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Topochemical analyses of furfuryl alcoholmodified radiata pine (*Pinus radiata*) by UMSP, light microscopy and SEM

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Abstract: Furfurylation is one of the wood modification techniques via catalytic polymerization of the monomeric furfuryl alcohol (FA) in the impregnated cell wall. Little is known about the topochemistry of this process. Brown rot degradation begins with lignin modification and therefore, the reactions between FA and lignin was one focus of this research. Furfurylated radiata pine (*Pinus radiata*) with three different weight percent gains (WPGs of 57%, 60% and 70%) after FA uptake was observed by cellular ultraviolet microspectrophotometry (UMSP) to analyze chemical alterations of the individual cell wall layers. Moreover, light microscopy (LM) and scanning electron microscopy (SEM) analyses were performed. The ultraviolet (UV) absorbance of the modified samples increased significantly compared to the untreated controls, indicating a strong polymerization of the aromatic compounds. Highest UV absorbances were found in areas with the highest lignin concentration. The UMSP images of individual cell wall layers support the hypothesis concerning condensation reactions between lignin and FA.

Keywords: brown rot, furfurylation, light microscopy (LM), lignin modification, scanning electron microscopy (SEM), ultraviolet microspectrophotometry (UMSP), wood modification

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Introduction

Brown rot basidiomycetes cause the most destructive type of wood decay, even before any mass loss (ML) of the wood is detectable (Schultze-Dewitz 1966; Bariska et al. 1983; Winandy and Morrell 1993). Cellulose and hemicelluloses are mainly metabolized, leaving a modified lignin behind (Danninger et al. 1980; Eriksson 1990; Goodell 2003). Lignin maintains its macromolecular nature throughout the brown rot decay (Kirk 1975; Niemenmaa et al. 2008; Yelle et al. 2008), but its structure is modified substantially by hydroxyl radicals produced by the fungi (Yelle et al. 2008, 2011; Arantes et al. 2009, 2011; Martínez et al. 2011). The lignin modification begins in the outermost part of the secondary cell wall (S2) and the compound middle lamella (CML) (Fackler et al. 2010; Rehbein and Koch 2011).

Wood preservation based on technologies with low environmental impact are mandatory today. Different approaches are known in this regard, for instance, modifications with melamine formaldehyde (Stamm 1964), with acetic anhydride (Rowell et al. 1994), or thermal modification (Militz 2002), the effects of which are based on chemical, biological and physical interactions, resulting in increased durability and dimensional stability (Hill 2006; Ringman et al. 2014). Another promising approach is the furfurylation process. Furfuryl alcohol (FA) is obtained from renewable resources of agricultural crop waste (corn or sugar cane production) consisting of pentosans, which are easily hydrolyzed, and the pentoses obtained are dehydrated to furfural (F) (Hill 2006; Uppal et al. 2008; Moghaddam et al. 2016), which is then converted to FA. F and FA are by far the most important furan derivates (Gandini and Lacerda 2015), which can be polymerized by several organic and mineral acids, such as by Lewis acids (Lande et al. 2004a). Ünver and Öktem (2013) investigated controlled cationic polymerization of FA, visualized by the UV bands at $\lambda_{228 \text{ nm}}$ and $\lambda_{286 \text{ nm}}$.

Furfurylation begins with FA impregnation of wood followed by *in situ* polymerization, which leads to a permanent swelling of the cell wall (Goldstein 1955, 1960; Stamm 1977; Schneider 1995; Bryne et al. 2010). In the last decade, FA impregnation was, for example, applied to

