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Vivian Merk, Munish Chanana*, Sabyasachi Gaan and Ingo Burgert* Mineralization of wood by calcium carbonate insertion for improved flame retardancy

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Abstract: Wood can be considered as a highly porous, three-dimensional organic scaffold. It can be mineralized to create hierarchically structured organic-inorganic hybrid materials with novel properties. In the present paper, the precipitation of CaCO₂ mineral in Norway spruce and European beech wood has been studied by alternating impregnation with aqueous and alcoholic electrolyte solutions. Microstructural imaging by SEM and confocal Raman microscopy shows the distribution of calcite and vaterite as two CaCO₂ polymorphs, which are deposited deep inside the cellular structure of the wood. The confined microenvironment of the wood cell wall seems to favor a formation of vaterite, as visible by XRD and Raman spectroscopy. In view of a practical application, the mineralization of wood opens up ways for sustainable wood-based hybrid materials with a significantly improved fire resistance, as proven via pyrolysis combustion flow calorimetry and cone calorimetry tests. Beyond that, this versatile solute-exchange approach provides an opportunity for the incorporation of a broad range of different mineral phases into wood for novel material property combinations.

Keywords: calcite, calcium carbonate, cone calorimetry, fire retardant, hybrid material, mineral, mineralized wood,

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PCFC, polymorph, Raman spectroscopy, scanning electron microscopy, vaterite, wood modification

Introduction

Embedding of minerals into biological tissues is a very common process in nature, e.g. bone, enamel or nacre, and leads to hybrid materials with extraordinary properties due to the synergetic interplay of inorganic and organic components at various levels of hierarchy ranging from the molecular up to the mesoscale level (Mann et al. 1993; Fratzl and Weinkamer 2007; Dunlop et al. 2011). Similar to other organic matrices, the porous biopolymer network of wood is a remarkable platform for depositing inorganic matter in a compartamentalized micro-environment and the literature demonstrates that mineralized wood materials can be obtained with enhanced properties, which have a benefit from the structural integrity of natural wood.

From the plethora of minerals provided by nature, only few minerals have been incorporated via chemical strategies into the wood structure, yet. Examples are silica (Saka and Ueno 1997), titania (Hubert et al. 2010; Mahr et al. 2012) aluminosilicates (Dong et al. 2002), calcium phosphate (Tampieri et al. 2009) and iron oxide (Merk et al. 2014). Calcium carbonate (Merk et al. 2015) was also employed in wood mineralization as it is well known from biomineralization, geology, and industrial applications (Meldrum 2003). However, strategies for CaCO, mineralization of wood, as reported in the literature, often require a costly technical set-up for supercritical gases (Tsioptsias and Panayiotou 2011), or harmful precursors and reactions, which for example involve elemental calcium, methanol, and hydrogen (Klaithong et al. 2013). Furthermore, the main challenge of a solid wood treatment is to deposit minerals deep inside the wood specimen, avoiding an exclusive precipitation of the mineral at and directly below the wood surface.

In this paper, a simple mineralization strategy of wood will be reported, which involves a subsequental *in-situ* mineral formation based on a solution-exchange process. By employing alternating solution-exchange cycles wit CaCl₂ in ethanol and NaHCO₃ in water, a high mass uptake can be achieved, which can be suitable for flame-retardant

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applications, but also for other material properties related to the inorganic character of the created hybrid material.

Materials and methods

Mineralization: Blocks of Norway spruce (*Picea abies* (L.) Karst.) and European beech (*Fagus sylvatica* L.) ($1\times1\times1$ cm³, 5–15 sample replicas) sawn from a wood beam were conditioned at 20°C/65% RH. Calcium chloride dihydrate (Merck, EMSURE ACS Reag. PhEur, Darmstadt, Germany) and sodium hydrogen carbonate (Merck, EMSURE ACS Reag. PhEur, Darmstadt, Germany) were used without prior purification to prepare solutions in absolute ethanol (Scharlau, ACS, Reag. PhEur, Barcelona, Spain) and in deionized water. After vacuum-assisted impregnation in ethanolic CaCl₂·2 H₂O (c=1.0 mol I⁴) for 10 h, the wood samples were rinsed with ethanol followed by vacuum-assisted impregnation in aqueous NaHCO₃ (c=1.0 mol I⁴) for 14 h (pH≈8.2) and subsequent rinsing with deionized water. Having completed 1–4 reaction cycles, the mineralized wood samples were equilibrated at 20°C/65% RH to determine the mass gain and to avoid a transformation of the crystalline phases.

