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- 1 Short title: Wood aging observed by ¹H NMR relaxometry
- 2 3

Investigation of the effect of aging on wood hygroscopicity by 2D ¹H NMR relaxometry

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- Abstract: 2D ¹H NMR relaxometry is increasingly used in the field of wood sciences due to its great 14 potential in detecting and quantifying water states at the level of wood constituents. More precisely, 15 in this study, this technique is used to investigate the changes induced by "natural" and "artificial" 16 aging methods on modern and historical oak woods. Two bound water components are detected and 17 18 present differences in terms of association to the different wood polymers in cell walls: one is more strongly associated to wood polymers than the other. The evolution of the two bound water types is 19 discussed in regard to aging methods and is related to the structure of the cell wall, especially with 20 21 the S2 layer and the evolution of wood chemical composition (cellulose, hemicelluloses and lignin). The evolution of hydric strains is discussed taking into account the effect of aging methods on the 22 two bound water components too. The obtained results confirm the ability of 2D NMR relaxomety to 23 24 evaluate the effect of aging at the molecular level and on hydric deformation. Furthermore, this method shows that it is possible to determine the moisture content of wood without the necessity to 25 oven-dry the wood material. 26
- 27
- Keywords: Aging, deformation, extractibles, hydric cycles, oak wood, thermal treatment, 2D NMR
 relaxometry
- 30

31 Introduction

The built heritage shows that wood makes it possible to design sustainable and healthy buildings (Epaud 2007, Obataya 2007, Froidevaux 2012). Wood, as an envelope or structural material, is a material of interest in the field of construction both for new buildings and for the question of renovation of old buildings (Froidevaux 2012). However, to estimate lifetimes or to propose methods of maintenance, it is necessary to continue the research efforts to understand the mechanisms of physical aging of wood and its hygro-mechanical properties at the scale of its constituents.

Aging is a complex phenomenon that modifies physical, chemical and mechanical properties of a 38 polymeric based material, under the effect of its proper instability, environmental parameters or 39 mechanical strains (Fayolle and Verdu 2005). As a natural and biodegradable resource, wood is 40 mainly constituted of three biopolymers: cellulose, hemicellulose and lignin. These components can 41 42 be vulnerable to environmental factors such as temperature, solar radiation, humidity, which can degrade the wood's polymeric structure. Studies have shown that the degradation due to aging mostly 43 affect the hemicelluloses (which are the most vulnerable polymers), resulting in a better dimensional 44 45 stability of the wood material (Kranitz 2014, Gauvin 2014). A decrease of cellulosic material as well 46 as a slight decrease in lignin content are also observed in some historical woods (Kranitz et al. 2016). Heat treatments at high temperature show a common effect with natural aging on decreasing the 47 48 Equilibrium Moisture Content (EMC), leading to an improved dimensional stability of wood and are usually considered as artificial aging. Numerous studies have been conducted to develop industrial 49 50 processes varying in terms of temperature (generally at high temperature between 160°C and 280°C), duration and vector (gas, water vapour, oil...) (Tjeerdsma et al. 1998, Esteves and Pereira 2008, 51 52 Wentzel et al. 2018) to improve the durability of timbers. The higher the temperature of treatment -53 and the longer the treatment time - the greater the dimensional stability of wood (Inari et al. 2009; Chaouch 2011; Endo et al. 2016). However, these treatments' intensity considerably weakens the 54 mechanical properties of timbers (Hill 2006, Froidevaux 2012, Candelier 2016). Thus, thermal 55

treatments at lower temperature are proposed as a better alternative as they have a lower effect on the 56 57 mechanical properties of timber. However, it remains unclear if the thermal treatments in presence or absence of humidity is the closest to a 'natural aging' effect. Some authors show that hydrothermal 58 59 treatments in precise conditions of Relative humidity, Temperature and Pressure can mimic an aged wood in terms of its mechanical characteristics, colour change and decreased swelling rate 60 61 (Froidevaux 2012, Gauvin 2015, Endo et al. 2016). Whereas in terms of dry heat treatments, it seems 62 to be a better option than steam to reduce the hygroscopic capacity of wood, and improve its dimensional stability, with a minimum of thermal degradation (Obataya 2007, Matsuo et al. 2009, 63 Sandberg and Navi 2013). 64

65 The wood hygroscopicity plays a major role on the dimensional stability and mechanical properties of wood. This improved dimensional stability may be explained by a decrease in adsorption sites in 66 the polymers chains of the wood cell walls (Hill 2006, Froidevaux 2012, Murata et al. 2013). 67 68 However, in Rautkari et al. 2013, a poor correlation between EMC, sorption sites accessibility and theoretical hydroxyl group content was found following a thermal treatment and therefore additional 69 70 mechanisms should exercise control over the EMC. This shows that water interactions with wood (hygroscopicity and dimensional stability) are still not completely understood, which can be 71 explained by the difficulty to characterize adsorbed water in wood. The use of proton Nuclear 72 Magnetic Resonance (¹H NMR) relaxometry to detect bound water in the wood's cell walls is gaining 73 74 popularity in the research field as it is a non-invasive method allowing to study the same material subjected to different loadings. Here, we investigate the state of water adsorbed on the wood's cell 75 walls through ¹H low-field (LF) NMR relaxometry (Menon et al. 1987, Araujo et al. 1994, Labbé et 76 77 al. 2002, Fredriksson and Garbrecht Thygesen 2017, Beck et al. 2018). Recent studies (Cox et al. 2010, Bonnet et al. 2017) have used two-dimensional (2D) T₁-T₂ correlation spectra, allowing to 78 79 differentiate two different types of bound water, which have been assigned to water adsorbed to different wood polymers in the hygroscopic range (Bonnet et al. 2017). So far, this technique is the 80

sole tool capable to observe two different types of bound water in wood. Thus, 2D ¹H NMR correlation spectra is a great method that can provide an insight on the water environment inside the lignocellulosic matter and hints the influence of the microstructure and chemical composition on the hydrogen signal observed. Moreover, the influence of the two bound water components on dimensional properties has to be further investigated. According to Cox et al. (2010), only one component may contribute to wood swelling, whereas for Bonnet et al. (2017), the two components have an effect on the hydric strains.