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improve plasticization and fixation during the densification of wood (Pfriem et al. 2012), and new catalysts were developed (Li et al. 2015; Sejati et al. 2017). Not all the details of the furfurylation process are well understood. The FA solution enters the cell wall to some extent but also fills the cell lumens completely or partially (Schneider et al. 2000). Depending on wood species and FA concentration, the curing temperature ranges from 45°C to 140°C (Westin 2003). Furfurylated wood (W_{EA}) has better durability against biodegradation (Lande et al. 2004c; Esteves et al. 2011; Gascon-Garrido et al. 2013; Li et al. 2015; Sejati et al. 2017) and has a low environmental impact (van Eetvelde 1998; Lande et al. 2004a,b,c; Pilgård et al. 2010a, 2010b). Furfurylated softwoods and hardwoods are available on the market with the equivalent quality to woods protected by copper, chromium and arsenic (CCA) solutions. W_{EA} is darker than native wood and has an aesthetic appeal similar to that of natural durable (and dark) tropical timbers like teak, ipé or azobé. W_{FA} fulfills the requirements for hazard classes 3 and 4 applications according to EN 335. Furfurylated southern yellow pine and radiata pine are admitted as "proven" for window frames on the German market (VFF-Merkblatt HO.06-4 2016). Previous studies by infrared (Venås and Rinnan 2008) and fluorescence microscopy (Thygesen et al. 2010) on $W_{_{\mathrm{FA}}}$ have shown that FA polymerization takes place within the wood tissue. Higher fluorescence occurs in lignin-rich parts, mainly in the cell wall corner regions (CC) and the CML (Thygesen et al. 2010). Nuclear magnetic resonance (NMR) spectra of early stages of FA polymerization demonstrated that lignin model compounds form covalent bonds in a liquid-phase system with the FA polymer (Nordstierna et al. 2008). Chemical bonds between the FA polymer and wood tissue, however, have not yet been proven.

To learn more about furfurvlation, ultraviolet microspectrophotometry (UMSP) was applied in the present study. UMSP is an established analytical technique which enables the direct imaging of lignin distribution and lignin modification in the course of biodegradation and technological delignification within individual cell wall layers in situ (Fergus et al. 1969; Scott et al. 1969; Saka et al. 1982; Fukazawa 1992; Koch and Kleist 2001; Koch et al. 2003a; Koch et al. 2003b; Koch and Grünwald 2004; Rehbein and Koch 2011; Koch and Schmitt 2013; Aguayo et al. 2014; Kojima et al. 2014; Bianchi et al. 2016; Ehmcke et al. 2016). The π electron-rich lignin shows UV maxima at around $\lambda_{212\,\mathrm{nm}}$ and $\lambda_{280\;nm}$ (Scott et al. 1969; Goldschmid 1971; Fukazawa 1992). Softwood lignin is mainly composed of guaiacylpropane moieties with an absorbance maximum (A_{max}) at around $\lambda_{280 \text{ nm}}$ (Musha and Goring 1975; Takabe 1992). The UV-A_{max} data are lower in lignins with higher syringpropan

unit participation, seen on the increasing OCH₂/C₀ ratios (Musha and Goring 1975; Fujii et al. 1987). Modified timbers, such as heat-treated and N-methylol melamine-modified hardwoods (Kielmann et al. 2013; Mahnert et al. 2013; Sint et al. 2013), hydrothermal-modified wood (Andersons et al. 2016) and degraded furfurylated softwood (Ehmcke et al. 2016), were also studied by UMSP.

The aim of the present study was to investigate FAmodified radiata pine (Pinus radiata) by the UMSP method to search for indications of lignin modification via chemical bonds with FA.

Materials and methods

Radiata pine (P. radiata) sapwood boards were industrially furfurylated under different process conditions and distributed to project partners of an ongoing research project (Research council of Norway project 219294/O30). The furfurylation processes are based on a full-cell impregnation with different FA solutions, buffer agents and catalysts followed by steam curing and kiln drying. Water-based [process A, a weight percent gain (WPG) of 60%] and alcohol solutions (process B, 70% WPG) were applied, both containing FA and standard catalysts. Process C was performed with an experimental mix of a water-based solution and an alternative catalyst leading to a WPG of 57%. The wood from process A is a commercial product and available on the market. Each modified board in this project was matched by an untreated board (W,,,,t).

