening of the diffraction peaks in the BC- CaCl_2 crystalline and the amorphous regions are due to the structural changes caused by the addition of CaCl_2 crystals.

The structure composition of BC- CaCl_2 was determined by analyzing three peaks of C, O, and Ca in the XPS full spectrum. The structural composition of BC- CaCl_2 was determined by analyzing the three peaks of C, O, and Ca in the XPS full spectrum. As shown in Figure 3H,I, only the XPS spectrum of BC- CaCl_2 has a peak of Ca 2p, indicating the introduction of Ca^{2+} successfully formed ion bonds with –COOH in hemicellulose, and formed a high-density cross-linked structure under the interaction of hydrogen bonds, thus building a stable wood system.⁶⁷

3.4 | Morphology and mechanical properties of BC- CaCl_2

SEM images reveal that the cross-section of BC-E is clearly layered, and the fibers are compressed and tightly

connected after the cell wall collapses (Figure 4A). Hence, there are still some gaps between the fiber layers. Figure 4B shows that the internal cell structure of BC-P thermoformed after alkali treatment is destroyed, and the complete wood structure cannot be seen in the section.⁶⁸ Since part of the lignin has been removed, its internal structure is partially porous. As supported by Section 3.2, BC- CaCl_2 forms coordination crosslinks under the action of Ca^{2+} , and recombines the fibers under the action of intermolecular hydrogen bonds, showing a stable internal structure (Figure 4C).⁶⁹ Meanwhile, its internal structure is composed of closely connected states similar to building blocks, so there are almost no pores. In addition, BC- CaCl_2 becomes dense after hot pressing, reducing its thickness by more than 65% (Figure S2B). Although the mass of BC-P is reduced due to the removal of lignin, the high-density cross-linked structure formed in BC- CaCl_2 due to the addition of Ca^{2+} leads to a high density of 1.58 g/cm^3 (Figure S2A).

The mechanical tests show that the tensile strength of BC- CaCl_2 (470.5 MPa) is much higher than that of BC-E (206.7 MPa) and BC-P (263.5 MPa) (Figure 4D,F). Meanwhile, the flexural strength of BC- CaCl_2 (470.5 MPa) is also much higher than that of BC-E (206.7 MPa) and BC-P (263.5 MPa) (Figure 4G,I). In addition, the tensile modulus and flexural modulus of BC- CaCl_2 are higher than those of BC-E and BC-P (Figure 4E,H).⁷⁰ The results indicate that a stable high-density cross-linked structure is formed inside BC- CaCl_2 , which enhances the bonding strength between fibers. This discovery is attributed to the formation of coordination crosslinks between –COOH in hemicellulose and Ca^{2+} , which enhances the internal binding of BC- CaCl_2 under the combined action of intermolecular hydrogen bond. On the other hand, Figure 4J shows the excellent specific strength of BC- CaCl_2 , surpassing most typical structural materials, especially some metals and alloys.⁷¹ In summary, the excellent mechanical properties of BC- CaCl_2 provide a solid foundation for its application as a structural material in the building materials industry, such as anti-seismic buildings.

3.5 | Color difference and water resistance of BC- CaCl_2

The color difference experiment (Figure 5A) shows that Ba- CaCl_2 has a lower lightness than BC-E and BC-P. This indicates that BC- CaCl_2 has the highest degree of surface charring.⁷² This is because part of the lignin in BC- CaCl_2 is removed by the alkali solution, and the remaining lignin melts and overflows during the thermoforming process. However, its internal stable structure inhibits the recombination of lignin, so a dense protective layer is formed on