The main purpose of this study is to analyse the effect of two aging methods on the hygrocopicity of 88 wood material by using the 2D ¹H NMR relaxometry to evaluate the interactions between water and 89 90 the cell wall's physical and chemical changes that may have occurred. A method to evaluate the 'true dry mass" of wood material is also proposed, enabling the calculation of the "true equilibrium 91 moisture content" without the need of drying the samples at 103°C in an oven (which can lead to 92 93 irreversible damages or misinterpretations). The evolutions/modifications are also discussed against hydric strains, as well as the possible contribution of the two water components on the volume 94 95 deformation. A parallel objective of this work is to compare these changes between historical and modern oak woods. 96

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98 Materials and Methods

99 **Materials:** The experiments were carried out on modern and historical oak wood samples to compare 100 the effect of aging on two types of wood. The historical wood (350-year-old oak) was provided by 101 Atelier Perrault, and comes from a wooden door's frame of an old building of the 17th century in Saint 102 Georges street in Rennes, France. The historical oak wood has been subjected to natural aging. The 103 modern oak wood was recently provided from a sawmill located in the North of France and may be 104 considered without aging. Figure 1 provides an optical microscopy image of the transversal cross-105 section (plane RT) of both wood materials. The chemical composition and the proportion of the different kinds of cells (vessel, fiber, parenchyma...) were not quantified in this study, but theseimages allowed confirming differences between these two oak materials.

Small specimens of approximately 1 cm³ were prepared as 1 cm is the maximum height for optimal 108 NMR measurements. This was done using a band saw and they were all clear from visible defects. 109 They were sawn along the anisotropic directions (Longitudinal L, Radial R and Tangential T). All 110 the samples were taken side by side, to minimize the variability between samples of the same type 111 (modern wood or historical wood). The average density of all specimens was measured at 2% relative 112 humidity (RH) and is 0.64 (\pm 0.02) g cm⁻³ for modern oak wood and 0.47 (\pm 0.02) g cm⁻³ for historical 113 oak wood. The lower density of the historical wood is in accordance with the fact that there are more 114 vessels. 115

All samples were subjected to a common cycle of adsorption-desorption to equalize their hydric
history, using saturated salt solutions at 2% RH (silica gel) and 97% RH (potassium sulphate: K₂SO₄).
They were kept in desiccators until constant mass (about one week). All the experiments were carried
out at 20°C and on an adsorption cycle (starting from 2% RH) to avoid the influence of the hysteresis
in the observed phenomena.

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Aging protocols: Two aging methods were conducted, on the one hand repeated hydric cycles (considered as a "natural aging" method with gentle impact) and on the other hand mild thermal treatment (considered as an "accelerated aging" method). To compare the effect of these aging methods, the samples were all characterized through 2D ¹H NMR at equilibrium moisture content (Eq. 1):

$$EMC_{x\%RH} = \frac{M_{x\%RH} - M_{dry}}{M_{dry}} \times 100$$
(1)

where $EMC_{x\%RH}$ is the equilibrium moisture content at x% RH [%], $M_{x\%RH}$ the mass of the equilibrated sample at x% RH [g] and M_{dry} the mass of dry sample [g].

Repeated hydric cycles consist in applying repeated cycles of adsorption and desorption to wood 129 130 specimens. In this study, three samples of modern wood and three samples of historical wood were subjected successively to 2% RH and 97% RH using the method of saturated salt solutions to attain 131 EMC at room temperature (20°C). These hydric cycles were repeated during 6 months (corresponding 132 to a total of 12 cycles of adsorption-desorption) and samples were studied at 1, 3 and 6 months. In 133 order to study the evolution of the samples at each period, they were conditioned at 65% RH, 20°C 134 (using sodium nitrite (NaNO₂)) and analysed through ¹H NMR at EMC. In parallel, control samples 135 of modern and historical wood (three of each) were conditioned at 65% RH, 20°C, over the entire 136 period of the experiments. Deformation measurements and mass measurements were carried out 137 138 regularly on all samples to assess the EMC by weighing and the evolution of the hydric strains.

Thermal treatments (TT) were conducted in an oven at a temperature of 120°C (where RH is close to 139 0%) for three periods of time (24 hours, 3 days and 7 days). Prior to the TT, the initial state of each 140 specimen conditioned at 65% RH was characterized through 2D ¹H NMR at EMC at 20°C. After TT, 141 the specimens were again conditioned at 65% RH, 20°C for one week before a second ¹H NMR 142 143 analysis. This protocol allows to compare the EMC evolution for the same sample and to determine the mass loss induced by the TT. Dimensions were also measured before and after treatment to assess 144 145 the change in volume deformation as this provides information on the stability of the wood. Three 146 specimens of each wood were considered for the 24 hours TT to take into account the variability of wood material and to test the reproducibility and the repeatability of the used method. As the 147 variability was statistically tested with a Student T-test and showing significant changes after 24h TT, 148 only one specimen of each wood type was analysed for the 3 and 7 days TTs. 149