Sample preparation for UMSP: Small blocks of $W_{_{FA}}$ 1 mm \times 1 mm \times 5 mm (R \times T \times L) were prepared for each treatment type (W_{FA,A}, W_{FA,B} and $W_{\mbox{\tiny FA,C}})$ and for $W_{\mbox{\tiny unit}}.$ Selected wood blocks were dehydrated in a graded series of ethanol and then embedded in Spurr's (1969) epoxy resin (Electron Microscopy Sciences, Hatfield, PA, USA) under mild vacuum conditions. The polymerization process was catalyzed by thermal curing at 70°C for 12 h. The embedded specimens were trimmed with a razor blade (Plano GmbH, Wetzlar, Germany) to provide an area of approximately 0.5 mm² and sectioned with a rotary microtome (Leica, RM2265, Leica Mikrosysteme Vertrieb GmbH Wetzlar, Germany) equipped with a diamond knife (DiATOME histo, 4.0 mm, 45°, Diatome AG, Biel, Switzerland). The semi-thin sections (1 µm) were transferred to quartz microscopic slides (Plano GmbH, Wetzlar, Germany) and thermally fixed. The sections were immersed in a drop of non-UV-absorbing glycerine (glycerine/water mixture $n_{\rm p}$ = 1.46) and covered with a quartz cover slip. As reference, unmodified radiata pine sapwood specimens were also prepared, following the same embedding process.

In addition, small cubes (1 cm³) of $W_{\mbox{\tiny untr}}$ and $W_{\mbox{\tiny FA}}$ were prepared for light microscopy (LM) and scanning electron microscopy (SEM) analyses. For conventional LM, the cubes were sectioned with a sliding microtome (Leica SM 2000 R, Leica Mikrosysteme Vertrieb GmbH Wetzlar, Germany) and sections of the three anatomical directions (transverse, radial and tangential) were mounted on microscopic slides and embedded in Euparal (Carl Roth, Carl Roth GmbH+Co. KG, Karlsruhe, Germany). For SEM analysis, specimens were boiled in water and then manually cut with a thin razorblade to obtain a plain surface perpendicular to the axial direction. The specimens were stored overnight under vacuum and then sputter-coated by a coating unit (International Scientific Instruments Co, New Delhi, India) with a 10-15-nm gold film.

Cellular UMSP: The ultrathin sections were placed in a Zeiss UMSP 80 (Carl Zeiss AG, Oberkochen, Germany) equipped with a scanning stage. The principle of the UMSP investigation is based on the typical UV absorbance of lignin in the range of $\lambda_{\scriptscriptstyle 270-280\;nm}$ (Musha and Goring 1975). For scanning, a defined wavelength of $\lambda_{_{280\;nm}}$ (the $A_{_{max}}$ of softwood lignin) was selected. The scan program Automatic Photometric Analysis of Microscopic Objects by Scanning (APAMOS, Carl Zeiss AG, Oberkochen, Germany) digitizes square fields of a local geometrical resolution of 0.25 μm × 0.25 μm and a photometrical resolution of 4096 grayscale levels. To visualize the absorbance intensities, the grayscale levels were converted into 14 basic colors as described in detail by Koch and Kleist (2001). The scans can be depicted as two-dimensional (2D) or three-dimensional (3D) image profiles, and a statistical data evaluation results in a semi-quantitative lignin distribution. For control purposes, the lignin distribution within individual cell types and cell wall layers of three $\boldsymbol{W}_{\text{untr}}$ specimens were scanned.

In addition, the specimens were also subjected to photometric point measurements with a spot size of 1 µm2. The UV spectra of individual cell wall layers were recorded in the range between $\lambda_{_{250\,nm}}$ and $\lambda_{470\,\mathrm{nm}}$ and statistically evaluated. A TIDAS MSP 800 microscope spectrometer (J&M Analytik AG, Esslingen, Germany) equipped with TIDAS-DAQ software was available. The spectral characteristics of 15-20 point measurements of individual cell wall layers and FApolymerized deposits in the lumen were evaluated.

LM and SEM: To localize FA-impregnated cell elements, selected samples were analyzed by conventional LM (Axiophot, Carl Zeiss AG, Oberkochen, Germany) equipped with a digital camera (AxioCam, Carl Zeiss AG, Oberkochen, Germany) combined with AxioVision software (Carl Zeiss AG, Oberkochen, Germany). The SEM was equipped with a tungsten cathode (EVO 40, 10-12 kV, Carl Zeiss AG, Oberkochen, Germany).

