Natural and artificial ageing of spruce wood as observed by FTIR-ATR and UVRR spectroscopy

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Abstract

Spruce samples, naturally aged for 200, 400 and 500 years, artificially aged by a hydrothermal treatment (at 180, 160 or 130°C, relative air humidities of 14%, 40%, or 60% and for treatment times between 1 to 50 h), as well as reference samples, were analysed by Fourier transform infrared spectroscopy (FTIR) attenuated total reflection (FTIR-ATR) and ultraviolet resonance Raman (UVRR) spectroscopy. Natural ageing mostly affected the hemicelluloses and lignin, as observed from the FTIR-ATR and UVRR spectra, respectively. The UVRR spectra of the same samples after acetone extraction indicated that lignin was partially degraded and quinone structures were possibly formed. Artificial ageing at 160°C showed a significant change in the lignin structure, a well-known effect in the thermal treatment of wood, whereas treatment at 130°C did not alter the wood structure to any significant extent. Principal component analysis of the UVRR spectra confirmed that the spectra of artificially aged wood up to 160°C are dissimilar to naturally aged wood and which are also dissimilar to unaged wood.

Keywords: FTIR; heat-treated wood; PCA; UV resonance Raman spectroscopy (UVRRS); wood ageing.

Introduction

The slow deterioration of objects of cultural heritage is often called ageing; however, with appropriate storage and protection, ageing can be slowed down. Understanding how the mechanical and chemical properties of a material degrade over time can contribute to the knowledge of the most appropriate protection conditions. Ageing indoors must be differentiated from weathering outdoors, but even the former is a complex process, including physical and chemical changes on the surface, which is exposed to photo-modification and intensive oxidation, and in the bulk wood, i.e., below the surface. The interrelating factors triggering and influencing ageing are: the long-term exposure to fluctuating relative humidity; changes in ambient temperature; oxygen penetration into the bulk material; mechanical loading; and, in the worst case, fungi and bacteria. Deceleration and prevention of wood ageing is a topic of great interest to historians and conservationists.

Ageing and thermal treatment have similar effects (colour change, decrease of physical properties) as highlighted in a recent paper by Matsuo et al. (2011). Matsuo et al. compared the colour change of wood in historical buildings and heattreated wood and found that the colour change (ΔE^*_{ab}) after 921 years at ambient temperature was almost equivalent to that of heating, which can be considered as artificial ageing, approximately for 6.7 h at 180°C. Thus, the vast recent literature about thermal treating of wood gives a good opportunity to also gain insight into the slow processes of ageing (González-Peña and Hale 2009a,b; Bryne and Wålinder 2010; Bryne et al. 2010; Tuong and Li 2011). The relationship between chemical changes and mechanical properties of thermally treated wood is summarised by Windeisen et al. (2009). Also in the present study, artificial ageing by means of hydrothermal treatment was the preferred method.

Fourier transform infrared spectroscopy (FTIR) spectroscopy is an analytical technique well suited for studying the changes that take place in wood during ageing or weathering (Borgin et al. 1975; Fengel 1991; Horn et al. 1994; Müller et al. 2003; Pandey and Pitman 2003; Popescu et al. 2005; Ucar et al. 2005; Dobrica et al. 2008). It is recognised that a decrease in polyose content occurs during natural ageing as a result of a slow hydrolysis initiated by acetic acid which originates from cleaved O-acetyl groups of hemicelluloses. Similar observations were seen on thermal treated wood with infrared (IR) spectroscopy techniques (Pandey and Pitman 2003; Windeisen et al. 2003; Nuopponen et al. 2004; Tjeerdsma and Militz 2005; Windeisen and Wegener 2008). Thermal treatment also leads to the removal of extractives. During ageing, extractives move by evaporation, transportation by water movement, and diffusion towards the surface, and they are partly oxidised (Lavoie and Stevanovic 2006). The crystalline structure of cellulose in 300-400-year-old wood does not differ from fresh, unaged, wood. However, the degree of condensation of lignin and the number of polyconjugated systems increases as a result of oxidation and the amount of acid soluble lignin decreases (Borgin et al. 1975). In thermal treatment, a decrease in the amorphous polysaccharide content is observed, as well as condensation and

demethoxylation of lignin (Tjeerdsma et al. 1998; Nuopponen et al. 2004).