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151 NMR methods: The samples were measured by 2D ¹H NMR and to do so they were inserted in 152 18mm NMR tubes. The RH was controlled with a saturated salt solution at 65% RH during the NMR 153 experiments (Bonnet et al. 2017, Fourmentin 2015) placed in a small container on the top of the NMR

tubes. As the temperature of the magnetic unit is 40°C, a cooling system is used to maintain the 154 samples at 20°C (at the bottom of the NMR tubes). The temperature in the area of the saturated salt 155 container (outside the instrument) corresponds to a temperature of 25±2°C (measured during NMR 156 analysis). The theoretical variation of the saturated salt solution used is up to 2% decrease of RH at 157 30°C. To take into account possible variations on EMC, all samples were weighed before and after 158 NMR measurements and the mean value of these two measurements was used to calculate the EMC. 159 160 The difference in EMC calculated with the mass before NMR and the mass after NMR measurement was evaluated and resulted to be lower than 0.1%. Therefore, it is considered of a minor effect and 161 does not affect the obtained results and interpretations. The device used was a BRUKER MINISPEC 162 163 MQ20 spectrometer that operates at 0.5 T, corresponding to a resonance frequency of 20 MHz for 1 H. 164

The principle of NMR is based on the intrinsic nuclear spin that some atomic nuclei have (with an 165 166 odd number of protons, neutrons, or both). When an atomic nucleus with a nonzero spin is placed in a magnetic field \mathbf{B}_0 , the nuclear spin can be aligned in the same direction or in the opposite direction 167 to the field. These two types of nuclear spin alignment are characterized by different energies, and 168 their sum is called the magnetization (\vec{M}) . When a second magnetic field (with a characteristic energy) 169 **B**₁, perpendicular to **B**₀ is applied as a pulse, this magnetization is disturbed and the relaxation (=time 170 needed for the magnetization to come back to its initial state) is characteristic of the atom and of the 171 local environment (Kekkonen 2014). The two main relaxation times measured are: T₂, the spin-spin 172 or transversal relaxation time and T₁, the spin–lattice or longitudinal relaxation time. 173

The Inversion Recovery (IR) sequence coupled with the Carr-Purcell-Meiboom-Gill (CPMG) sequence (Carr and Purcell 1954, Meiboom and Gill 1985) was used to obtain 2D NMR correlation spectra of the T_1 and the T_2 relaxation values. The calculation of T_1 - T_2 correlation spectra from NMR data was performed with an in-house software, which essentially reproduces the 2D-Inverse Laplace Transform (ILT) algorithm of Song et al. (2002). For more details about this and about the acquisition method and parameters, see Bonnet et al. 2017. The T_1 - T_2 correlation spectra allow to show the NMR signal as a function of the two relaxation times, T_1 and T_2 , obtaining two different peaks for the adsorbed water (which is not possible with 1D T_1 or T_2 spectra due to peak overlapping). The volume under each peak of the 2D spectra is proportional to the amount of hydrogen (H) atoms. The T_1 and T_2 values of each peak are determined by means of the coordinates of their respective maxima. 2D spectra also give unambiguous access to T_1/T_2 ratios. The latter is characteristic of H atoms mobility and confinement and greatly helps spectrum interpretation.

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Determination of EMC: The moisture content can be calculated from the T_1 - T_2 spectra, as the area under the peaks is proportional to the amount of H atoms. Thus, the total moisture content can be evaluated through NMR as the sum of the area under the two peaks corresponding to bound water. In order to convert the NMR signal into a mass of adsorbed water, a standard curve is performed with different quantities of water. The bound water mass ($M_{NMR,x\%RH}$) determined by ¹H NMR at x% RH is defined by (Eq. 2):

$$q_{\rm NMR,x\%RH} = \alpha. M_{\rm NMR,x\%RH}$$
(2)

193 Where $q_{NMR,x\%RH}$ is the NMR signal intensity of peaks of interest and α the proportional coefficient 194 determined through the standard curve (in this study, $\alpha = 164.77$ NMR signal/g of water).

To calculate $EMC_{NMR,x\%RH}$, it is necessary to determine the mass of the dry sample, M_{dry}. To avoid irreversible structural damages or misinterpretations (in particular for thermal treatment), the studied materials are never oven-dried at 103°C as usually done before heat treatments (Rajohnson 1996, Obataya 2007, Candelier 2013). Therefore, ¹H NMR can be of good use to determine the "true dry weight" of wood. Total mass of wood is weighed at EMC with a scale of precision ±0.0002g and the amount of bound water at EMC is calculated through ¹H NMR as depicted previously. Thus, the mass of dry wood M_{dry} [g] can be calculated as follows (Eq. 3):

$$M_{dry} = M_{w,x\%RH} - M_{NMR,x\%RH}$$
(3)

Where $M_{w,x\%RH}$ [g] is the total mass (dry wood and bound water) determined by weighing at x% RH (which corresponds to the average value of weighing before and after NMR experiments) and $M_{NMR,x\%RH}$ [g] is the bound water mass determined by NMR at x% RH. The moisture content determined through ¹H NMR measurements (EMC_{NMR,x%RH}) is defined as $M_{NMR,x\%RH}$ divided by M_{dry} . The moisture content determined by weighing at x% RH (EMC_{w,x%RH}) is defined as $M_{w,x\%RH} - M_{w,2\%RH}$ divided by $M_{w,2\%RH}$. These two moisture contents verify the following relationship (Eq. 4):

$$EMC_{NMR,x\%RH} = EMC_{w,x\%RH} \frac{M_{w,2\%RH}}{M_{dry}} + \frac{M_{w,2\%RH} - M_{dry}}{M_{dry}}$$
(4)

The obtained data for $\text{EMC}_{\text{NMR},x\%\text{RH}}$, $\text{EMC}_{w,x\%\text{RH}}$, M_{dry} and $M_{w,2\%\text{RH}}$ are given in SI. Uncertainties on measurements for EMC are determined through the standard deviation values; they take into account the possible variability between samples' results and the precision of the balance (0.36% for the modern wood and 0.28% for the historical wood).