Results and discussion

Transverse sections of W_{untr} and W_{FA} from the FA modification processes were observed by cellular UV microscopy at $\lambda_{280 \text{ nm}}$. Preliminary tests showed that the standard thresholds for the UMSP analyses of untreated wood tissue (min 10% and max 80%) were not suitable to evaluate W_{FA} due to the high absorbance intensities (Ehmcke et al. 2016). To obtain a comparable evaluation and image generation, the thresholds were therefore individually adapted to a minimum of 1.0% and a maximum of 67.5% for all investigated specimens.

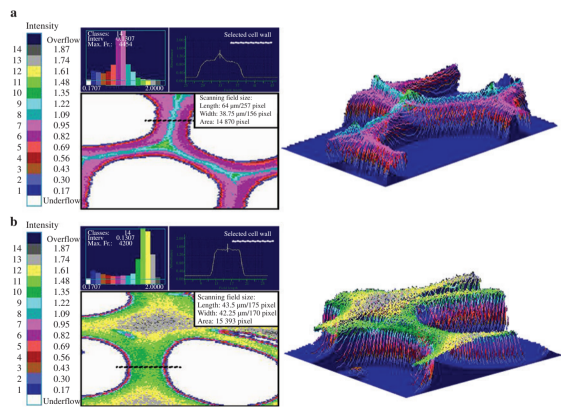


Figure 1: UV microscopic scanning profiles of untreated (a) and furfurylated (b) radiata pine (process A, 60% WPG). The data are statistically evaluated from their corresponding histograms. The colored pixels mark the absorbance intensity at $\lambda_{280\,\mathrm{nm}}$. A selective line scan of an individual cell wall region intuitively visualizes the local lignin concentration and chemical cell wall modification.

UV microscopic scanning profiles

In Figure 1, representative UV microscopic scanning profiles of W_{untr} and $W_{FA,A}$ (60% WPG, commercial product) are presented. The colored pixels mark the absorbance intensities at $\lambda_{280~nm}$. The data of each UV profile are statistically evaluated from its corresponding histogram. The high resolution (0.25 μ m² per pixel) enables a detailed differentiation of the UV absorbance within the individual cell wall layers (Koch et al. 2003a). In addition to the standard 2D UV micrograph, a selective line scan of an individual cell wall region intuitively visualizes the local lignin concentration and the chemical cell wall modification. For spatial illustration, the scanning area is also presented as a 3D image profile.

A representative profile of natural lignin distribution in cell walls of W_{untr} is presented in Figure 1a. The highest absorbances occur in the area of the CML and the CC as previously described for W_{untr} (Koch and Kleist 2001; Koch and Grünwald 2004) and also for thermally (Mahnert et al. 2013) and hydrothermally modified wood (Andersons et al. 2016). For the W_{untr} tissue, the following UV absorption data ($A_{280~nm}$) are typical: 1.22 for the CML, 1.48 for the CC and lower, slightly varying values for the S2 layer (0.69–0.95). The fluctuating values for the S2 layer were already described by Takabe (2002). The line scan of a double cell wall in the radial direction and the corresponding 3D image illustrates the high absorbance intensities of the CML as a pronounced peak and a highly absorbing band.

For comparison, the UV scanning profiles of W_{untr} and $W_{FA,A}$ (60% WPG) are presented in Figure 1. The profiles (Figure 1b) show significantly higher absorbances of the entire cell wall compared to W_{untr} . With regard to the typical lignin distribution, both UV micrographs of W_{untr} and W_{FA} reveal a similar spectral behavior, where the absorbance intensities are increasing from the S3 layer towards the area of the CML and the CC. For W_{FA} , a clear shift is visible

to green-, yellow-, and gray-colored pixels, representing higher UV intensities in the range of 1.22–1.74 ($A_{280~\mathrm{nm}}$), as it was already documented by Ehmcke et al. (2016). With reference to fluorescence studies (Thygesen et al. 2010), the highest amount of polymerized FA was detected in the lignin-rich areas. The line scan in Figure 1b visualizes a distinct increase of the absorbance values from the outer cell wall towards the CML. On the other hand, the CML evinces a less-pronounced peak compared to W_{untr} in Figure 1a. The CML in Figure 1b is associated with a broad plateau of high UV intensities representing the S2 layer ($A_{280~\mathrm{nm}}$ 1.22–1.35), which indicates an intensive modification of the cell wall.