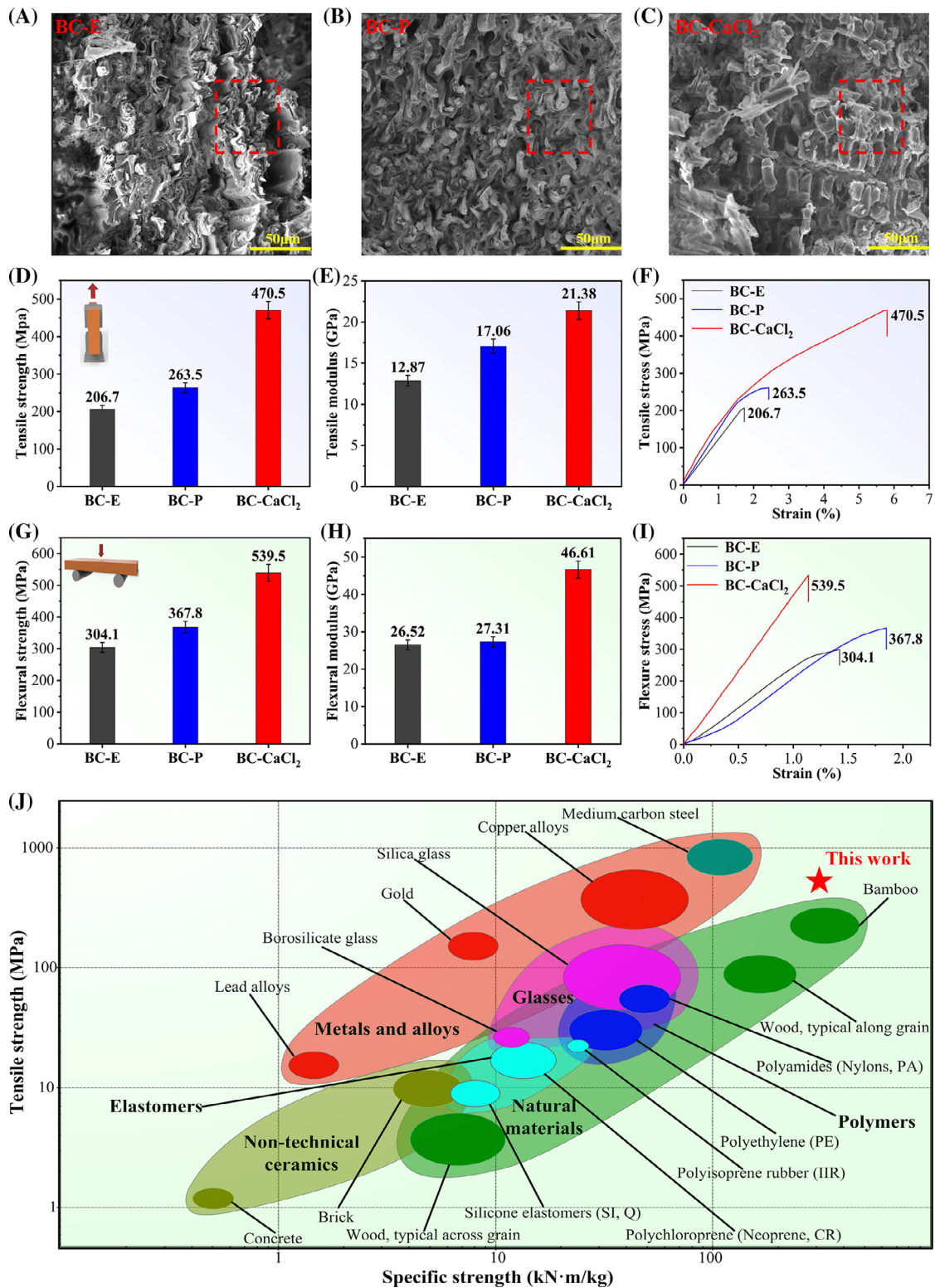


FIGURE 4 SEM images of the cross section of (A) BC-E, (B) BC-P and (C) BC-CaCl₂. (D) Tensile strength, (E) tensile modulus, (G) flexural strength and (H) flexural modulus of BC-E, BC-P and BC-CaCl₂. (F) Tensile and (I) flexural stress–strain curves of BC-E, BC-P and BC-CaCl₂. (J) Comparison diagram of tensile strength-specific strength between BC-CaCl₂ and other materials.

the surface of BC-CaCl₂ after cooling. The SEM observations show that the surface of BC-E and BC-P is full of cracks, while the surface of BC-CaCl₂ is flat and smooth,

making it difficult for moisture to enter the interior of the material (Figure S3).⁷³ This phenomenon is attributed to the protective layer structure formed by lignin.