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Volume measurements: In order to assess the changes in regard to the two aging methods, the samples' dimensions were measured with a Mitutoyo electronic calliper with a precision of \pm 0.01mm. These measurements were performed at each step of the experiments for the three directions R, T and L allowing the calculation of the samples' volume: during the conditioning, at EMC, right before and after NMR measurements.

218 Measurements of the samples' dimensions allow to determine the deformation between two states, 219 2% RH and x% RH (with x=65% or 97% RH). Hydric strains at x% RH ($\varepsilon_{x\%}$) are related to the state at 2% RH, as the volume of the dry wood was not experimentally measured as previously explained,
according to (Eq. 5):

$$\varepsilon_{\rm x\%} = \frac{V_{\rm x\%RH} - V_{\rm 2\%RH}}{V_{\rm 2\%RH}} \times 100 \tag{5}$$

222 Where $V_{x\%RH}$ and $V_{2\%RH}$ are the volumes of a sample at x% RH and 2% RH respectively.

Note that the determination of the volume deformations was done using the state at 2% RH right
before the conditioning at 65% RH for samples subjected to hydric cycles.

Uncertainties on deformation measurements are determined through standard deviation values; they take into account the possible variability between samples' results and the precision of the electronic calliper. For these experiments, the variability is about 0.39% for the modern wood and 0.30% for the historical wood. Also note that the EMC values for hydric strains may slightly differ from those during the NMR experiments due to possible variations of RH and temperature and because the dimension measurements are performed outside the desiccators.

231

232 **Results and discussion**

T₁-T₂ spectra and peak assignment: Figure 2 shows T_1 - T_2 correlation spectra recorded from historical and modern oak wood samples (only one sample of each wood is represented here), before aging processes. Regardless of small differences in the shapes of the peaks observed in the cartographies, the global scheme is similar for both wood materials.

To interpret these spectra, it is necessary to understand the influence of water mobility and environment on the relaxation times T_1 and T_2 . T_1 and T_2 depend on the local environment of the measured hydrogen atoms (H) (i.e. from water molecules and wood polymers in this work), that is to say: size of the molecules they are bound to, affinity with molecules interacting and pore size. T_2 relaxation time decreases as pore size is reduced. Moreover, T_2 will also be shortened when molecular tumbling (mobility) decreases. If H atoms are in molecules with restricted mobility due to viscosity,

or the interactions being in macromolecules or solid phase, T₂ values will decrease gradually 243 244 depending on the degree of this restriction. The case of T_1 is a bit more complicated than T_2 as it first decreases when molecular tumbling decreases (from liquid phase) but it increases when molecular 245 tumbling is low (in gel and solid like molecules). Fortunately, T_1/T_2 ratios are characteristic of the 246 mobility of H atoms and greatly help spectrum interpretation. Depending on the value of these ratios, 247 248 water can be assigned to unconfined water, mobile water in a free or adsorbed state and bonded 249 immobile water molecules and solid macromolecules. It can be noticed that T_1/T_2 ratios lower than one are physically not permitted (Bonnet et al. 2017). 250

Based on this knowledge, the peaks observed in the T₁-T₂ spectra can be identified based on analysis 251 252 previously done in Cox et al. (2010) and Bonnet et al. (2017). Peak A is labelled as liquid water in 253 the lumen as it has the highest T_1 and T_2 values and it is close to the $T_1=T_2$ line. In some spectra, this 254 peak is located slightly above the $T_1=T_2$ line, which certainly owes to the way the signal acquisition 255 was made. Indeed, the CPMG period of the NMR sequence was deliberately truncated after 12ms to prevent excessive radio frequency (RF) power deposition in the sample and thus avoid heating the 256 sample. As a result, the measurement of T₂ value for peak A is not complete and lacks accuracy, 257 leading to a widening and 'mislocation' in T₂ direction. In our case, the amplitude of the peak is also 258 very low and it is therefore not of interest in this study. Peaks B and C correspond to bound water in 259 260 two specific and distinct water reservoirs in the wood cell walls. The T_1/T_2 ratio is higher for peak B than for peak C, indicating a more restricted molecular motion in compartment B and thus more 261 strongly bonded water molecules in that compartment. In contrary, the low T_1/T_2 ratio for peak C 262 indicates less restricted molecular motion and less strongly bonded water molecules. In some spectra, 263 peak C seems to split into two peaks, one with lower T₁ and T₂ values than the other peak but having 264 almost the same T_1/T_2 ratio. This could be interpreted as two types of C-water with small differences 265 on their environment (e.g. surface-to-volume ratio). In general, the peak with higher T_2 and T_1 values 266 is most intense, but in few cases, it is the other one showing higher amplitude. Thus, for easier 267

interpretation, the peak taken into account for the calculation of T_1 and T_2 values of peak C is always the one with higher T_1 and T_2 values. It would be interesting to understand the nature of this splitting but it is out of scope of this work. Finally, peak D represents H atoms from the wood polymers since the T_2 value relaxes very fast (in the order of μ s).