 $W_{\rm FA,B}$ (70% WPG) and $W_{\rm FA,C}$ (57% WPG) are presented in Figures 2 and 3. The cell wall layers of $W_{\rm FA,B}$ (Figure 2) reveal relatively homogenous absorbance intensities in the green color range ($A_{\rm 280~nm}$ 1.22–1.35). The CC and CML are somewhat more pronounced and characterized by rare yellow pixels ($A_{\rm 280~nm}$ 1.48). Figure 3 shows lower intensities with bluish-green colors for $W_{\rm FA,C}$. The cell wall layers of $W_{\rm FA,C}$ show increasing intensities beginning with the cell lumen ($A_{\rm 280~nm}$ 0.17–0.56) to S2 ($A_{\rm 280~nm}$ 1.09–1.22), to the CML and the CC ($A_{\rm 280~nm}$ 1.35–1.61), where the absorbance intensities culminate. Compared to $W_{\rm untr}$ (Figure 1a) and all other treated materials (Figures 1b, 2 and 3), the wood tissue of $W_{\rm FA,A}$ (Figure 1b) shows the highest absorbances with a clear shift to green, yellow and gray pixels ($A_{\rm 280~nm}$ 1.22–1.74).

The mean UV absorbances did not increase proportionally to the loadings (i.e. to the WPGs) of the analyzed samples (Table 1). The highest loading was generated by process B with 70% WPG, representing the lowest A $_{\rm 280\ nm}$ with 1.07, compared to process A with A $_{\rm 280\ nm}$ 1.16 and C with A $_{\rm 280\ nm}$ 1.09, but these differences are not significant. The slight deviations could be explained by differences in the anatomical structure and the high resolution of the individual measuring fields. In general, the individual parameters of the FA modification (in terms of catalyst concentration,

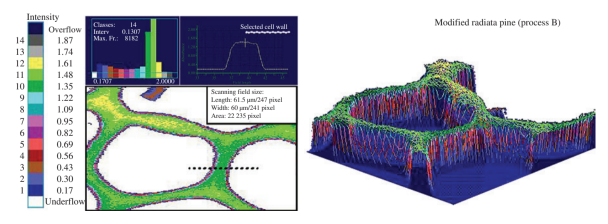


Figure 2: UV microscopic scanning profiles of furfurylated radiata pine (process B, 70% WPG).

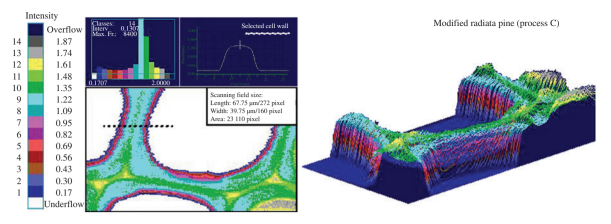


Figure 3: UV microscopic scanning profiles of furfurylated radiata pine (process C, 57% WPG).

Table 1: Mean UV absorbance (A_{280nm}) of untreated and modified radiata pine and maximum UV absorbance values (A_{max}) of polymerized FA deposits (processes A, B and C, threshold minimum 1.0% and maximum 67.5%).

		UMSP-sca	UV-Abs. spectra				
Process	WPG (%)	A _{280 nm}	StD	(pixel) ^b	λ(nm)	A _{max}	StD
Untr.	0	0.74	0.05	12 733	-	-	_
Α	60	1.16	0.07	19 752	278	1.24	0.07
В	70	1.07	0.06	14 426	256	1.21	0.07
C	57	1.09	0.09	23 103	278	1.20	0.06

^aLumens filled with FA, ^bField size. WPG is for weight percentage gain. StD is for standard deviation.

processing temperature and time and pH value of the solution) strongly affect the final product by the degree and type of chemical bonds (Lande et al. 2004a).