The usefulness of FTIR combined to ultraviolet resonance Raman spectroscopy (UVRRS) was demonstrated by Nuopponen et al. (2004, 2005, 2006). UVRRS is based on the sensitive interaction of lignin with UV light (Halttunen et al. 2001; Saariaho et al. 2003; Mononen et al. 2005; Nuopponen et al. 2006; Pandey and Vuorinen 2008), whereas the UV absorption by the hemicelluloses and cellulose is negligible.

Today, methods of chemometrics are an integral part of the utilization of modern hyphenated techniques. They make visible, even small changes, not otherwise visible at the first glance (Nuopponen et al. 2004, 2005; Martin et al. 2005). Chemometrics are frequently combined with FTIR and UVRRS; one speaks of multivariate data analysis with the approach partial least quares (PLS) or principal component analysis (PCA). PCA is suited for grouping of spectra, as well as for defining bands responsible for their similarities and dissimilarities (Ferraz et al. 2000; Saariaho et al. 2005; Esbensen 2009).

The aim of this study was to compare, both visually and with PCA, FTIR and UVRR spectra from un-aged wood, naturally aged wood and wood artificially aged by means of hydrothermal treatment. Qualitative and semi-quantitative analyses were performed. This approach should help draw conclusions as to the suitability of the different artificial ageing processes to reproduce the chemical changes occurring during natural ageing.

Materials and methods

Wood samples

Heartwood samples of spruce (*Picea abies*) wood were the focus of the study, namely: naturally aged old wood (OldW), wood artificially aged by hydrothermal treatment (HtW) and recent (reference) wood (RefW). The 400-year-old sample (400yW) was taken from a floorboard showing some radial cracks and visible insect attack, but no visible fungal attack. The 200- and 500-year-old samples (200yW and 500yW) were taken from beams used as structural elements in roofs. The age of the OldW samples was determined by dendrochronology. The parameters of artificial ageing of HtW samples are summarised in Table 1. The samples were stabilised to their equilibrium moisture contents (EMC) at 20°C and 50% relative humidity (RH).

All specimens were taken not less than 7 mm away from the wood surface to avoid any influence of photo degradation or surface weathering. The specimens for FTIR spectroscopy were cut with a

Table 1 Description of the artificial ageing processes by hydro-treatment (HtW) performed at 1 bar.

Series no.	Temp. (°C)	RH (%)	Time (h)	Abbreviation HtW _{Temp/RH/Time}
1	180	60	30	HtW _{180/60/30}
	160	14	50	HtW160/14/50
	160	60	30	HtW160/60/30
2	130	40	1	HtW _{130/40/1}
	130	40	2	HtW _{130/40/2}
	130	40	10	HtW _{130/40/10}

blade in the radial direction to obtain thin samples with dimensions of $0.05-0.1\times5\times20$ mm³ (t×1×r). Five pieces were prepared for each sample type. For the UVRRS measurements, eight wood specimens with dimensions of $0.7\times5\times25$ mm³ (t×1×r) were prepared from each sample type. They were cut with a chisel following the radial direction and the freshly cut surfaces were reserved for the measurement.

Extractions

Samples were extracted for 6 h with acetone in a Soxhlet apparatus. After evaporation of the acetone, the extracted specimens were dried at room temperature before further analyses.

FTIR measurements

Instrument: Spectrum 100 (Perkin Elmer, Waltham, MA, USA) spectrometer equipped with an attenuated total reflection (ATR) unit. Scanning range: 4000 and 650 cm⁻¹; 32 scans; resolution 4 cm⁻¹. Baseline correction on four points (3960, 1784, 1545 and 782 cm⁻¹); normalization to the highest absorption band between 1550 and 1500 cm⁻¹. The spectral range 1800–600 cm⁻¹ was evaluated. Each specimen was measured at five different positions on the freshly cut radial face.

UVRR spectroscopy

The specimens were attached to a microscope slide with doublesided tape with the freshly cut radial face on top and gently pressed with another slide to ensure good adhesion. Instrument: Renishaw 1000 UV Raman Spectrometer (Renishaw, Goucestershire, UK) equipped with a Leica DMLM microscope (Leica Microsystems, Wetzlar, Germany). Light source: Innova 90°C FreDTM frequencydouble Ar+ion laser (Coherent Inc., Santa Clara, CA, USA) tuned to an excitation wavelength of 244 nm. Focus on the specimen was initially achieved with white light and a 40×objective lens. Then, the focus was adjusted to 30 μm deeper, so as to get the UV laser beam to focus on the specimen surface. The laser power was kept at 1 mW and the specimens were moved slowly along a line perpendicular to the annual rings with a motorised sample stage. The acquisition time was 30 s (line distance 6 mm), consisting of 2 CCD readouts of 15 s. The spectral data were analyzed by GRAMS/32 software (Thermo Galactic, Woburn, MA, USA). The spectra were normalised on the most intense band at 1602 cm⁻¹ for easier comparison.