Among the recent results, Bonnet et al (2017) have proposed to relate the two components B and C 272 to the structure of the S2 layer of the cell wall, because this layer may explain the mechanical 273 274 properties and especially the hydric strains. Based on a schematic structure of the S2 layer inspired by Boyd (1982) and Salmén and Burgert (2009), it has been explained that the population of water 275 from component B (strongly bonded) may be located in the macrofibrils composed mainly of 276 277 microfibrils of crystalline and amorphous cellulose embedded in a matrix of hemicelluloses (primarily glucomannan for softwoods). While the population of water from component C (weakly 278 279 bonded) may be in a lignin and hemicelluloses (xylan) matrix corresponding to the intra-macrofibril 280 region. The attribution of the two bound water components B and C to these two regions is related to the fact that water affinity is higher in the region composed of cellulose and hemicelluloses than in 281 the region composed of lignin and hemicelluloses. However, the proportion of polymers differ 282 between hardwoods and softwoods (Chaouch et al. 2010). Hardwoods contain mainly glucuronoxylan 283 and low amount of glucomannan. Thus, for our studied materials, it can be suggested that xylan 284 285 (glucuronoxylan) may be present in the two regions.

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NMR results for samples at the initial state: Table 1 presents the average results obtained from NMR measurements on the control samples of hydric cycles and on the samples at the initial state before TT. EMC values (deduced from the sum of peaks B and C) are the same order of magnitude for both woods. Furthermore, peak B reveals a higher moisture content than peak C for both woods, meaning that this compartment contains more adsorbed water than the other one. Compartment C

shows higher water content for the historical wood than for the modern wood, while there is lesswater in compartment B for historical wood.

294 These differences in terms of relative moisture content in B and C could provide new insights in terms 295 of evolution in the structure and composition of the wood cell wall. The higher moisture content in compartment C of the historical wood appears to indicate a higher amount of available adsorption 296 297 sites in that region compared to the modern wood. The higher content of adsorption sites could be 298 due to an evolution of the polymeric structure such as a change in the polymers conformation. Indeed, the lignin undergoes structural changes due to oxidation during aging but, as shown in Kranitz 2014, 299 300 the absolute amount remains the same. Therefore, these structural changes in the lignin could free 301 some previously inaccessible hydroxyl sites for water adsorption.

NMR results give an additional indication in favor of the hypothesis made on the local environment 302 303 of water in compartment B, which is mainly composed of cellulose and hemicelluloses. As mentioned 304 earlier, the hemicelluloses are the first polymers to degrade during wood aging and a decrease in amorphous cellulose is observed in naturally aged woods (Kranitz 2014), thus leading to a decreased 305 306 number of adsorption sites and therefore lower moisture content in that region for historical wood compared to the modern wood. However, it may be difficult to quantitatively compare these two oak 307 wood types without chemical characterization. This is a common issue found when historical and 308 309 modern wood are to be compared, as the composition and proportion of polymers in the cell wall before aging are not known. The environmental conditions and geographical location are also a source 310 of variability in the wood development and evolution. Thus, to improve our understanding of the 311 relative differences on the amounts of B and C 'water' between historical and modern oak wood, a 312 313 detailed chemical analysis could be useful (e.g. characterization and quantification of wood polymers) in future works. 314

316 **NMR results for repeated hydric cycles:** Table 2 presents the moisture content measured during 317 the repeated hydric cycles as well as the characteristic relaxation times T_1 and T_2 for all samples. A 318 graphical representation of the evolution in total moisture content is also presented in Figure 3.

First observation shows that hydric cycles did not really affect the adsorption behaviour of the 319 samples, meaning that no significant increase or decrease in EMC is observed in regards to the 320 standard deviation values. A paired Student T-test was performed on all the moisture content 321 measures (EMC_{NMR,65%RH}(B + C), EMC_{NMR,65%RH}(B) and EMC_{NMR,65%RH}(C)) showing no significant 322 evolution of the EMC (p-values > 0.05) along the 6 months of repeated hydric cycles. Moreover, the 323 evolution of the T₁/T₂ ratio is globally constant. This indicates no change in the mobility of water 324 pools, thus no change in the polymeric structure of the cell wall, i.e. no change in the hygroscopicity 325 326 of the wood material. Such results were expected since the room temperature and humidity are fixed 327 and controlled along the experiment, which is not the case in natural conditions of aging. Furthermore, mass of samples was recorded during the cycles and permits to estimate the mass loss through the 6 328 months cycles, showing no significant mass loss through the whole experiment period. 329

In conclusion, although these repeated hydric loads may have an influence on the hydric properties over the long term, we suppose that the duration of tests is not sufficient to induce a significant effect on the hygroscopic properties. This result will be discussed with the hydric deformation measurements below.

334

NMR results for moderate heat treatment: NMR data are presented in Table 3 and a graphical representation of the evolution in total moisture content before and after the heat treatments is presented in Figure 3.

A noticeable loss in total $EMC_{NMR,65\%RH}(B + C)$ is observed for both modern and historical wood samples and the longer the treatment, the higher the loss in hygroscopicity, as expected. Thermal treatments also show a total higher effect on the historical wood than on modern wood. There is a higher EMC loss for historical wood the longer the treatment time (13% for modern wood against
19% for historical wood, in average of the 72 h and 168 h TT). The loss of water in compartments B
and C can be calculated separately, showing for the 24 h TT a higher percentage of water loss in peak
C (18.0%) than peak B (6.9%) for historical wood. While it seems to be the contrary for modern oak
wood, with higher loss in compartment B (11.5%) than C (9.9%). The same conclusions were
obtained for the 72 h and 168 h TTs.

