Solar-assisted fabrication of large-scale, patternable transparent wood

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Transparent wood is considered a promising structural and light management material for energy-efficient engineering applications. However, the solution-based delignification process that is used to fabricate transparent wood generally consumes large amounts of chemicals and energy. Here, we report a method to produce optically transparent wood by modifying the wood’s lignin structure using a solar-assisted chemical brushing approach. This method preserves most of the lignin to act as a binder, providing a robust wood scaffold for polymer infiltration while greatly reducing the chemical and energy consumption as well as processing time. The obtained transparent wood (~1 mm in thickness) demonstrates a high transmittance (>90%), high haze (>60%), and excellent light-guiding effect over visible wavelength. Furthermore, we can achieve diverse patterns directly on wood surfaces using this approach, which endows transparent wood with excellent patternability. Combining its efficient, patternable, and scalable production, this transparent wood is a promising candidate for applications in energy-efficient buildings.

INTRODUCTION

The growing demand of optical components in energy-efficient buildings (1–3), optoelectronics (4–6), solar cells (7), and solar desalination devices (8) has fueled the search for high-performance, sustainable optical materials. Toward this aim, transparent wood has emerged due to its unique hierarchical structure (9–10), high specific strength (11–12), and favorable light management properties (13–14), demonstrating a broad range of applications including optoelectronic devices (15–18), energy-efficient building materials (9, 19–20), light management layers for solar cells (21–22), and touch panels (23–24). To fabricate transparent wood, two steps are typically involved: (i) completely removing the light-absorbing lignin from the cell walls of natural wood by a solution-based immersion method and (ii) infiltrating a refractive index matching polymer into the delignified wood matrix to minimize light absorption and scattering, respectively (9, 13, 25–27). However, this solution-based delignification process impairs the mechanical strength of wood and is also relatively energy and chemical intensive. Furthermore, it is necessary to fully remove the dispersed lignin from the wood scaffold, which lengthens the processing time (6 to 12 hours) (9–10, 18, 28–29). Last, the generated liquid waste is difficult to recycle and creates an additional environmental burden.

To address these processing issues as well as obtain desirable optical and mechanical properties, recent studies have begun to optimize the preparation of transparent wood by controlling the degree of delignification. For example, Qin et al. (30) prepared a laminated transparent wood of 1.5 mm in thickness (~82% optical transmittance) using a two-step partial delignification method (NaClO2/H2O2), in which 80% of lignin was removed and the processing time was reduced to 3 hours. Meanwhile, Wu et al. (31) prepared a transparent wood (~61% optical transmittance) from basswood (0.42 mm in thickness) with improved mechanical properties by reducing the degree of delignification (75% lignin content was removed) using a NaClO2 solution method. Although these partial delignification methods reduce the processing time and improve the mechanical characteristics of transparent wood, they require toxic chemicals that are difficult to dispose, as well as large solution volumes to enable complete immersion of the materials. In addition, further simplifying the reaction process with even shorter processing time would improve the scalability and efficiency of transparent wood production.

We note that the chromophore groups in lignin are mainly responsible for the brownish color and light absorption properties of natural wood (32). Recently, researchers have found that lignin can be decolorized by altering the chromophoric composition (33–36) rather than removing the lignin structure entirely. For example, Li et al. (36) reported a transparent wood that retained its lignin content and instead removed the lignin chromophore using an alkaline H2O2 hydrothermal solution to reduce the processing time. H2O2 as a desirable oxidant produces only water as a by-product, which notably reduces the waste liquid production. However, like previous techniques, this method requires relatively large chemical, water, and energy consumption. Therefore, it has remained challenging to process transparent wood in a fast, scalable, low cost, and environmentally friendly way.