PCA analysis of UVRR spectra

PCA of normalised UVRR spectra was carried out by the Unscrambler X (CAMO Software AS., Oslo, Norway) software. Algorithm: NIPALS with systematic cross-validation and ascertainment test. A correlation optimised warping (COW) was performed to reduce the differences between the explained variance calibration and validation curves. Loadings and score were then plotted.

Results and discussion

Qualitative analysis of FTIR ATR spectra

Five FTIR-ATR spectra were collected from the recent reference sample (RefW) to check the spectral variation within the sample (Figure 1). Intensity differences were observed but, of course, no shifts of the principal bands were seen



Figure 1 Five FTIR ATR spectra of the reference sample RefW in the fingerprint region. Numbers correspond to band assignment as described in Table 2.

(Figure 1). The differences in the band intensities are partly as a result of the heterogeneity of the measured material and differences in the surface quality, which also influence the quantification. To reduce errors from artefacts, the most representative spectra were selected from among the five spectra collected from each sample. The assignment of the 22 characteristic bands between 1800 cm⁻¹ and 600 cm⁻¹ are listed in Table 2 based on the literature (Faix 1992; Pandey and Pitman 2003; Müller et al. 2008). A low frequency band at 664 cm⁻¹ could not be assigned but it was considered in further analysis.

Figure 2 shows the characteristic FTIR ATR spectra of five different wood samples. The main changes in the spectra as a result of natural ageing (differences between the spectra of RefW and 400yW) were observed in the range 1800-1500 cm⁻¹ concerning the bands 1 (1735 cm⁻¹), 3 (1641 cm⁻¹) and 4 (1617 cm⁻¹). Natural ageing resulted in a decreasing intensity of bands 1 and 3, whereas band 4 increased. These spectral changes were interpreted as modification of hemicelluloses. The 400yW sample is the only one in which the intensity of band 5 (1508 cm⁻¹) was lower than that of band 4 (1617 cm⁻¹). Bands 4 and 5 are attributable to the aromatic skeletal vibration of lignin. The artificially aged (HtW) samples did not show remarkable changes in this regard. A similar effect in the intensity of bands 4 (1617 cm⁻¹) and 5 (1508 cm⁻¹) was also observed after UV treatment of wood (Müller et al. 2003; Pandey and Vuorinen 2008) and was interpreted as lignin decay combined with the formation of new carbonyl groups evidencing photo-induced oxidation of the wood surface.

Artificial ageing by hydrotreatments $HtW_{180/60/30}$, $HtW_{160/14/50}$ and $HtW_{160/60/30}$ (the indices are for temperature/RH/ time, respectively) also modified the band intensities in the 1800–1500 cm⁻¹ range. A significant decrease in band 1 (1735 cm⁻¹) and simultaneous increase in band 2 (1700 cm⁻¹) was visible. Accordingly, esterified carbonyl structures in xylan decreased and free acid groups increased. The small shift of band 5 (1508 cm⁻¹) to a lower wavenumber has been

Table 2FTIR band assignment in the $1800-600 \text{ cm}^{-1}$ region for different wood species (Faix1992; Pandey and Pitman 2003; Müller et al. 2008).

Band	Observed (cm ⁻¹)	Lit. (cm ⁻¹)	Compound or chemical group	
1	1735	1732	C=O Stretch in xylan (hemicelluloses)	
2	1700		Shift of 1735 after Py, wood welding, and TM*	
3	1641	1649	Absorbed O-H and conj. C-O; in Py: loss of water	
4	1617	1593	Arom. skeletal vibr. in L	
5	1508	1505	Arom. skeletal vibr. in L	
6	1463	1460	C-H def. in L and CH	
7	1452		C-H def. in L and CHe	
8	1423	1424	Arom. skeletal vibr. in L with C-H def.	
9	1368	1367	C-H def. in cellulose and hemicell.	
10	1335	1328	S ring and G ring condensed	
11	1317	1318	C-H vibration in cellulose	
12	1265	1267	C-O stretch lignin ring in G units	
13	1231	1235	C-O stretch in G units	
14	1204	1245-1220	Higher condensate structure in TM wood	
15	1157	1157	C-O-C vibration in cellulose and hemicell.	
16	1104	1122	Arom. skeletal vibr. and C-O stretch	
17	1051		C-O vibr. in cellulose and hemicell.	
18	1027	1031	C-O vibr. in cellulose and hemicell.	
19	897	897	C-H def. in cellulose	
20	877	859	C-H out-of-plane arom. in L	
21	808	818	C-H out-of-plane arom. in L	
22	664		Unknown	