Compared to W,,,, higher UV absorbances were also observed by UMSP on thermally modified wood (160°C-180°C) (Mahnert et al. 2013). Andersons et al. (2016) reported similar results at lower temperatures, but at 140°C for 1 h only a slight impact on the UV absorbance was seen. The furfurylation occurs below 140°C (Westin 2003). The chemical reactivity of the lignin is induced above a temperature of 80°C (Koch et al. 2003a) and real thermal changes of lignin are expected to begin at 150°C (Fengel and Przyklenk 1970). Ünver and Öktem (2013) showed that polymerized FA is characterized by a UV- $\!A_{\rm max}$ at around $\lambda_{280 \text{ nm}}$, i.e. which is also the case for untreated softwood lignins. Accordingly, the presented results concerning the absorbance increment at 280 nm can be unambiguously interpreted as a manifestation of the preferred deposition of (condensed) UV-active compounds at places of high lignin concentration in the cell wall. Most likely chemical bonds between FA and lignin were also produced, but of course, this assumption is not yet proved definitively.

UV absorption spectra

UV spectra with a spot size of 1 µm² in the wavelength range from 240 to 600 nm were recorded to study the spectral behavior of $W_{\scriptscriptstyle FA}$ on a cellular level. The spectral characteristics of the polymerized FA deposits in the cell lumen of W_{FA} tissues are given in Figure 4. The UV spectra of the FA deposits from all three modification processes show approximately similar profiles. Common to all three modifications are (i) the high absorbance intensities, (ii) a distinct shoulder in the wavelength range between $\lambda_{\text{340 nm}}$ and $\lambda_{380\,\mathrm{nm}}$ and (iii) certain absorbance levels in the visible light range resulting from large chromophoric structures that were formed during the furfurylation processes. The absorption maxima of the FA deposits detected in WFA.A and $W_{FA,C}$ around $\lambda_{278 \text{ nm}}$ are similar to the findings of Ünver and Öktem (2013). Comparable peaks around $\lambda_{270 \text{ nm}}$ are also described by Gandini and Belgacem (1997), who studied the photopolymerization and photocrosslinking

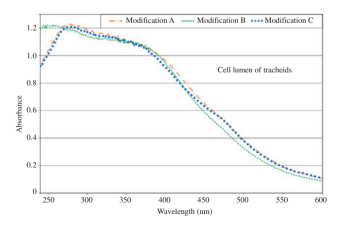


Figure 4: Representative UV absorbance spectra of polymerized FA deposited from all three modification processes in the cell lumens of radiata pine.

0.6

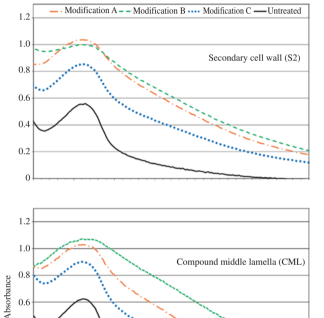
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of simple furans and F. The A_{max} of the deposits detected in $W_{_{FA\;B}}$ are less pronounced and shifted hypsochromically (Nic et al. 2014) to $\lambda_{256\;nm}$. This slight variation of the spectral behavior may have been caused by the individual process parameters of the furfurylation and are independent of sample loading (Table 1).

In Figure 5, representative lignin spectra of the individual cell wall layers of W_{instr} (black line) and the three furfurylated wood types are presented. The cell wall layers (S2, CML and CC) of W_{untr} show typical UV spectra of lignified softwood tracheids, with the characteristic A_{max} at



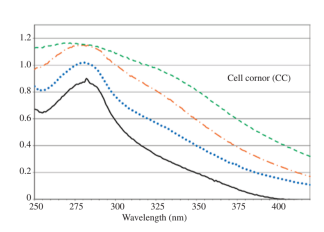


Figure 5: Representative UV absorbance spectra of the individual cell wall layers (S2, CML and CC) of untreated and furfurylated radiata pine (processes A, B and C).

 $\lambda_{280 \text{ nm}}$ attributed to the strongly absorbing guaiacyl-type units (Musha and Goring 1975; Fujii et al. 1987). The different absorbance intensities at $\lambda_{_{280\,nm}}$ are strongly correlated to the lignin concentration, with the highest amount in the CC (Koch and Kleist 2001; Koch and Grünwald 2004). In this case, the absorbance in the CC of W_{untr} is 1.6 times higher than in the S2 (Table 2). The spectra of the modified cell wall layers show similar characteristics, with significantly higher absorbance values for the individual cell wall layers (Figure 5) as was already demonstrated by the UV scanning analyses.