Light microscopy: Cross-cut wood specimens were observed under a Leica (Wetzlar, Germany) M165C stereomicroscope (3.2x objective) coupled to a Basler GigE Vision camera (Ahrensburg, Deutschland). Images were recorded with a control plugin (PHASE GmbH) for the software ImageJ 1.47c.

Environmental scanning electron microscopy (ESEM) was carried out on a FEI Quanta 600 (Eindhoven, The Netherlands) in the low-vacuum mode (water vapor, 0.53 Torr) driven by 20 kV acceleration using a SSD detector.

For the Raman measurements 20 µm cross-sections were prepared with a rotary microtome (Leica Ultracut, Wetzlar, Germany), placed on a microscopic slide with a few drops of D_2O and sealed with a cover slip and nail polish to avoid evaporation. All Raman spectra were recorded with a confocal Raman microscope (Renishaw InVia, Wotton-under-Edge, UK) equipped with a Nd-YAG laser (λ =532 nm), a 1800 grooves mm⁴ grating and a 100× oil immersion objective (Nikon, NA=1.4), a step width of 300 nm and an integration time of 0.30 s. The data were processed and scanned for cosmic rays with the software Wire 4.1. Single spectra were plotted with the program OriginPro 8.1 (Northampton, USA). Additionally, for the extraction of hyperspectral data, Vertex Component Analysis (VCA) was performed in the spectral range 900–1700 cm⁴ with the MatLab-based software CytoSpec v. 2.00.01 (Peter Lasch, Berlin, Germany) assuming six endmembers.

XRD spectra were collected from chopped mineralized wood (sample volume ~0.25 cm³) in triplicate on a Bruker D8 Advance (Karlsruhe, Germany) instrument ranging from 2θ =10–80° equipped with a copper source [λ (CuK α 1)=0.154 nm, 40 kV voltage, 40 mA current] and a rotating flat disk sample holder using a scan speed of 0.15° min³.

Pyrolysis combustion flow calorimetry (PCFC): The heat of combustion of wood/CaCO₃ composites and reference wood (conditioned at 20°C/65% RH) was determined by PCFC (Fire Testing Technology Instrument, East Grinstead, UK) according to (Lyon and Walters 2004) with a pyrolysis temperature of 85–750°C and a 80:20 N₂/O₂ gas mixture and operated at a heating rate of β =1 Ks⁴ and a combustion

temperature of 900°C. The PCFC measurements were replicated at least five times per sample with a weight of ca. 5 mg. The HRR curves were baseline-corrected and fitted with multiple Gauss curves using the software OriginPro 8.1. Herein, the resulting peak sum (the total heat release) displayed residual values close to 1. The maximum heat release divided by the constant heating rate β =1 Ks⁻¹ gives the heat release capacity. Samples were weighed before (m₀) and after (m₁) micro-calorimetry testing. The char yield is calculated from μ =(m₀-m₀)·100%/m₀.

In cone calorimetry testing (Fire Testing Technology Instrument, East Grinstead, UK), wood samples (five samples per variation) were exposed to a constant uniform irradiation of 50 kW m² (T~750°C) in the horizontal position according to the ISO 5660-1 norm. The heat release rate (HRR) was calculated based on the oxygen consumption. Bulk densities of the samples were determined at 20°C/65% RH prior to chemical modification. After a vacuum-drying step at 65°C, the samples (L≈100×R≈100, T≈10 mm³) were treated in four mineralization cycles, where the relative mass gain were 17±2% (spruce) and 18±2% (beech). Then the samples were washed with water for 30 min, dried in a vacuum kiln, and conditioned to 20°C/65% RH. The samples were mounted onto the loading cell in an aluminum boot and a standard metal frame.

FT-IR spectra of char residues obtained from cone calorimetry testing were collected with a Tensor 27 spectrometer (Bruker, Fällanden, Switzerland) equipped with a Pike ATR cell in a spectral range of 4000–350 cm⁻¹ with 4 cm⁻¹ spectral resolution and averaged over 100 scans. The spectra were recorded with the program OPUS and plotted with OriginPro 8.1.