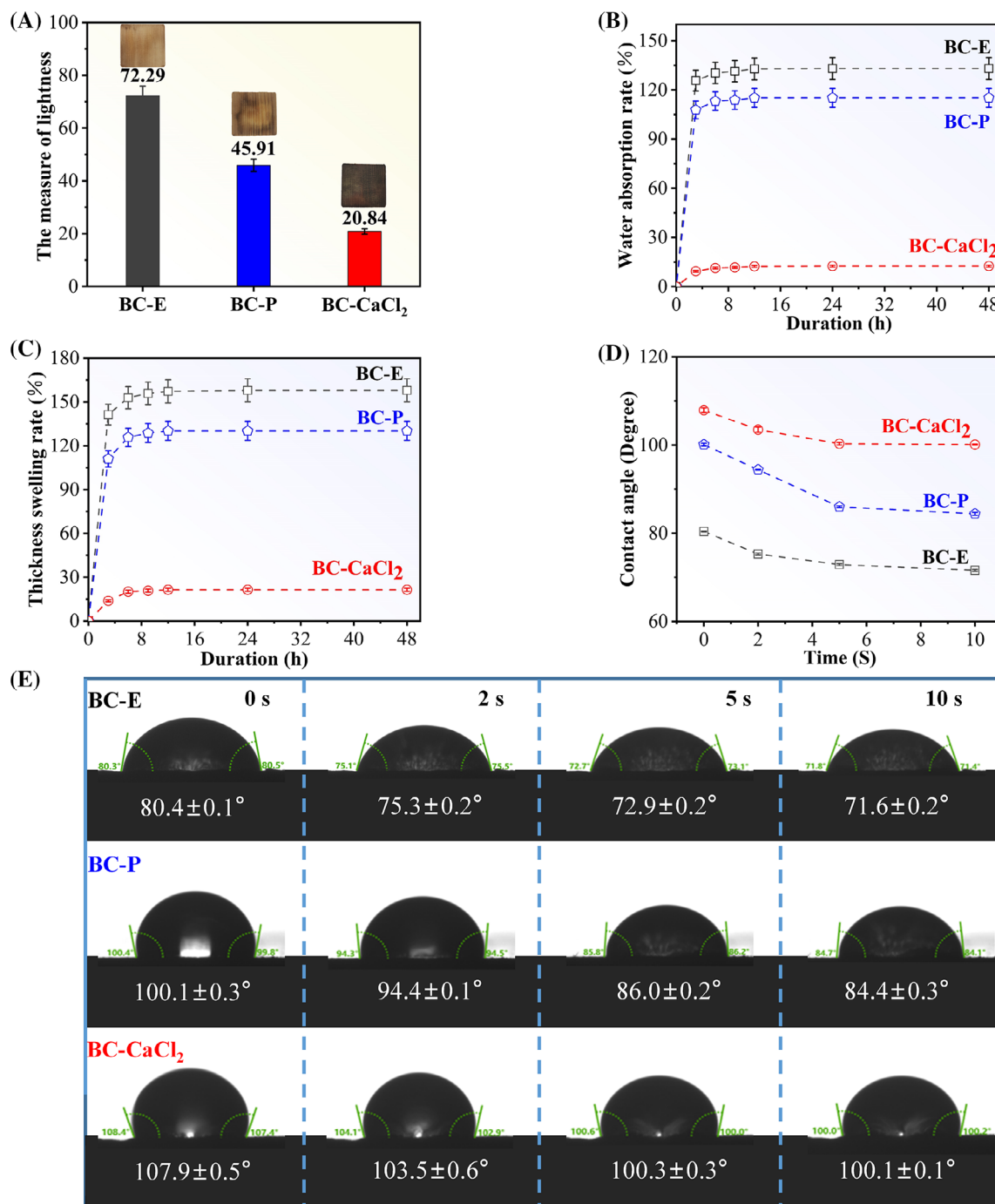


FIGURE 5 (A) Lightness comparison of BC-E, BC-P, and BC-CaCl₂. (B) Water absorption rate comparison of BC-E, BC-P, and BC-CaCl₂. (C) Water absorption thickness swelling rate comparison of BC-E, BC-P, and BC-CaCl₂. (D) Contact angle-time curve of BC-E, BC-P, and BC-CaCl₂. (E) Contact angle photographs of BC-E, BC-P, and BC-CaCl₂ at 0, 2, 5, and 10 s.

Furthermore, the stable building blocks structure inside BC-CaCl₂ also hinders the absorption of water (Figure 4C). Therefore, the water resistance test of BC-E, BC-P, and BC-CaCl₂ soaked in water for 2 days shows that the water absorption rate and water absorption thickness expansion rate of BC-CaCl₂ (12.48%, 21.38%) are much lower than those of BC-E (133.05%, 157.97%) and BC-P (115.12%, 130.15%) (Figure 5B,C).

Meanwhile, the contact angle test shows that BC-CaCl₂ has the highest contact angle and is stable above 100° (Figure 5D,E),⁷⁴ showing that BC-CaCl₂ has the best water stability. This property is attributed to the reduction of hydrophilic group –OH under the treatment conditions of alkaline solution, which corresponds to the FT-IR results (Figure 3G). Meanwhile, the protective layer formed by lignin on the surface of the materials