Here, we report on a technique for rapidly fabricating transparent wood that involves chemical brushing, rather than immersion, combined with solar illumination to remove the lignin chromophore. Figure 1A shows the fabrication process of the transparent wood via a two-step approach. First, we modify the lignin structure by brushing H2O2 across the wood surface, followed by ultraviolet (UV) light illumination that can be conducted using natural sunlight to remove the light-absorbing chromophores of lignin. Refractive index matching epoxy can then be easily infiltrated into the microporous wood structure to prepare the transparent wood, which features a dense microstructure and low light scattering. The hierarchically porous structure of natural wood promotes fast H2O2 solution infiltration/diffusion and UV light trapping to efficiently remove the light-absorbing chromophore of lignin, significantly reducing its light absorption (<4%). Compared to lignin-removed wood (0.4 MPa), the lignin-modified wood also shows a substantially higher tensile strength (20.6 MPa) due to the presence of the modified lignin.
binding with the well-oriented cellulose fibrils, which provides a strong and robust scaffold for polymer infiltration. Note that transparent wood can be made from both transversely and longitudinally cut natural wood, respectively (Fig. 1, B and C). The resulting transparent wood exhibits a high transmittance (up to ~90%), excellent tensile strength (> 46 MP), and favorable light guiding effect. In addition, the transparent wood can be easily patterned (Fig. 1C) using this chemical brushing combined with UV light illumination method. Compared with traditional solution-based immersion processes, our method requires fewer chemicals and energy consumptions, thus greatly reducing the preparation cost and liquid waste. This environmentally friendly, scalable, patternable, and low-cost transparent wood with favorable optical and mechanical properties holds great potential in energy-efficient building applications and light management devices.

RESULTS AND DISCUSSION
Preparation of transparent wood and its structural characteristics
Figure S1 shows the preparation process of transparent wood via this simple yet effective approach. First, ~15 ml of H$_2$O$_2$ [30 weight % (wt %) concentration] was brushed on a natural balsa wood sample (200 mm by 10 mm by 0.6 mm), followed by exposure to UV light (a UV lamp was used as the solar UV light simulator) for 1 hour until the natural wood color turned completely white. The resulting lignin-modified wood displays a low absorptivity (<4%) (fig. S2) and a high reflectivity (>88%) (fig. S3) in the range of 400 to 800 nm, suggesting the removal of the light-absorbing chromophores in lignin. Meanwhile, in the Fourier transform infrared (FTIR) spectrum of the lignin-modified wood, we attribute the absorption peaks at approximately 1595, 1505, and 1435 cm$^{-1}$ to the aromatic vibrations...
of lignin (Fig. 2A) (37–38), indicating that the process preserves the aromatic backbone of lignin despite degrading its chromophore. The peak of 1734 cm⁻¹ is assigned to the carboxyl groups in hemicellulose (xylan/glucosannan). The peak of 1235 cm⁻¹ belongs to the uronic acid groups of the hemicellulose or the ester linkage of the carboxyl groups of lignin and hemicellulose (18, 25, 39). The disappearance of 1734 cm⁻¹ peak and the decrease of peak intensity of 1235 cm⁻¹ in lignin-modified wood indicate the partial dissolution/removal of hemicellulose from natural wood after treatment. In addition, the lignin content of the natural wood and lignin-modified wood samples was ~23.5 and ~19.9%, respectively (Fig. 2B), which further suggests that the most of lignin structure was well preserved after the treatment.

We note that the preserved lignin can act as a binder to strengthen the mechanical properties of the lignin-modified wood as well as provide a robust wood scaffold for subsequent polymer infiltration (40–41). The lignin-modified wood exhibits a tensile strength of 20.6 MPa under wet conditions, which is ~50 times higher than lignin-removed wood (0.4 MPa) prepared by the traditional NaClO₂ solvent-based method (fig. S4). In addition, the lignin-modified wood shows excellent flexibility along the longitudinal (parallel to the fiber direction, L) and transverse directions (perpendicular to the fiber direction, T) (fig. S5). In contrast, the lignin-removed wood prepared by the traditional NaClO₂ solvent-based method is very fragile and easily cracked (fig. S6). Benefiting from the strong mechanical properties, we further prepared a large lignin-modified wood sample with a length of ~1 m using this chemical brushing combined with UV light illumination method (Fig. 2C).

Using this lignin-modified wood, we then infiltrated it with epoxy by vacuum to obtain the final transparent wood product. Figure 2 (D to I) shows scanning electron microscopy (SEM) images of the natural wood, lignin-modified wood, and transparent wood. The natural wood shows a three-dimensional hierarchical and interconnected porous structure (Fig. 2, D and G, and fig. S7), featuring microchannels with diameters that range from ~15 to 300 μm. This unique porous structure is beneficial for fast H₂O₂ solution infiltration/diffusion and efficient UV light trapping inside the wood microchannels, which allows us to achieve efficient removal of light-absorbing chromophores during the process (fig. S8). Meanwhile, the diameters of the microchannels in the lignin-modified wood range from 10 to 270 μm (Fig. 2, E and H), demonstrating that lignin-modified wood preserves the hierarchical, interconnected porous structure after treatment. Last, SEM images of the transparent wood show the epoxy is well infiltrated into the pores of the lignin-modified wood (Fig. 2, F and I, and fig. S9). A dense and compact structure is formed in the transparent wood after the infiltration, which helps suppress light scattering and improves the optical transmittance (11, 14).