Results and discussions

Calcium carbonate can be inserted deep into the wood structure by employing alternating solution-exchange cycles of highly concentrated electrolytes (Figure 1a) with CaCl₂ in ethanol and NaHCO₃ in water. The electrolytes contain ionic precursors that react instantaneously to anhydrous CaCO₃ polymorphs. Wood is incubated successively with salt solutions, thus CaCO₃ mineral formation is primarily governed by the high supersaturation.

A main advantage of a successive impregnation is the versatility regarding the mineral composition and the high degree of mineralization, which is easily tunable by the number of reaction cycles (Figure 1b,c). After four cycles, high relative mass gains (MG) of over 35% in spruce and 25% in beech (Figure 1b) were achieved, but it should be noted that besides $CaCO_3$ the samples most probably contain side products, such as NaCl, or unreacted salts in smaller amounts. For the small samples, the absolute MG (Figure 1c) is similar for spruce and beech, though they have different bulk density and cellular structure, and this reflects the versatility of this approach. Despite of the high total MGs in the samples, there is a MG gradient due to the fast reaction between ionic precursors, which is manifested by a higher mineral accumulation near the



Figure 1: (a) Scheme of mineralization *via* liquid exchange of alternating impregnations with $CaCl_2$ in ethanol and $NaHCO_3$ in H_2O . Averaged relative (b) and absolute mass gain (c) of cubic spruce and beech specimens (1 cm³ volume) after cycled solution-exchange process with 1 M $CaCl_2 \cdot 2 H_2O$ in ethanol and 1 M $NaHCO_3$ in H_2O .

surface readily visible as whitish $CaCO_3$ precipitate from the outside.

Both the mineral composition and the solvents can be varied. The initial infiltration is slightly better, when CaCl₂ is supplied as alcoholic instead of water solution (data not shown), because this approach allows trapping the NaHCO₃ precursor salt in the wood structure by taking advantage of its low solubility in ethanol. The mineral precipitation occurs spontaneously and quantitatively due to its poor

solubility in both solvents. After four solution-exchange cycles, however, the mineral uptake reaches the saturation level irrespective of the employed solvent. Hence, a purely water-based mineralization can also be envisaged from the environmental and economic point of view.

Light microscopy at low magnification shows large amounts of mineral precipitates in large cavities, namely spruce earlywood (EW) tracheids and beech vessels (Figure 2a,d). Closer observation with SEM reveals the



Figure 2: Light microscope and SEM images of $CaCO_3/wood$ composites obtained in the backscattered electron mode, beech (a–c) and spruce (d–f) cross sections. The scale bar corresponds to 500 μ m (a, d) and 20 μ m (b, c, e, f).

prevailing deposition of polycrystalline $CaCO_3$ in the lumina of beech vessels and fibers (Figure 2b,c) or spruce tracheids (Figure 2e,f). The deposition pattern of $CaCO_3$ mineral seems to coincide with the microporous structure that facilitates fluid transport in wood.

CaCO₃ precipitates in rays (Figure 2b,e) or in interconnecting pit channels of neighboring beech fiber cells (Figure 2a–d) indicate, in addition to that in micropores of the axial tissues, that an ion transport occurs perpendicular to the wood fiber direction. Furthermore, in the case of beech, high amounts of electron-lucent material are also visible in the cell corners and the compound middle lamellae (CML) of the fibers (Figure 2b,c). This might be either explained by the adsorption of calcium cations to negatively charged and metal-complexing surfaces within the cell wall regions rich in pectin and lignin (Grant et al. 1973; Jarvis 1984; Cooper 1998; Petric et al. 2000; Caffall and Mohnen 2009) or by an increased ion transport to the CML from the lumen across the pit membranes (Cooper 1998; Petric et al. 2000; Gindl et al. 2003).

The mineral distribution in the wood structure differs substantially from the mineralization in a one-pot method (Merk et al. 2015). In the quoted work, a neutral molecule (dimethyl carbonate) served as carbonate source and was hydrolyzed to CO_2 upon adding excess base. It was observed that $CaCO_3$ was predominantly formed in the cell walls and the lumina were hardly filled. From the present work, it can be assumed that capillary effects of pores in the cell walls are involved in controlling the deposition pattern.

Judging from the mineral morphology, in the first instance, the CaCO₃ precipitates seem to be constituted mainly of calcite. However, a closer look reveals that the resultant mineralization product is a mixture of calcite and vaterite according to X-ray powder diffraction (Figure 3) and Raman spectroscopy (Figure 4) both in spruce or beech. Under the present reaction conditions (pH≈8.2), the initially formed metastable vaterite is expected to gradually transform into calcite (Spanos and Koutsoukos 1998; Sheng Han et al. 2006; Bao et al. 2011). Vaterite, however, is remarkably stable in dry wood samples over several months, which might be explained by the effect of confined spaces in the wood microstructure.