G, guaiacyl; S, syringyl; *TM, thermal modification, Windeisen and Wegener (2008); L, lignin. CH, carbohydrates. Py, pyrolysis.



Figure 2 Qualitative analysis of the FTIR-ATR spectra of un-aged samples (RefW), 400-year-old sample (400yW) and hydrothermally treated samples of series 1 (HtW_{180/60/30}), HtW_{160/14/50}, HtW_{160/60/30}). The arrows show the relative changes of intensity or the position of the several absorption bands.

attributed to the splitting of the aliphatic chains of lignin (Faix 1992; Uçar et al. 2005). An increase in the intensity of band 11 (1317 cm⁻¹) as well as the shift of band 14 (1204 cm⁻¹) has been attributed to the formation of condensed structures (Faix 1992). Artificial ageing by hydro-

treatment leads to the same types of chemical changes that were observed following the thermal treatment of wood. The mildest treatment in the case of $HtW_{160/14/50}$ (i.e., treatment at 160°C, 14% RH for 50 h) resulted in spectral changes most similar to that of natural ageing.

Semi-quantitative analysis of FTIR ATR spectra

Semi-quantitative analysis was performed on the five spectra described above concerning the ratios of the 22 typical band intensities listed in Table 3. The intensity of the peak (height from the baseline) was corrected by calculating the height of the peak from the base of the peak. Ratios between the corrected intensity of band 1 (1735 cm⁻¹, acetyl groups in xylan and other non-conjugated carbonyls), band 5 (1508 cm⁻¹, the aromatic skeletal vibration of lignin), band 9 and 19 (1368 and 897 cm⁻¹, carbohydrates) were calculated (Figure 3). All aged samples had lower carbonyl/carbohydrate ratios (A1735/ A_{1368} and A_{1735}/A_{897}) than the reference samples. The lignin/ carbohydrate ratios (A1508/A1368 and A1508/A897) were significantly higher for hydrothermal treatments up to 160°C. The higher intensity of the aromatic skeletal vibration was probably the result of carbohydrate decomposition or the splitting of aliphatic chains in lignin and/or condensation reactions increasing the aromatic band intensity (Faix 1992; Uçar et al. 2005; Windeisen and Wegener 2008). The ratio of A1508/A1368 was slightly higher for sample 400yW, which may also be explained by the formation of a higher number

 Table 3
 Relevant FTIR ATR absorption bands average corrected intensity of the studied samples and the reference (RefW).



Figure 3 Corrected intensity ratio of the absorption bands which are characteristic for carbonyl, carbohydrate and lignin for unaged samples (RefW), 400-year-old samples (400yW) and hydro-thermally treated samples of series 1 (HtW_{180/60/30}, HtW_{160/14/50}, HtW_{160/60/30}).

		Base-line corrected band intensities							
Band			Wood aged by hydrothermal treatment (HtW _{Temp/RH/Time})						
(cm^{-1})	RefW	$400 \mathrm{yW}_{\mathrm{old}}$	160/14/50	160/60/30	180/60/30				
1735	0.451	0.344	0.310	0.003	0.017				
1700	0.000	0.000	0.020	0.375	0.506				
1640	0.212	0.163	0.013	0.010	0.011				
1616	0.306	0.500	0.295	0.466	0.661				
1508	0.800	0.860	0.789	0.805	0.860				
1462	0.089	0.080	0.085	0.088	0.074				
1452	0.097	0.109	0.106	0.107	0.118				
1423	0.421	0.469	0.407	0.356	0.342				
1368	0.438	0.442	0.426	0.316	0.319				
1334	0.052	0.075	0.055	0.084	0.089				
1316	0.215	0.291	0.213	0.306	0.302				
1264	0.527	0.633	0.477	0.442	0.444				
1230	0.069	0.062	0.052	0.045	0.050				
1204	0.091	0.145	0.110	0.177	0.260				
1157	0.592	0.790	0.781	0.799	0.776				
1103	0.366	0.563	0.613	0.657	1.033				
1051	0.528	0.775	0.728	0.841	0.958				
1026	0.977	1.044	1.037	0.853	0.900				
897	0.166	0.175	0.163	0.122	0.108				
876	0.011	0.012	0.000	0.040	0.040				
808	0.155	0.283	0.176	0.158	0.112				
664	0.196	0.238	0.189	0.229	0.277				