**The optical and mechanical performance of transparent wood**

Transparent wood along T and L directions exhibits excellent optical properties. Images of the transparent wood in the L (Fig. 3A)
and T directions (Fig. 3B) are demonstrated, in which we can clearly see the background through the materials. In addition, we can facilely prepare thick transparent wood (L, 1.5 mm; T, 3.5 mm) with the excellent optical transparency using chemical brushing combined with UV light illumination method (figs. S10 and S11). We measured the optical transmittance of the natural wood and transparent wood from 200 to 2000 nm (Fig. 3C). The transparent wood along the L and T directions has a high optical transmittance of ~90% over the visible wavelength range (400 to 800 nm) compared to the transmittance of the natural wood (L < 6%, T < 36%).

The absorptivity of the transparent wood (near 0%) is also much lower than that of the natural wood (L < 83%) in visible wavelengths (Fig. 3D) due to the removal of the light-absorbing chromophore of lignin, which allows almost all visible light to pass through the transparent wood. Figure S12 shows the transparent wood exhibits a high transmittance haze of ~60 to 80% over the wavelength range of 400 to 800 nm. Transparent wood with excellent optical transparency can also be made from other wood species with different density, such as oak and poplar, suggesting the universality of this approach (fig. S13). In addition, the transparent wood features preserved vertically aligned microchannels, which allow light propagation to be guided along the channel direction. As shown in Fig. 3E, we used a 650-nm red single mode laser to perpendicularly illuminate the transparent wood along the L and T directions. Our results show the beam propagates along the direction of the wood channels, indicating the transparent wood has light guiding capacity and anisotropic optical transmittance (10, 14, 42–43).

We measured the mechanical properties of the natural wood and transparent wood at different tensile directions. The tensile strengths of the natural wood (stress along the L and T directions) were 24.5 and 0.7 MPa, respectively (Fig. 3F). Meanwhile, the tensile strengths of the L- and T-transparent wood samples were 46.2 and 31.4 MPa, respectively, which corresponds to an enhancement of 1.8 times and 44.8 times higher than the natural wood (L and T). The L- and T-transparent wood also have a significantly improved toughness of 0.93 and 1.64 MJ m⁻³ compared to the natural wood (L, 0.26 MJ m⁻³; T, 0.03 MJ m⁻³), as shown in Fig. 3G. The toughness of the L-transparent wood is lower than that of the T-transparent wood because of the smaller elongation at break of the L-sample (3.4% < 7.4%) (Fig. 3F). We attribute the improved mechanical strength of the transparent wood to its lignin-reinforced anisotropic structure as well as the infiltrated epoxy, the tensile strength of which is ~87 MPa (fig. S14). In addition, the low volume fraction of wood scaffold also contributes to the improved mechanical properties of transparent wood (fig. S15). Benefiting from the high mechanical strength, the transparent wood is also quite flexible (fig. S16). The transparent wood has both excellent optical properties (high optical transmittance, high haze, and light guiding effect) and excellent mechanical strength, which suggest its application in energy-efficient buildings and light management devices.

Fig. 3. The optical and mechanical properties of the natural wood and transparent wood. Digital images of the transparent wood along the (A) longitudinal direction (L; 400 mm by 110 mm by 1 mm) and (B) transverse direction (T; 70 mm by 30 mm by 1.5 mm). (C) The transmittance and (D) absorption of the natural wood and transparent wood (the volume fraction of lignin-modified wood scaffold: L, ~30%; T, ~33.2%). (E) Guided light propagation by the transparent wood. Photo images of the scattered laser light spot of the L- and T-transparent wood materials. (F) The tensile strength and (G) toughness of the natural wood and transparent wood. Photo credit: Qinqin Xia, University of Maryland, College Park.
Optically transparent wood with excellent patternability