Figure 4 shows confocal Raman microscopic mappings of $CaCO_3/wood$ composites. Vertex component analysis (VCA) is a powerful multivariate method which allows for discerning the polymeric wood cell wall components and different $CaCO_3$ mineral polymorphs, as described elsewhere (Gierlinger 2014; Keplinger et al. 2015; Merk et al. 2015). The respective endmember mappings show the distribution of carbohydrates in the S2 cell



wall layer (Figure 4a,e,i,m) and different CaCO₂ phases (Figure 4b,c,f,g,j,k,n,o). On the basis of Raman signals, the calcitic and vateric form of CaCO₃ can be easily distinguished. Calcite gives rise to an intense v_1 vibration band at 1085 cm⁻¹ (Rutt and Nicola 1974), whereas the v_1 mode of vaterite splits into a doublet at 1089 cm⁻¹ and 1074 cm⁻¹ (Behrens et al. 1995; Kontoyannis and Vagenas 2000). A smaller portion of CaCO₃ is localized within the cell walls and along the CML (Figure 4b, f, j, n). In the nanoporous wood cell walls, the formation of the more soluble polymorph vaterite seems to be favored over calcite. This could be due to the high local supersaturation of ions in the confined cellular pores (Falini et al. 1996), which coincides with a lower pH drop. In various natural organic/ inorganic hybrid materials, oversaturation in confined space governs the formation of the less stable mineral polymorph (Falini et al. 1996; Falini et al. 2000; Falini et al. 2002; Rodriguez-Navarro et al. 2007; Meldrum and Cölfen 2008; Cantaert et al. 2013; Kirboga and Oner 2013). For instance, sustained supersaturation is decisive for the formation of vaterite in solution and for inhibiting its transformation into calcite (Spanos and Koutsoukos 1998; Sheng Han et al. 2006).

Apart from pore confinement effects, negatively charged groups on the surface might also mediate crystallization (Mann et al. 1991). The high density of calcium-complexating surface sites, especially lignin and pectin in the CML (Grant et al. 1973; Jarvis 1984; Caffall and Mohnen 2009) likely promotes the formation of CaCO₂ (Figure 4b,f,j,n). In primary cell walls, pectin





Figure 4: Raman mapping of $CaCO_3$ precipitates in beech and spruce (h–o). Vertex component analysis showing spatial distribution of cell wall carbohydrates (a, e, h, l), $CaCO_3$ polymorphs vaterite (b, f, i, m, n) and calcite (c, g, j) and corresponding endmember spectra (d, h, k, h). The scale-bar corresponds to 10 μ m.

(homogalacturonan) forms complex networks ionically cross-linked by calcium (Cosgrove 2005). In solution, (esterified) pectin slightly favors the formation of more soluble $CaCO_3$ phases or liquid precursor phases by destabilizing prenucleation clusters and inhibiting mineral nucleation (Rao et al. 2014). Furthermore, a transformation of the thermodynamically metastable vaterite to the stable calcite polymorph might occur more easily in larger cavities in a range of several tens of μ m (beech vessels, spruce EW tracheids) due to better fluid convection and mixing. In aqueous media, vaterite transforms into calcite by dissolution and crystallization from solution (Kralj et al. 1994). This seems to be less likely inside the compartmentalized nano-environment of the wood cell walls.

As previously demonstrated, a high content of incombustible $CaCO_3$ mineral renders wood less flammable (Merk et al. 2015). To evaluate the flammability of these $CaCO_3$ /wood composites on the micro- and the benchscale, the pyrolysis combustion flow calorimetry (PCFC) and cone calorimetry were applied. In micro-calorimetry, samples are pyrolized in an inert N₂ gas stream according to a temperature ramp followed by high temperature oxidation of the gaseous combustion products in a gas mixture $80/20 (N_2/O_2)$ (Lyon and Walters 2004; Sonnier et al. 2011). In Figure 5a, the heat release rate (HRR) of mineralized (colored curves) and untreated spruce (black curve) is plotted against the temperature. In good accordance with calorific values reported in the literature (Shafizadeh 1982; White 1987), the greater lignin content in softwoods gives rise to higher total heat release (THR), heat release capacity (HRC), and char residue values than in hardwood references. In CaCO₂/wood composites, the THR (Figure 5b), HRC (Figure 5c) and temperature of highest mass loss (T_{max}) are significantly lowered compared to the natural wood. Additionally, the THC and HRC remain more or less similar for various treatment cycles. A slightly more pronounced THR and HRC decrease has been found for spruce, presumably due to its higher relative mineral content. In the PCFC tests, the char yield of mineralized wood is approximately doubled as compared to unmodified wood (Figure 5d). A small amount of CaCO₂ seems to be disproportionally effective in increasing the char yield