makes it difficult for water to penetrate. Furthermore, the high-density cross-linked structure inside BC-CaCl₂ can improve its dimensional stability, thereby greatly enhancing its water resistance. Hence, BC-CaCl₂ can be used in indoor areas with high humidity, such as toilets, bathrooms and kitchens as well as in the envelope structure of outdoor buildings to prevent the infiltration of rain, snow and groundwater.

3.6 | Thermal conductivity and thermal stability of BC-CaCl₂

Thermal conductivity tests show that the thermal conductivity of BC-CaCl₂ is higher than that of BC-E and BC-P (Figure 6A,B). Meanwhile, the infrared thermal

imaging experiment shows that the thermal diffusion degree of BC-CaCl₂ is the largest within the heating time of 3 min, which is consistent with the result of the highest thermal conductivity (Figure S4). This is because Ca²⁺ is a benign conductor of heat. It collides with free electrons in BC-CaCl₂, and the unbound free electrons move freely inside the material, which can conduct heat. Therefore, the excellent thermal conductivity makes BC-CaCl₂ suitable for indoor floor heating systems and heat dissipation of high-power devices such as communication in place of thermal conductive rubber.⁷⁵

The thermal stability of BC-CaCl₂ and other materials is evaluated by TGA analysis (Figure 6C). Three stages of pyrolysis in air are observed, that is, dehydration, oxidative pyrolysis and carbonization stages. The thermal degradation of BC-CaCl₂ mainly occurs at about 230–400 °C,