Figure 4 UVRR average spectra. (a) 200, 400, and 500-year-old spruce wood (200yW, 400yW, and 500yW) compared to recent sample (RefW). (b) Spruce wood hydrothermally treated samples of series 1 ($HtW_{180/60/30}$, $HtW_{160/14/50}$, $HtW_{160/60/30}$) up to 160°C compared to recent sample (RefW). (c) Spruce wood hydrothermally treated samples of series 2 ($HtW_{130/40/1}$, $HtW_{130/40/10}$) at 130°C compared with unaged sample (RefW). (d) Acetone-extracted naturally aged (ext_400yW), hydrothermally treated (ext_ $HtW_{180/60/30}$ and ext_ $HtW_{130/40/10}$) spruce wood samples compared to acetone-extracted recent sample (ext_RefW). The main absorption bands showing differences are indicated with arrows.

Table 4	UVRR	band	assignme	nt in	the 1	700 - 1	100 cn	1 ⁻¹ reg	gion f	for	different
wood spe	cies (Ag	arwal	1999; Sa	ariho	et al.	2003;	Pandey	and	Vuor	inen	1 2008).

Band	Found (cm ⁻¹)	Lit. (cm^{-1})	Compound or chemical group
1	1646	1645	Unsaturated structures of resin acids
2	1602	1605	Sym. aryl ring stretch. of G and S
3	1495	1502	Asym. aryl ring stretch. vibr. of S
4	1365	1368	Phenolic and/or cond. structure of L
5	1267	1270	Arom. ether in G
6	1150	1152	Delta phenol mode of G

G, guaiacyl; S, syringyl.

of condensed structures in the lignin of naturally aged (old) samples.

UVRRS of non-extracted samples

For each sample, an average spectrum was calculated from eight UVRRS. The band assignment based on the literature in presented in Table 4. In all cases, the absorption band at 1605 cm⁻¹ was the most prominent (aromatic ring vibrations).

The intensities of the bands within 1300–1500 cm⁻¹, attributed to the guaiacyl groups of the lignin, were higher in OldW than in RefW (Figure 4a). The small increase of the relative intensities in this region compared to the reference peak at 1602 cm⁻¹ may be as a result of a slight increase in unsaturated structures resulting from lignin degradation. In the case of UV irradiated wood, the intensity changes were similar, but more intense (Pandey and Vuorinen 2008). The 400yW sample showed a band at 1365 cm⁻¹ with slightly higher intensity in comparison with the spectra of 500yW and 200yW. Consequently, differences observed between the OldW spectra are not related to the age of the sample. Other ageing factors (such as changes in moisture content or mechanical loading), independent of time, may be responsible for the small differences observed. As the 400yW specimen was taken from a floorboard, it could well have been more mechanically stressed than the 200yW or 500yW samples, which were taken from roof beams. Repeated adsorption and desorption resulting from washing and drying of the floor might be also be a reason for such differences (Froidevaux et al. 2011).

Changes in the spectra of HtW samples treated up to 160°C (Figure 4b) were similar to the changes observed in thermally treated wood, but were less pronounced (Nuopponen et al. 2005). There was a significant increase in the absorption band at 1365 cm⁻¹ and a significant decrease in the band at 1262 cm⁻¹ for treatment types HtW_{180/60/30} and HtW_{160/60/30}. This indicates that guaiacyl structures were modified at high temperatures. Moreover, the band at 1646 $\rm cm^{\text{-1}}$ was decreased and the bands at 1602 $\rm cm^{\text{-1}}$ and in the region 1700–1550 cm⁻¹ were broadening for the samples HtW_{180/60/30} and HtW_{160/60/30}. Pandey and Vuorinen (2008) observed the same intensity changes in UV irradiated spruce and explained it by the formation of o- and p-quinone type structures. The same was true for thermomechanical pulps (Agarwal 1998; Jääskeläinen et al. 2006). Treatment type HtW_{160/14/50} exhibited the same characteristics but were less pronounced than the treatments HtW180/60/30 and HtW160/60/30.