347 A Student T-test was performed on these percentages of loss in EMC for the 24 h TT, showing no significant difference between the loss in EMC_{NMR.65%RH}(B) and EMC_{NMR.65%RH}(C) for modern wood. 348 349 For historical wood, the statistical tests confirmed a significant difference between peaks B and C, with a higher loss for EMC_{NMR.65%RH}(C). As mentioned previously, the strongly bound water 350 (component B) may be located in the macrofibrils (cellulose-glucuronoxylan-glucomannan matrix) -351 and the weakly bound water in the lignin-glucuronoxylan matrix (component C). It is suggested in 352 Salmén and Burguert (2009) that there might be a highly acetylated xylan which is closely associated 353 to a less condensed type of lignin, whereas low substituted xylans are associated with cellulose and a 354 condensed type of lignin. It can be noticed that xylan units of hemicelluloses in hardwoods are 355 generally strongly acetylated, generating a higher sensitivity of these materials to thermal treatments, 356 as the xylan units could lead to higher kinetic of thermo-degradation in hardwoods (Chaouch et al. 357 2010). Therefore, one could assume that the highly acetylated xylan are linked with the component 358 C, while the low-substituted xylan is linked to the component B. These associations could be of major 359 importance in regard of the selective evolution of wood adsorption. 360

For the historical wood, it seems that the dry mass loss (Table 4) is greater with treatment time. Indeed, the dry mass loss for 72 and 168 h TTs was higher than for the 24 h treatment, but as only one sample was used for these two durations, observations are mostly qualitative. For the modern wood, the same observation was made except for the 168 h TT that seems to be an outlier due to its low value, but which is due to the fact that it is the dry mass loss of a single sample. Recent tests confirmed an increased dry mass loss with treatment time for modern wood, up to 2.3%. Nevertheless,
these percentages of dry mass loss are quite low and according to Kollmann and Fengel (1965) and
Esteves and Pereira (2008) wood degradation begins only at 130-150°C for oak wood. Thus, by
heating at 120°C, there should be no degradation of polymers and the dry mass loss observed may be
due to a loss of extractives during heat treatment as discussed in Esteves and Pereira 2008.

371 In our case, the extractives degradation cannot fully explain the loss in EMC observed after TT, 372 knowing that not all the extractives are volatile (Hillis 1971, Esteves and Pereira 2008, Jankowska et al. 2017). Indeed, the T_1 - T_2 correlation spectra reveal a global increase of the T_1/T_2 ratio after thermal 373 treatments (Table 3), indicating that there are stronger bonds between adsorption sites and water 374 375 molecules. The ratio increase also indicates the confinement of the H atoms detected. The evolution of H atoms mobility could be due to a restricted space between polymers therefore changing the local 376 377 environment of H atoms. When wood is oven-dried in a dry atmosphere, various chemical changes 378 take place producing a "shrunken" state for which the intermolecular space is minimized (Obataya 2007). Thus, when wood is reconditioned at 65% RH, the restricted intermolecular space will not 379 380 allow the adsorption of the water molecules in several layers as it used to before TT. This hypothesis can also be interpreted by looking at the evolution of peak C for instance (Figure 4), which can be 381 divided into two parts to highlight two populations of water. The lower part corresponds to the least 382 mobile H atoms and therefore the most strongly linked water molecules (directly linked to the 383 adsorption sites). The upper part corresponds to the most mobile H atoms and therefore the least 384 strongly linked water molecules (upper additional layers of adsorbed water) (see Figure 4b). 385 Following the TT, peak C is shifted downward (T₂ decreases and T₁ increases) showing a larger 386 population of H atoms more strongly bound to the polymeric matrix, but there has been a loss of the 387 less strongly bound H population. Consequently, there is a loss of additional layers of adsorbed water 388 389 molecules due to the smaller intermolecular space following TT. Note that this observation is similar for peak B. The shrinking of the polymeric matrix could be a plausible explanation for the increasein the confinement of bound water molecules and therefore for the decrease in water adsorption.

392

Discussion in relation with deformation: The hydric strains of studied materials either subjected to 393 repeated hydric cycles or to thermal treatments are given in Figure 5 and are compared to those 394 without aging (control samples and samples at the initial state). They are expressed as a function of 395 total EMC (EMC_{NMR.65%HR}(B + C)) determined by ¹H NMR from the components B and C. Note 396 that, if the hydric strains (related to the state at 2% RH) are plotted against EMC (related to dry mass) 397 398 it is observed that the hydric strains for modern and historical wood materials vary linearly with the moisture content, as expected for wood materials (see SI for more details). The slope β of the obtained 399 400 linear law expressed as a function of EMC is 0.50 for modern wood 0.45 for historical wood. It can be noticed that the value of the slope is related to the density of the studied materials (Bonnet 2017) 401 402 and the slope β decreases with decreasing density. According to the above results (see Table 2 and Table 3), the densities of the samples after "natural" or "artificial" aging do not significantly vary, so 403 the slope remains the same after aging. Thanks to this linear law between hydric strains and EMC, it 404 405 is possible to discuss the influence of each bound water compartments (B or C) on the decrease of 406 volume deformation. This evolution is related to the decrease of EMC(B) and EMC(C) which is 407 related to the decrease of EMC(B+C) for samples subjected to aging, see SI for more details.

The hydric strains for samples subjected to repeated hydric cycles seem to be "globally" stable through the 6 months cycles (Figure 5) for modern and historical wood materials. This is in coherence with a "globally" stable EMC. Note that the hydric strains of the samples at one month are higher than those of the control samples, in coherence with higher EMC (Table 2). Between 1 month and 6 months cycles there is surprisingly a slight decrease of the volume deformations with no decrease of the EMC. Further studies are necessary to confirm these unexpected results (at other durations). Moreover, it seems to be more important for modern wood than for historical wood (Figure 5) in 415 coherence with the fact that the hemicelluloses are supposed to be mostly degraded in historical wood
416 (Kranitz 2014, Obataya 2007), thus leaving a material more stable in terms of volume deformation.