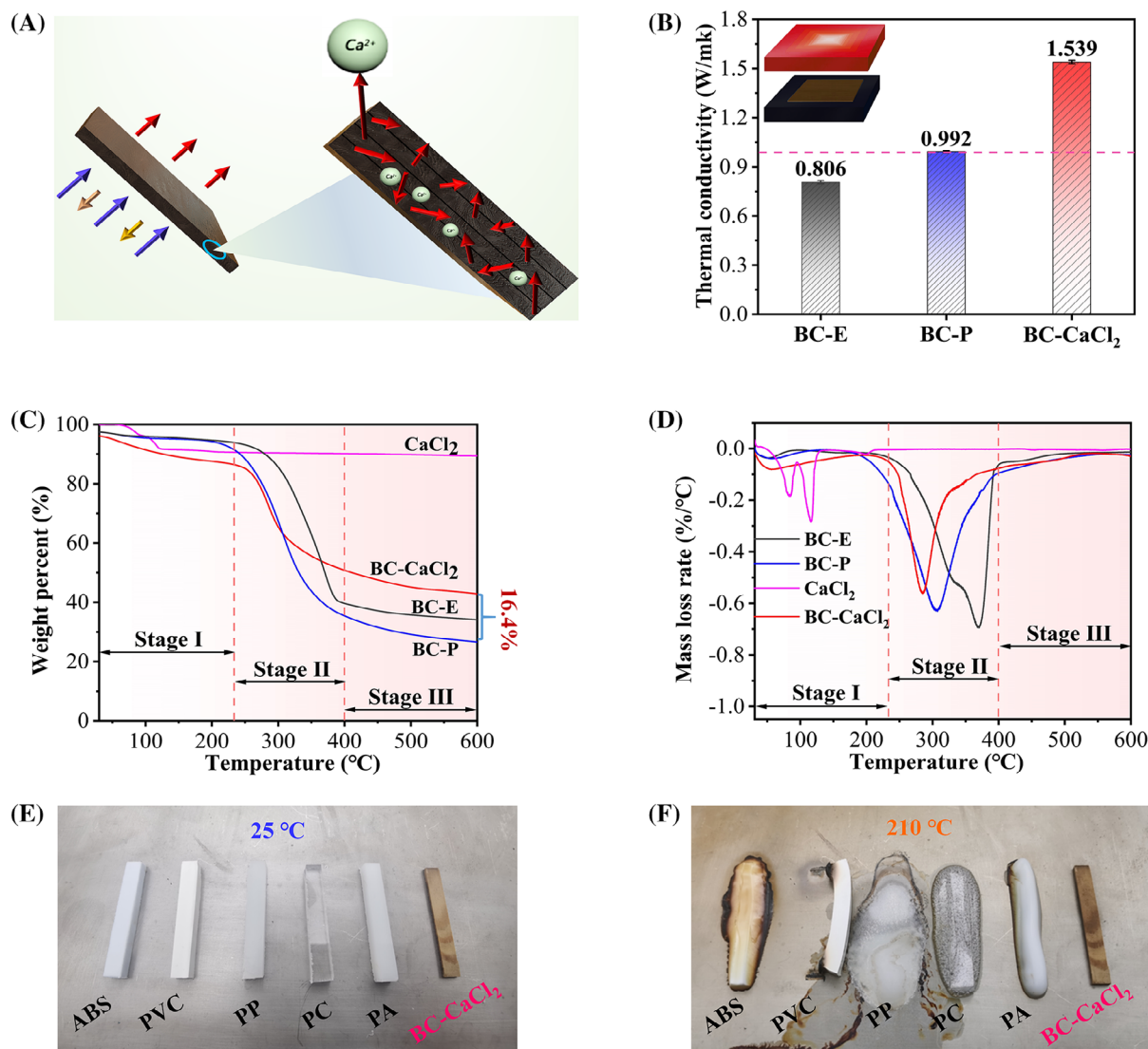


FIGURE 6 (A) Schematic diagram of thermal conductivity mechanism of BC-CaCl₂. (B) Thermal conductivity comparison of BC-E, BC-P, and BC-CaCl₂. (C) TG and (D) DTG curves of BC-E, BC-P, BC-CaCl₂, and CaCl₂. (E) Comparison diagram of thermal stability of BC-CaCl₂ and several normal used plastics at 25 °C. (F) Comparison diagram of thermal stability of BC-CaCl₂ and several normal used plastics at 210 °C.

which is the degradation stage of cellulose and hemicellulose. Since crystal CaCl_2 is not easily thermally degraded, BC- CaCl_2 has the lowest mass loss and the final residue is 16.4% more than that of treated BC-P. However, the maximum degradation temperature of BC- CaCl_2 is lower than that of BC-E and BC-P. This is due to the formation of coordination crosslinks between $-\text{COOH}$ in hemicellulose and Ca^{2+} . During the heating process, ions are not bound but move freely, making the internal structure of material unstable.