The spectra of $W_{FA,C}$ (57% WPG) appear close to the course of the W_{untr}, with generally higher absorbance intensities. The highest absorbances can be detected for the CC with 1.17 $(A_{272 \text{ nm}})$ and for the CML with 1.07 $(A_{_{280\ nm}})$ for $W_{_{FA,B}}\text{, i.e.}$ for specimens with the highest WPG (70%) (Table 2). The spectra of these strongly modified cell wall layers display a bathochromic shift and shoulders with lower intensities around 320 nm (compared to W,,,,...). This is probably due to the formation of conjugated double bonds. The higher degree of conjugation stabilizes π - π * transitions resulting in bathochromic shifts (Goldschmid 1971). In comparison, the spectra of the CML and CC show a course more similar to the spectra of polymerized FA in the lumina ($\lambda_{300 \text{ nm}}$ and $\lambda_{370 \text{ nm}}$, Figure 4), whereas the spectra of the S2 show only a shoulder with lower intensity. The strong increase from 0.56 (S2 in W_{untr} , $A_{282 \text{ nm}}$) to 1.04 (S2 in $W_{FA,A}$, $A_{278 \text{ nm}}$) and the shift of the bands of the $W_{_{F\!A}}$ cell walls indicate a higher condensation of carbonyl groups as a possible reaction of FA with the guaiacyl units of softwood lignin (Lande et al. 2004c).

Light microscopy and SEM

The LM and SEM images give insight into the cellular deposition of polymerized FA in the tissues. Figure 6 presents the LM observations in the three anatomical directions (transverse, radial and tangential) of W_{mater} and W_{EA} samples from the three process variants. The cross and radial sections include earlywood (EW) and latewood (LW). LW is easier to treat due to a larger amount of nonaspirated bordered pits compared to the EW (Phillips 1933; Liese and Bauch 1967; Zimmer et al. 2009). The sections in W_{FAA} (60% WPG, Figure 6d) and W_{FAA} (57% WPG, Figure 6j) show filled lumens in the LW. The transverse section of $W_{\text{FA.B}}$ (70% WPG, Figure 6g) unexpectedly displays less filled lumens in the LW tracheids. As the treating solutions have the same pathways as in the living tree (Nicholas and Siau 1973), it was assumed that

Table 2: Mean UV absorbances of the secondary cell wall (S2), compound middle lamella (CML) and cell corners (CC) of untreated and A-, B- and C-type modified radiata pine.

Process	WPG (%)	S2			CML			ССС		
		λ (nm)	Abs.	Δ	λ (nm)	Abs.	Δ	λ (nm)	Abs.	Δ
Untr.	0	282	0.56		281	0.62		282	0.90	
Α	60	278	1.04	0.48	280	1.03	0.41	278	1.15	0.25
В	70	280	1.00	0.44	280	1.07	0.45	272	1.17	0.27
C	57	280	0.86	0.30	280	0.90	0.28	281	1.02	0.12

The data are averages of 15–20 individual spectra. Δ is the absorbance difference between the untreated control and the FA-modified samples.