Figure 5: Pyrolysis combustion flow calorimetry. (a) Heat release curves for natural (black line) and mineralized spruce (colored lines) averaged over five measurements. Total heat release (b), heat release capacity (c) and char yield are plotted as a function of the mineralization cycle for spruce and beech composites.

and the fire retardance of wood composites. These results are highly consistent with our previously published data (Merk et al. 2015), and it can be concluded that similar or even identical thermal decomposition mechanism are valid, and hence, the fire retarding performances are also similar.

Cone calorimetry tests of mineralized and pristine wood samples have been performed to simulate the combustion behavior in a developing fire (Schartel et al. 2005; Schartel and Hull 2007). In comparison to PCFC, physical fire-retardant mechanisms are more amenable to be studied in cone calorimetry (Sonnier et al. 2011). In this flammability test, wood samples are subjected to a constant heat flux of 50 kW m² from a conical radiant panel under well-ventilated conditions. As soon as sufficient pyrolysis gasses have evolved from the wood specimen, piloted ignition starts. In our studies, the observed time to ignition (Table 1) is hardly affected by the mineralization treatment.

Bimodal heat release profiles of beech and spruce timber are displayed in Figure 6a,b. Peaks of heat release rate (PHRR) correspond to flaming combustion, which is a cause of rapid fire spread in the environment. The initial PHRR (visualized in Figure 6c, t=40 s) originates from the oxidation of volatile pyrolysis products (Lowden and Hull 2013). As combustion proceeds, lowering of HRR indicates the formation of an insulating layer (Schartel et al. 2005; Batiot et al. 2014). When the protective char structure breaks up, the HRR undergoes a second maximum followed by gradual glowing and smoldering combustion (Lowden and Hull 2013). The thermo-oxidative decomposition of the condensed phase mainly occurs through cracks on the char surface as a result of the built-up gas pressure in deeper layers (Batiot et al. 2014). Only in native wood, major deformation was observed during the afterglow phase due to residual stress (Figure 6c, t=450 s). In mineralized wood, both peak HRR are substantially decreased and shifted to longer exposure times (Table 1), which indicates a lower fire growth rate and enhanced char stabilization. In essence, the PHRR values are reduced to a similar extent as in other fire-retardant systems reported in the literature (Simkovic et al. 2005; Simkovic et al. 2007; Hagen et al. 2009; Grześkowiak 2012; Mahr et al. 2012). As the mass loss values of unmodified and mineralized woods are comparable (Table 1), it can be assumed that the improved fire retardance of mineralized wood results from a combination of both physical and chemical effects. Incombustible minerals embedded into wood tissues most likely dilute the amount of combustible material and constitute a barrier for heat and mass transport during the pyrolysis. As depicted in Figure 7a, native wood yields brownish-black, highly disintegrated residues, whereas compact residues with partially intact cellular ultrastructure are seen in mineralized samples. Herein, the residue yield clearly exceeds the amount of incorporated mineral. Together with the lowered T_{max} recorded in PCFC, this indicates a base-catalyzed pyrolysis of cellulose (Shafizadeh 1982; Di Blasi et al. 2009; Merk et al. 2015).

Differences in the chemical composition of the heattreated woods studied were observed by FTIR spectroscopy (Figure 7b). In reference wood, the absence of a O-H

Parameters	Untreated beech	Mineralized beech	Untreated spruce	Mineralized spruce
t _{Ignition} (s)	28±4	30±5	18±3	18±1
Initial mass (g)	61±2	76±1	42±2	49±2
Total mass loss (%)	60±1	59±2	41±2	39±2
THR (MJ m ⁻²)	129±23	84±20	81±17	63±18
t _{1 PHRR} (s)	60±20	49±7	36±4	30±6
1. PHRR (kW m ⁻²)	190±13	96±10	173±16	105±18
t _{2 PHPP} (s)	307±15	464±10	354±11	420±13
2. PHRR (kW m ⁻²)	402±61	152±18	151±5	113±11

Table 1: Cone calorimetry data obtained at 50 kW m⁻² averaged over five samples per variation.