Meanwhile, the DTG curve (Figure 6D) shows that the maximum degradation rate corresponding to the maximum degradation temperature of BC- CaCl_2 is the lowest.⁷⁶ Moreover, the weight loss temperature of BC- CaCl_2 between 230 and 300°C is lower than that of BC-E and BC-P due to the decrease of lignin content. Hence, this property can be applied to develop materials with good thermal insulation properties, such as insulation materials in construction. The combustion experiment after 10 s shows that BC-E and BC-P emit strong flames and spread rapidly while BC- CaCl_2 is hardly ignited (Figure S5a). In addition, most of BC-E and BC-P have been charred and turned black after 10 s, while BC- CaCl_2 still maintains its structural integrity (Figure S5b).⁷⁷ It can be directly observed from Figure 6E,F that BC- CaCl_2 can maintain the initial stable shape on the heating stage at 210°C, while other commonly used plastics have all softened and deformed.⁷⁸ The above results show that BC- CaCl_2 has excellent thermal stability, compared to existing petrochemical plastics. This is because lignin forms a dense protective layer on the surface of BC- CaCl_2 , which hinders the flame

combustion of the material. Therefore, good thermal stability of BC- CaCl_2 makes it possible to replace petrochemical-based materials, such as plastics for pipes and packaging materials. It further tackles the pollution problem that waste plastics are difficult to handle from the source, which is in line with the concept of clean production.

3.7 | UV resistance, corrosion resistance and antibacterial properties of BC- CaCl_2

Figure 7A shows a schematic diagram of the UV resistance, corrosion resistance and antibacterial properties of BC- CaCl_2 . BC- CaCl_2 absorbs almost all ultraviolet light (94%), including UVA (315–400 nm), UVB (280–315 nm), and UVC (200–280 nm) (Figure 7B). Meanwhile, the UV transmittance of BC- CaCl_2 is the lowest meaning that BC- CaCl_2 can completely shield UV radiation. This is because the surface of BC- CaCl_2 is covered with a protective layer composed of lignin, and the lignin molecules contain a large number of phenylpropane units with phenol hydroxyl groups, which can strongly absorb UV light. The excellent UV resistance enables BC- CaCl_2 to replace the UV resistant fiber for weaving outdoor products such as sunshades and outdoor UV resistant polymers.⁷⁹

The antibacterial test after 24 h shows an obvious inhibition zone around BC- CaCl_2 , while the rest of the samples had no obvious changes (Figure 7C, S6A).⁸⁰ The phenomenon demonstrates excellent antibacterial properties of BC- CaCl_2 because Ca^{2+} will affect the metabolism of microorganisms.^{81,82} This makes BC- CaCl_2