Concerning the samples subjected to thermal treatments, the hydric strains significantly decreased 417 after TT and with treatment duration. However, the effect seems to be weaker for the modern wood 418 than for historical wood. The contribution of each component (B or C) on the hydric strains is not the 419 same for both studied materials. For historical wood material, the contribution of compartment C on 420 421 volume deformation decreases with treatment time while the contribution of component B increases (see Table 5). Furthermore, the component C contributes the most in the loss of volume deformation 422 for historical wood, which means that for higher treatment times, the lignin-xylan matrix might 423 424 undergoes some chemical changes that reduces the hygroscopicity of the wood cell wall. As for the modern wood, inverse phenomena was observed with a decrease in contribution of component B in 425 the loss of volume deformation, while component C seems to increase. Nevertheless, the contribution 426 427 of compartment C in modern wood presents higher percentage meaning that it has a higher influence on the decrease in volume deformation. 428

To conclude, the use of NMR by quantifying the two components B and C and relating them to the possible modification of hygroscopicity due to aging, appears to be an available and accurate method to evaluate the decrease of hydric strains. Furthermore, these results confirm that the two components B and C have an effect on hydric deformation, especially for thermal treatments.

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434 Conclusions

This study shows the potential of the 2D ¹H NMR correlation spectra to study modifications in wood
hygroscopicity by being the sole technique able to quantify two types of adsorbed water in two

different chemical environments in the wood cell walls. This method also permits to determine themoisture content of wood without the necessity to oven-dry the wood material.

Two aging methods have been conducted on two oak wood materials (modern and historical wood), 439 on the one hand "natural" aging with repeated hydric cycles and on the other hand "artificial" aging 440 with mild thermal treatment. Results show that the duration of the "natural" aging seems to be not 441 sufficient to induce evolution on the hygroscopic properties, but unexpected results over time were 442 443 observed for the hydric strains. More investigations are necessary, especially with other durations. Concerning "artificial" aging, a clear reduction in total moisture content was observed after heat 444 treatment and reconditioning at 65% RH for the two types of oak wood, accompanied with low dry 445 446 mass loss. The observed decrease of water content is in accordance with the decrease of the swelling 447 strains between 2% RH and 65% RH. Moreover, T_1 - T_2 NMR experiments allow us to measure the effect of heat treatment for the two types of adsorbed water by showing a higher percentage of water 448 449 loss in one compartment than in the other for the historical oak wood. The reason of this selective loss of water adsorption can be explained in relation with the chemical composition of the studied 450 materials. The global decrease of EMC is the result of the loss of extractives but also of structural 451 modifications such as restricted space between polymers leading to a decrease of adsorbed water 452 layers. This is shown by the increase of the T_1/T_2 ratio which indicates a decrease on the averaged 453 454 mobility of H atoms for each water pool after TT and stronger interactions (bonds) between wood and adsorbed water molecules. All these results could be completed with ¹³C NMR or IR investigation 455 to determine the changes that occurred in wood polymers more precisely. 456

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TABLES

Table 1: Density $[g/cm^3]$ and 2D ¹H NMR results (EMC_{NMR,65%} noted B+C [%] and the ratio T₁/T₂ [-])

for the specimens at 65% RH (samples without hydric or thermal loadings). Average values (mean) and

standard deviation (SD) are given.

	Modern wood	Historical wood
	Mean (SD)	Mean (SD)
Density g/cm ³	0.67 (0.02)	0.50 (0.02)
B+C [%]	12.10 (0.36)	11.51 (0.29)
B [%]	9.24 (0.34)	7.36 (0.39)
T1/T2	88.39 (3.46)	76.06 (7.39)
C [%]	2.86 (0.20)	4.15 (0.58)
T1/T2	3.34 (0.12)	3.83 (0.67)

Table 2: Density $[g/cm^3]$ and 2D ¹H NMR results (EMC_{NMR,65%} noted B+C [%] and the ratio T_1/T_2 [-]) of the specimens subjected to repeated hydric cycles. Average values (mean) and standard deviation

(SD) are given.

	<u>.</u>		Modern wood Mean (SD)				Historical woo Mean (SD)	d
	Control samples	1 month	3 months	6 months	Control samples	1 month	3 months	6 months
Density g/cm ³	0.65 (0.04)	0.66 (0.02)	0.65 (0.02)	0.64 (0.02)	0.49 (0.02)	0.48 (0.01)	0.47 (0.01)	0.47 (0.01)
B+C [%]	12.08 (0.11)	12.46 (0.04)	12.45 (0.07)	12.47 (0.02)	11.59 (0.35)	11.85 (0.23)	11.90 (0.35)	12.23 (0.36)
B [%]	9.16 (0.13)	9.20 (0.18)	9.28 (0.09)	9.36 (0.12)	7.46 (0.43)	7.83 (0.50)	7.89 (0.79)	7.12 (1.11)
T1/T2	91.16 (0.00)	82.90 (5.43)	87.84 (5.75)	84.52 (5.75)	84.61 (7.37)	81.20 (0.00)	81.29 (4.70)	75.20 (2.49)
C [%]	2.92 (0.23)	3.27 (0.15)	3.17 (0.08)	3.11 (0.10)	4.13 (0.77)	4.02 (0.35)	4.01 (0.65)	5.11 (0.99)
T1/T2	3.37 (0.20)	3.80 (0.39)	4.17 (0.28)	3.94 (0.27)	4.24 (0.97)	4.10 (0.37)	4.19 (0.52)	4.03 (0.47)

578	Table 3: Density $[g/cm^3]$ and 2D ¹ H NMR results (EMC _{NMR,65%} noted B+C [%] and the ratio T ₁ /T ₂
579	[-]) for the specimens subjected to moderate heat treatment. Average values (mean) and standard

deviation (SD) are given.