The UVRR spectra of the HtW samples of series 2 at 130°C (HtW_{130/40/1}, HtW_{130/40/2}, and HtW_{130/40/10}) were similar to RefW (Figure 4c). Only a slight increase in the band at 1365 cm⁻¹ was visible. The lignin structures did not change following this treatment compared to RefW.

UVRRS of acetone extracted samples

Soxhlet extraction should help defining which lignin band is overlapping with acetone extractable substances. The UVRR spectra of the acetone extracted samples, RefW, 400yW, $HtW_{180/60/30}$, and $HtW_{130/40/10}$ are presented in Figure 4d.

With the exception of the $HtW_{180/60/30}$ spectra, all spectra are nearly identical and the slight differences observed on nonextracted samples are not perceptible. Consequently, the slight UVRR spectral differences described above (between OldW, RefW, and HtW) were caused by acetone-extractable compounds. These are probably lignans or lignin degradation products (Rogers et al. 1967; Kawamura et al. 1998). Such compounds are also generated during the photo-discoloration of wood under UV-light. As the specimens were taken from layers not penetrated by UV-light, the amount of acetonextractables could increase as a result of a sub-surface degradation caused by longer wavelength light that still has sufficient energy to degrade wood (Hon et al. 1980; Kataoka et al. 2007).

PCA of UVRR spectra

The UVRR spectra of all non-extracted samples were evaluated by PCA. The first principal component (PC1) explained 49% and the second principal component (PC2) explained 33% of the differences between the spectra. Other successive PC explained no more than 5% of the dissimilarities and, consequently, the analysis was limited to PC1 and PC2. The plotted scores of PC1 vs. PC2 (Figure 5) shows the two largest variations in magnitude within the data and the spectra are separated on the plot. Spectra of the sample HtW_{180/60/30} and HtW_{160/60/30} form individual groups which were well separated from the other samples. They both had positive values of PC1 and PC2, which was not the case for other groups of compounds. The other groups were overlapping with each other, which means that they were more similar. The position of OldW (PC1>0 and PC2<0) is opposite



Figure 5 Principal component analysis from UVRR spectra collected for all sample sets.



Figure 6 (a) Loading of the first principal component (PC1) of the UVRR spectra for all sets. (b) Loading of the second principal component (PC2) of the UVRR spectra for all sets.

to that of RefW (PC1<0 and PC2>0). Nevertheless, careful interpretation is needed, as the groups are partly overlapping.

The loadings plot of PC1 and PC2 provides information on the bands which are responsible for grouping of the samples. The loading for PC1 (Figure 6a) shows that bands at 1627 cm⁻¹, 1567 cm⁻¹ (*o*- and *p*-quinone type structures) and 1368 cm⁻¹ (phenolic and/or condensed structures of lignin) are mainly responsible for the grouping of sample types HtW_{180/60/30} and HtW_{160/60/30}. The loading of PC2 (Figure 6b) reveals that the band at 1620 cm⁻¹ (*o*- and *p*-quinone type structures) gives a positive loading and is also responsible for the grouping of samples in discussion. Grouping of the OldW samples (PC2<0) seems to be caused by the bands at 1582 cm⁻¹ (broadening of the absorption band at 1602 cm⁻¹) and at 1274 cm⁻¹ (aromatic ether units in guaiacyl structures).

Conclusions

Spectroscopic data of naturally aged wood samples are not changed to a great extent. According to FTIR spectra, hemicelluloses are deteriorated in naturally aged spruce samples. The UVRR spectra of naturally aged wood show that the small differences compared with a recent reference wood are caused by acetone-extractable compounds. These compounds could include degradation products of lignin or unsaturated wood extractives. No specific correlation could be found between the age from 200 to 500 years and spectral changes, confirming that time alone is not the only factor affecting the chemistry of wood during ageing. Additional factors, such as humidity, insects, fungi and the origin of the wood influence the changes. For artificial ageing by hydrotreatment up to 160°C, the changes in lignin were significant, as is the case for thermally treated wood. Samples from hydrotreatment at 130°C do not differ from the reference sample. Probably, the analytical power of FTIR and UVRRS is limited, and thus additional methods, such as ¹³C NMR or wet chemical analyses, are needed to gain more insight into ageing.

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