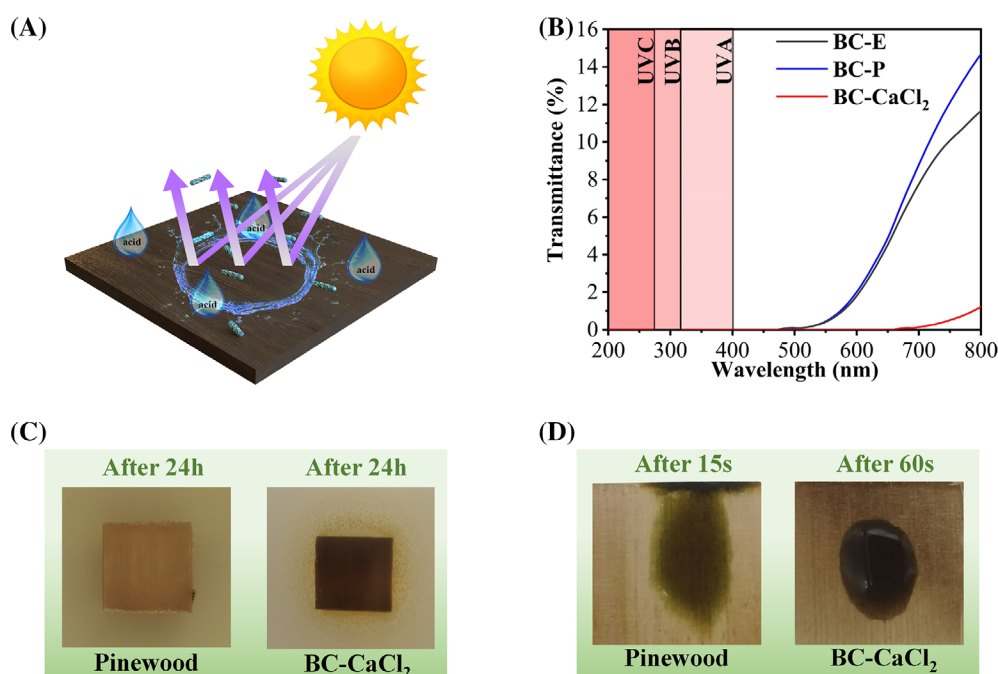


FIGURE 7 (A) Schematic diagram of UV, antibacterial and corrosion resistance of BC- CaCl_2 . (B) UV transmittance curves of BC-E, BC-P, and BC- CaCl_2 . (C) Comparison of antibacterial resistance between pinewood and BC- CaCl_2 . (D) Comparison of acid resistance between pinewood and BC- CaCl_2 .

suitable for food and pharmaceutical packaging. In addition, BC-CaCl₂ has a good corrosion resistance due to its excellent water resistance, which prevents sulfuric acid from entering the interior of the material (Figure 7D, S6B). Coupled with the outstanding mechanical properties of BC-CaCl₂, it can be used as an anti-corrosion material in comparison to stainless steel in the field of installation engineering.^{83–85}

4 | CONCLUSIONS

This study develops an efficient process for facile fabrication of high-performance, sustainable wood-based structural functional materials BC-CaCl₂, based on natural pinewood. Ca²⁺ was used to build a high-density cross-linked structure in pinewood, including high-density hydrogen bonds between cellulose microfibril and hemicellulose, and coordination crosslinks between Ca²⁺ and –COOH. The stable internal structure endows BC-CaCl₂ with outstanding tensile strength (470.5 MPa) and flexural strength (539.5 MPa), providing structural advantages for the application of the material. Moreover, BC-CaCl₂ exhibits excellent water resistance, thermal conductivity, thermal stability, UV resistance, corrosion resistance and antibacterial properties. The prepared high-performance wood-based composites address the inefficient use of pinewood and achieve the high-value utilization of natural wood. Meanwhile, the simple preparation process of one-step thermoforming is conducive to the further development and practical application of wood-based composites in the field of structural and functional materials. The potential application areas for BC-CaCl₂ include waterproof enclosure structure of buildings, indoor underfloor heating, outdoor UV resistant protective cover and anti-corrosion materials for installation engineering, and so forth. The as-prepared BC-CaCl₂ may have the potential to replace traditional petrochemical-based materials for a wide range of applications, such as replacing plastics for pipelines and packaging materials. However, wood-based structural functional materials often require special treatment and protection to improve their performance and durability. Future research will improve the durability of the material by applying protective layers. Due to the limitations of hot-pressing technology and solid wood as raw materials, the material is also subject to processability limitations in practical applications. Hence, multiple molds can be developed in the future to suit a variety of application scenarios.

AUTHOR CONTRIBUTIONS

Haoran Ye and Yang Shi carried the experimental work and data processing, Wei Fan, Zhongfeng Zhang, Daniel

M. Mulvihill and Pengju Shi helped in data analysis, Ben Bin Xu design the work program with Zhanhu Guo, Xuehua Zhang, Ximin He and Shengbo Ge. All authors contributed in the manuscript drafting.

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CONFLICT OF INTEREST STATEMENT

We confirm that the work submitted is not under consideration elsewhere. Prof Ximin He is an Editorial Board member of EcoMat and a co-author of this article. To minimize bias, she should be excluded from all editorial decision-making related to the acceptance of this article for publication. Otherwise, no further conflict of interest in this submission.

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SUPPORTING INFORMATION

Additional supporting information can be found online in the Supporting Information section at the end of this article.

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