THR, Total heat release; PHRR, peak heat release rate.



Figure 6: Cone calorimetry data. Heat release profiles of unmodified (orange) and mineralized beech (blue) (a), pristine (black) and modified spruce (turquoise) (b). (c) Photos of cone calorimeter measurements of beech. Incandescent flaming is observed in reference beech (orange frame). The PHRR in untreated beech is shifted from 300 s to 450 s in mineralized beech (blue frame).

stretching vibration band (~3300 cm⁻¹) (Nishimiya et al. 1998) indicates a complete dehydration to carbonaceous char, which does not occur in mineralized wood. Infrared spectra of mineralized wood residues display modes originating from untransformed calcite (1796, 872, 847 and 712 cm⁻¹) (Xyla and Koutsoukos 1989) and remnant aromatic constituents (1400 and 1630 cm⁻¹) (Nishimiya et al. 1998).

Apart from the heat release rate, other reaction-to-fire characteristics, e.g. the generation of toxic gasses or soot, are relevant for evaluating the fire hazard. Upon endothermic decomposition (Satterfield and Feakes 1959), CaCO₃ releases carbon dioxide which potentially dilutes and cools flammable combustion products. In correlation with

a lowered HRR, smoke production is suppressed in $CaCO_3/$ wood composites (Figure 7c,d).

The various sample preparation procedures and test conditions have to be taken into account (Sonnier et al. 2011), when comparing HRC (Figure 5c) and PHRR results (Table 1) obtained by PCFC or cone calorimetry. For instance, PCFC requires only milligram quantities, whereas cone calorimetry samples have a uniform volume of $\sim 10 \times 10 \times 1$ cm³. Despite a lower relative mass uptake (17±2% in spruce, 18±2% in beech) in cone calorimetry samples, a substantially improved fire retardance was achieved compared to reference wood. HRC values are reduced to ca. 65% (beech) and 49% (spruce) after four



Figure 7: (a) Photographs of char residues of unmodified (orange) and mineralized beech (blue) (a), pristine (black) and modified spruce (turquoise) after combustion. (b) FTIR spectra of char residues. Dynamic smoke production rate of beech (c) and spruce (d) as a function of time.

mineralization cycles, while PHRR values are decreased to 38% and 61%, respectively. The higher bulk density of the beech samples (ρ =688±17 kg m³) compared to the spruce samples (ρ =458±19 kg m³) explains the higher THR and PHRR values of beech in cone calorimetry tests. When higher amounts of fuel are available, embedded fire-proof minerals might be more effective in slowing down the decomposition by limiting the oxygen and volatile supply. Barrier effects do not become apparent in non-flaming PCFC testing, as pyrolysis processes in the solid state are separated from the vapor phase (Lyon and Walters 2004).

Conclusions and outlook

In focus was the *in-situ* formation of CaCO₃ mineral by subsequent impregnation of wood with aqueous NaHCO₃ and alcoholic CaCl₂ electrolytes. The SEM, XRD, and Raman microscopic studies showed the deposition of high amounts of polycrystalline calcite and vaterite inside the wood lumina and to a lesser extent in adjacent cell walls. By performing micro-calorimetry and cone-calorimetry tests, a significantly improved fire resistance of CaCO₃/wood composites was demonstrated. Hence, the embedding of incombustible CaCO₃ into the wood can be

considered as an eco-friendly fire-retardant treatment. The probability is high that the presented approach can be upscaled to an industrial level.

The solute-exchange approach offers a high degree of versatility towards the mineral composition, mineral phase, and amounts of mineral in the wood structure as well as the choice of solvents. Various mineral compositions comprising different cations (earth alkali, light and heavy metals) and anions (carbonates, phosphates, sulfates, etc.) or mixtures thereof could be inserted into the wood structure, in analogy to the impregnation protocol as depicted for calcium carbonate formation in this study. From a fundamental research perspective, the mineralization of woody tissues sheds light on basic processes such as ion transport, crystal nucleation and growth. Further control over the mineral distribution, phase and morphology and their impact on the mechanical performance of wood-based hybrid materials will be subject of future studies.

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