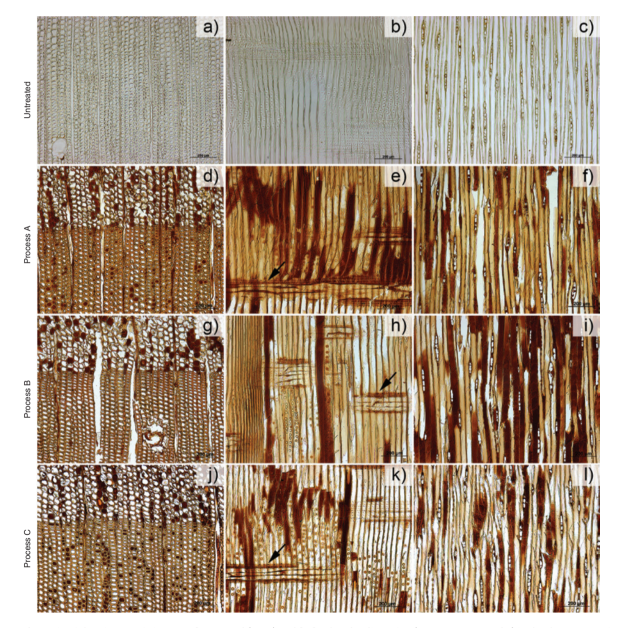


Figure 6: Light microscopic images of untreated (6a-c) and furfurylated radiata pine (processes A, B and C) in the three anatomical directions (transverse 6a, d, g, j; radial 6b, e, h, k; tangential 6c, f, g, l).

Compared to the untreated material, the furfurylated tissue appears darker with polymerized FA deposits in cell lumens. Filled ray tracheids are marked by arrows.

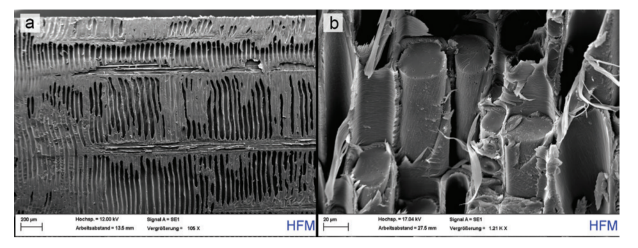


Figure 7: SEM images of furfurylated radiata pine. The overview image shows the cellular distribution of polymerized FA deposits in the radial section (a). The polymerized FA is spatially displayed as a compact, lumen filling substance (b).

polymerized FA would be located in the easily accessible parenchyma cells of the wood rays. Radiata pine investigated by Wardrop and Davies (1961) showed an apparently selective penetration of ray parenchyma and less intensively penetrated ray tracheids. However, the radial and tangential sections of the $W_{\rm FA}$ samples revealed that the FA was accessorily deposited in the ray tracheids (Figure 6e, h, k). One reason could be that the FA solution enters the ray tracheids predominantly via the connecting pits of the EW tracheids. The variability in filled cell lumens in the axial and radial directions supports the findings of Lande et al. (2010) or Zimmer et al. (2014) and are manifestations of varying permeability in Scots pine sapwood. This seems to also apply for radiata pine sapwood.

Aspirated tracheid bordered pits are assumed to function as physical barriers in case of impregnation (Stamm 1946; Zimmer et al. 2014). UMSP indicates a concentration decrement of polymerized FA from CML to S2. This can be explained by the early observation of Wardrop and Davies (1961) that aspirated bordered pits lead directly to the CML and serve as entrance openings during impregnation of the cell wall.

An uneven distribution and differently filled cell types can be seen. The permeability of W_{untr} may vary due to various physiological parameters in the living tree and also by the drying parameters (Booker 1990; Booker and Evans 1994; Zimmer et al. 2009, 2014; Lande et al. 2010). For successful wood protection system performance, a homogenous distribution of the treating solution in the wooden tissue is essential (Zimmer et al. 2012). Degradation studies reveal the importance of a consistent penetration and polymerization of FA during the production of

 $W_{\scriptscriptstyle FA}$ (Venås 2008). Certainly, more systematic studies are needed in this context.

The SEM images (Figure 7) confirm the UMSP observations and provide more detailed information on the cellular distribution of polymerized FA deposits. The cell lumens of the ray parenchyma cells appear empty (Figure 7a), as already observed under LM. In Figure 7b, the polymerized FA is spatially displayed as a compact lumen-filling substance. These deposits are obviously attached to the tracheid walls.

Conclusions

UMSP is an appropriate method for topochemical studies of $W_{\rm FA}$ subjected to various processes of modification. The measurements clearly show that the entire wood cell wall is modified, especially in the regions with the highest lignin concentration. The analyzed samples are characterized by strongly increased UV absorbances as a result of the formation of (condensed) aromatic compounds. This leads to the assumption that chemical bonds between FA and lignin may have been present. More specific knowledge of the nature of these bonds would probably be helpful in improving the biological resistance of furfurylated wood.

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