		Modern wood Mean (SD)			Historical wood Mean (SD)	l
Time of treatment	24h	72h	168h	24h	72h	168h
Density [g/cm ³]	0.68 (0.01)	0.65	0.65	0.50 (0.02)	0.48	0.46
B+C [%]	10.90 (0.21)	10.04	10.20	10.34 (0.32)	9.02	8.97
B [%]	8.34 (0.20)	7.69	8.06	6.82 (0.46)	6.34	6.30
T1/T2 [-]	111.73 (5.27)	128.99	128.99	100.38 (9.85)	114.90	128.99
C [%]	2.57 (0.16)	2.35	2.14	3.52 (0.61)	2.68	2.67
T1/T2 [-]	4.05 (0.18)	4.25	3.78	4.41 (0.29)	3.57	4.01

- **Table 4:** Mass of specimens [g] before and after heat treatments and the relative mass loss [%]. *this value may be an outlier.

Thermal	Modern wood Mean (SD)			Historical wood Mean (SD)		
Treatment time	Before	After	Relative %	Before	After	Relative %
une	treatment [g]	treatment [g]	mass loss	treatment [g]	treatment [g]	mass loss
24h	0.5731 (0.06)	0.5715 (0.06)	0.27 (0.19)	0.4313 (0.06)	0.4292 (0.05)	0.46 (0.54)
72h	0.5781	0.5746	0.61	0.4303	0.4265	0.87
168h	0.4634	0.4623	0.23*	0.3874	0.3783	2.36

587	Table 5: Contribution of components B and C [%] on the variation of hydric deformation at 65%HR
588	before and after aging. This contribution is given by $\Delta EMC(B \text{ or } C)$ divided by $\Delta EMC(B + C)$.

	Modern	n wood	Historica	l wood
Thermal treatment time	B [%]	C [%]	B [%]	C [%]
24h	78.6	21.4	39.3	60.7
72h	70.8	29.2	41.3	58.7
168h	55.2	44.8	51.4	48.5

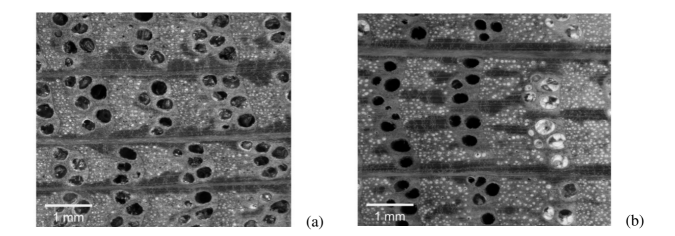


Figure 1: Optical microscopy image of transversal cross-section of the historical oak wood (a) and
the modern oak wood (b). Fibers were possibly filled with wood powder during the polishing (in
white in the fibers).

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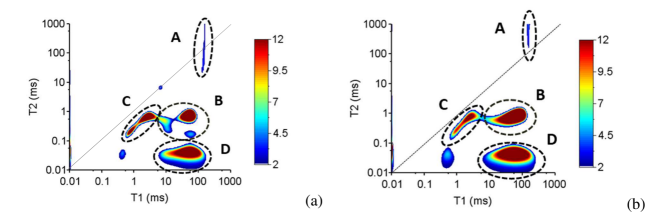


Figure 2: T_1 - T_2 correlation spectra of the historical oak wood (a) and the modern oak wood (b) at the initial state before aging at 65% RH, 20°C. Diagonal lines correspond to T_1 = T_2 .

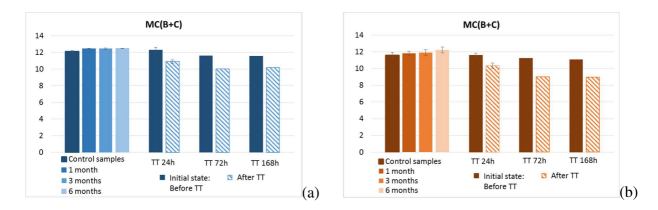


Figure 3: Evolution of the total moisture content $EMC_{NMR,65\%RH}(B+C)$ noted MC(B+C) [%] for (a) modern and (b) historical wood samples subjected to repeated hydric cycles and moderate thermal treatment.

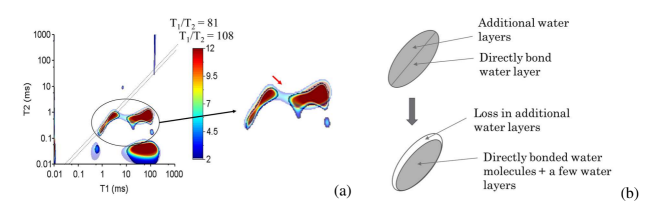


Figure 4: (a) Superposition of two T_1 - T_2 correlation spectra of the same modern wood sample before (in transparency) and after (in bright colour) heat treatment. (b) Schematic representation of the evolution of "peak C" following a thermal treatment.

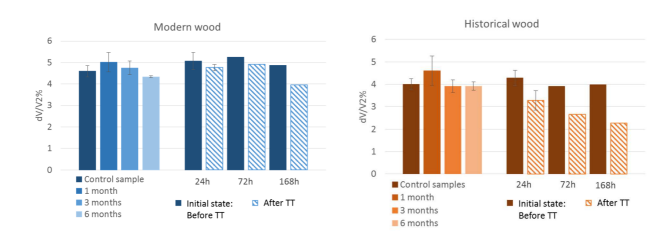


Figure 5: Volume deformations of samples (according to Eq. 5) without aging and through the 6
months repeated hydric cycles and for the thermal treatments.

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