Traditional solution-based delignification methods generally involve entirely immersing the wood samples in chemical solutions, which makes it difficult to bleach selective areas of the material. In contrast, our chemical brushing combined with UV light illumination method allows us to selectively bleach designated areas of the wood samples, which enables us to easily prepare transparent wood with diverse patterns (movie S1). The transparent wood can be selectively and precisely patterned by taking advantage of the different optical transmittance between natural wood (6 to 36%) and transparent wood (~90%), as shown in fig. S17. First, the desired patterns drawn with a brush on the natural wood samples using H₂O₂ as an “ink.” Then, these regions were illuminated with UV light, which turns them white. We then infiltrated epoxy into the microchannels of the lignin-modified wood to obtain transparent wood with desired patterns (Fig. 4, A and B, and fig. S18). T-transparent wood samples with number (“4”) and letter (“A”) patterns are shown in Fig. 4 (C and D). In this case, we wrote the number 4 in the brownish color of the natural wood, which was defined by the patterned transparent area, while the transparent letter A was patterned in the reverse. We also demonstrated L-transparent wood samples with more complex geometries, including two transparent circles minus a nontransparent diamond and a yin-yang symbol (Fig. 4, E and F), showing transparent wood with arbitrary patterns can be achieved using our chemical brushing combined with UV light illumination method. Figure 4 (C to F) shows that the patterned transparent wood can be made from transversely and longitudinally cut natural wood, respectively, suggesting the high versatility in direction selection and pattern designability of our method.

Demonstration of solar-assisted fabrication of transparent wood

UV light (100 to 400 nm) on Earth originates from the Sun (an inexhaustible source), with more than 95% of the wavelengths that reach the planet’s surface being in the UVA range (315 to 400 nm) (44). Thus, we use this solar UV light as a driver of wood decoloration to achieve the rapid fabrication of transparent wood. The schematic diagram in Fig. 5A demonstrates the potential large-scale fabrication of transparent wood based on the mature rotary wood cutting method in wood industry and the solar-assisted chemical brushing process. The lignin-modified wood can be continuously and rapidly fabricated in this eco-friendly and energy-saving solar-driven manner for further scalable fabrication of transparent wood with high reliability and productivity. As shown in Fig. 5B, three large pieces of balsa wood with a length of 1 m can be rapidly produced in 1 hour by solar UV radiation [Global Solar UV Index: 7 to 8, Maryland, College Park (latitude: 39.00; longitude: −76.75) on 9 August 2019 at 13:00]. In our lab scale production, we obtained 400 mm by 110 mm by 1 mm of the transparent wood with high transmittance (Fig. 5C) after infiltrating epoxy into above lignin-modified wood.

We further evaluated the energy consumption, cost, and chemical emission of this production process for transparent wood and compared it with the NaClO₂ solution-based delignification methods (Fig. 5D). Our solar-assisted chemical brushing process only removes the chromophores of lignin and does not involve completely breaking the covalent bonds between cellulose and lignin. Thus, our method neither involves extensive usage of chemicals through harsh reaction nor high temperature treatment. Other solution-based methods generally require samples to be entirely...
immersed into adequate chemicals for an effective reaction, causing large amounts of chemical consumption and waste disposal. From the energy/chemical consumption point-of-view, the transparent wood fabricated by solar-assisted chemical brushing approach features greatly reduced energy (solar versus electrical heating) and chemical costs (3.14 versus 7.31 $/kg) compared to the lignin-removed transparent wood. From the waste emission point-of-view, our transparent wood also generates much less waste liquid (30 versus 960 ml) and no toxic gas (0 versus 960 ml). Thus, compared with the traditional solution-based delignification methods, the solar-assisted chemical brushing technique can produce transparent wood with the following advantages (Fig. 5E): (i) fast, low-cost, sustainable, and scalable fabrication; (ii) high lignin retention (~81%); and (iii) patternable transparent features.

In summary, we demonstrate a rapid, cost-effective, and sustainable method to fabricate patternable transparent wood based on a scalable solar-assisted chemical brushing method. This process, the light-absorbing chromophore groups of lignin were removed, which allowed us to improve the optical properties of the resulting transparent wood without destroying the aromatic structure of lignin entirely. Both the longitudinally and transversely cut transparent wood with a thickness of ~1 mm show a high tensile strength of 31.4 to 46.2 MPa, excellent optical transmittance of >90%, low optical absorption of <4%, and high optical haze of 60 to 80%. This transparent wood demonstrates excellent optical properties without significantly sacrificing the material’s mechanical strength due to the high lignin preservation. Moreover, the solar-assisted chemical brushing method can selectively treat designated areas of wood samples, imparting the transparent wood with excellent designable patterning capabilities. Compared to solution-based delignification processes, our solar-assisted chemical brushing has higher production efficiency, lower cost, and is more sustainable and controllable. This inexpensive and highly efficient fabrication of transparent wood can also use natural solar energy, expanding the technique’s application to large-scale industrial production.

**MATERIALS AND METHODS**

**Materials and chemicals**

American balsa wood with the density of 0.15 to 0.26 g/cm³ was purchased from Specialized Balsa Wood LLC. Sodium hydroxide substitute.
(NaOH) (Carolina Biological Supply Company), hydrogen peroxide (H2O2, 30% solution; EMD Millipore Corporation), and UV lamp (380 to 395 nm of wavelength; China) were used for lignin modification of the balsa wood. The epoxy resin (#300 resin and #21 Non Blushing Cycloaliphatic Hardener, AeroMarine Products Inc.) was further used for infiltration.

**Fabrication of the transparent wood**

A balsa wood log was cut along the transverse and longitudinal directions to form wood slices (0.6 to 3.3 mm in thickness). Then, the wood slices were brushed H2O2 (30 wt %), followed by UV illumination until the samples became completely white. We used a UV lamp (380 to 395 nm of wavelength) to simulate UV light in solar radiation. Note that a trace amount of NaOH (10 wt %) was coated on the wood surface before brushing H2O2 to improve the oxidation efficiency of the H2O2. This process removes the chromophores in lignin, causing the color of the wood to change from brown to white. The treated wood pieces were then immersed in ethanol for 5 hours to remove any remaining chemicals and then transferred to toluene so as to exchange the ethanol in the wood. Later, the samples were impregnated with epoxy (AeroMarine 300/21 epoxy) by vacuum infiltration for 1.5 hours. Last, the epoxy-impregnated wood samples were stored at room temperature until the epoxy was completely cured.

**Fabrication process of the lignin-removed wood**

NaClO2 was added into water to prepare a 5% solution, followed by the addition of acetic acid to change the pH to ~4.6. The natural wood was immersed in this boiling NaClO2 solution for 8 hours until the samples became completely white.

**Characterization**

The morphologies of the natural wood, lignin-modified wood, and transparent wood were observed on an SEM (Hitachi SU-70). FTIR spectra were recorded using a Nexus 670 FTIR Spectrometer over the range of 650 to 4000 cm⁻¹. The x-ray photoelectron spectroscopy (XPS) experiments were calibrated by the binding energy of C1s as a reference of 284.6 eV. Eclipse V2.1 data analysis software supplied by the VG ESCA-Lab200I-XL instrument manufacturer was applied in the manipulation of the acquired XPS spectra.

The content of lignin (Klason lignin) in wood samples was measured according to TAPPI method (45). First, 0.1-g lignin-modified wood was reacted with 1.5 ml of 72% H2SO4 at room temperature with stirring for 2 hours. Second, the solution was diluted with deionized water to a 3% H2SO4 concentration and refluxed for 4 hours. Last, the mixture was filtered by sand core funnel, and the acid-insoluble lignin was determined by gravimetric analysis. The percentage of acid-insoluble lignin presents the content of the preserved modified lignin. Three samples of each material were tested to obtain the averaged values.

The optical spectra, including absorption, transmission, and haze, were measured by a UV-Vis Spectrometer Lambda 35 (PerkinElmer, USA). An integrated sphere was used to collect the reflected and transmitted light. The dimensions of the natural wood and lignin-modified wood used for optical test were approximately 60 mm by 60 mm by 1 mm. The dimensions of transparent wood along the longitudinal and transverse directions used for optical test were approximately 60 mm by 60 mm by 1 mm, respectively. Note that multiple samples of transparent wood, lignin-modified wood, and natural wood were tested.

**Mechanical tensile testing**

The tensile properties of the natural wood, lignin-modified wood, lignin-removed wood, and transparent wood samples were measured using a Tinius Olsen H5KT tester. Five experiments were carried out for each sample. The dimensions of the samples were approximately 50 mm by 5 mm by (0.8 to 1.4) mm. The samples were stretched along the longitudinal and transverse directions of the wood until they fractured with a constant test speed of 5 mm min⁻¹.

**The volume fraction of lignin-modified wood**

The volume fraction of lignin-modified wood in the transparent wood was calculated according to the following equations (46)

\[
V_f = \frac{W_f x \rho_c}{\rho_f}
\]

(1)

\[
\rho_c = \frac{W_f}{W_m} + \frac{W_m}{\rho_m}
\]

(2)

where \(V_f\) is the volume fraction of lignin-modified wood, \(\rho_c\) is the density of composite, \(\rho_m\) is the density of lignin-modified wood (~0.2 g/cm³), \(W_f\) is the weight fraction of lignin-modified wood, and \(W_m\) is the weight fraction of the epoxy.

**Supplementary materials**

Supplementary material for this article is available at http://advances.sciencemag.org/cgi/content/full/7/5/eabd7342/DC1

**References and Notes**


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