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1 Effect of extractions on dynamic mechanical properties of

2 white mulberry (Morus alba L.)

3 **Original article**

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 mulberry wood (*Morus alba* L.).
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20 Abstract:

21 Vibrational properties of wood are affected by several parameters, of which extractives can be 22 one of the most important ones. Wood for European musical instruments has been often 23 studied, but traditional Middle Eastern ones had been left unnoticed. In this study white mulberry (Morus alba L.), the main material for long-necked lutes in Iran, was extracted by 24 25 five solvents of various polarities (water included). Free-free bar forced vibrations were used to measure longitudinal (L) loss tangent ($\tan \delta$), storage (elastic) modulus (E') and specific 26 modulus (E'/γ) in the acoustic range. Their anisotropy between the 3 axes of orthotropy was 27 28 determined by DMA (dynamic mechanical analysis). Native wood had a quite low $E_{\rm L}/\gamma$ but 29 its tan δ was smaller than expected, and the anisotropy of tan δ and E'/γ was very low. Removal 30 of extractives caused $\tan \delta$ to increase and moduli to decrease. Acetone, the most effective 31 solvent on damping despite a moderate extraction yield, increased $\tan \delta_L$ by at least 20% but 32 did not modify E'/γ as much. When used successively, its effects masked those of solvents 33 used afterwards. Anisotropy of E'/γ was nearly unchanged after extraction in methanol or hot 34 water, while $tan\delta$ was much more increased in R than in T direction. Results suggest that in 35 white mulberry, damping is governed more by nature and localization of extractives rather 36 than by their crud abundance.

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42 Introduction:

43 Wood has always been considered as a multifunctional material. Each of its versatile 44 properties makes it appropriate for a particular final use. Wood historical use in musical 45 instruments is in relation with its dynamic mechanical, or vibrational, properties, including 46 mechanical damping (loss tangent) $\tan \delta$, storage (elastic) modulus E' and specific modulus 47 E'/γ . As a quasi-orthotropic material at the macroscopic scale, wood has different properties in longitudinal (L), radial (R) and tangential (T) directions (Backman and Lindberg¹; Nairn²; 48 49 Brémaud et al³). In the case of wood used for Western classical string instruments, it is 50 generally accepted that a low tan δ and high E'/γ in axial direction, and a high anisotropy, are 51 convenient for obtaining a soundboard of good quality. However, few studies have addressed 52 the vibrational properties of wood used in Middle-East instruments. String instruments in Iran can be traced back to 970 B.C. Long-necked lutes Tar, Setar and Kamancheh are specifically 53 54 made from white mulberry (Morus alba L.), which is also used for the Japanese short-necked lute *Biwa* (Yoshikawa⁴). 55

56 Sources of variations in dynamic mechanical properties of wood include cell arrangement and grain angle, microfibril angle within the cell wall, proportions of constitutive polymers, as 57 well as the presence of additional compounds (Ono and Norimoto^{5,6}; Obataya et al⁷; Noda et 58 al⁸; Bucur⁹). Although in lower quantities than cell wall constitutive polymers, extractives can 59 60 have a considerable effect on mechanical and damping properties of wood. Exceptionally low 61 damping of some musically important woods was somehow correlated with their extractives. Extractives impact can be studied either through correlations between their amounts and 62 63 vibrational properties variation resulting by their removal, or by the re-injection of extracts 64 into a "neutral" wood. Methanol extraction increased the tan δ of heartwood by 15 to 37% in 65 red cedar (*Thuja plicata*) and up to 60% in African Padauk (*Pterocarpus soyauxii*), while E'/γ

slightly decreased (Yano¹⁰; Brémaud et al.¹¹). Impregnation of water-soluble extractives of 66 pernambuco (*Caesalpinia echinata*) into spruce decreased tan δ by a half (Matsunaga et al.¹²). 67 The formation of hydrogen bonds between extractives and cell wall matrix was proposed as a 68 possible cause of tan δ reduction (Matsunaga et al.¹³). Impregnation of Sitka spruce with 69 70 isolated key chemical compounds from muirapiranga (Brosimum rubescens), a Moraceae as is mulberry, decreased tan δ by up to 25%, both in axial and radial directions. This was explained 71 by motion restriction of matrix macromolecules due to extractives entering (Minato et al.¹⁴). 72 73 Contrary to these studies, water-soluble extractives of Reed (Arundo dunax L.) increased both $\tan\delta$ and E' (Obataya and Norimoto¹⁵; Obataya et al.¹⁶). Extractives can also change wood 74 equilibrium moisture content (EMC) (Hernandez¹⁷). Higher EMC induces higher tan δ and 75 lower E'/γ (Akitsu et al.¹⁸; Obataya et al.¹⁹). The potential effect of extractives on vibrational 76 properties could come from their chemical structure, cellular location and relation with basic 77 polymers (Matsunaga et al.¹³; Brémaud et al.¹¹). Removing extractives using solvents of 78 79 different polarities could give a first idea about their polarity and possible cellular location.

Our work aims at evaluating the dynamic mechanical properties of white mulberry and 80 81 understanding the possible contribution of extractives. Basic properties of samples from different regions of Iran were compared in a previous study (Se Golpavegani²⁰; Pourtahmasi 82 and Se Golpavegani²¹). In the present article, mulberry wood was extracted in five solvents of 83 84 different polarities. In order to separate the overlapping effect of various solvents, they were 85 used both as independent extracting agents, and in a successive order on the same specimens, 86 to determine changes in acoustical properties along the grain. Furthermore, the effects on the anisotropy of tan δ , E' and E'/ γ were also studied after extraction by the two solvents with 87 highest vields. 88

90 Material and Methods

91 Material

Mulberry trees were cut in Iran and kept for one month in outdoor conditions. Wood was selected there by a professional instrument maker. Several $500 \times 50 \times 50$ mm (L×R×T) rods were cut from the middle of the heartwood and sent to France, in three batches within one year. The 1st batch (used for powder samples and independent extractions on vibrational specimens) and 2nd batch (successive extractions on vibrational specimens) came from the same tree; the 3rd (specimens for DMA) came from another tree.

A portion of each rod was ground, the fraction passing through 40 mesh and retained on 60 mesh sieves was kept for powder extractions (TAPPI standard method T207_cm²²). For vibrational tests, the rods were cut into approx. 200 axial specimens of dimensions $150 \times 15 \times 2 \text{ mm}^3$ (L×R×T), from which 77 specimens were kept for extractions and controls (Table 1). For DMA (Dynamic Mechanical Analysis), specimens were cut to a cross section of 4×3 mm² and a length of 42-48 mm (Fig. 1). 14 specimens were prepared in each direction of orthotropy leading to a total number of 42.

105 Conditioning and physical measurements

106 The specimens were oven dried at 60°C for 48 h. This procedure ensured that the equilibrium 107 was reached in adsorption. Residual moisture content, as compared with oven drying at 108 103°C, was around $1.36\pm0.49\%$. Specimens were then kept for three weeks in controlled 109 conditions of $20\pm2°$ C and $65\pm5\%$ RH. This procedure was repeated before any mechanical 110 measurement. Specific gravity γ (both oven dried and air dried) and EMC were recorded. 111 EMC of native wood was of $7.9\pm0.8\%$. Although mechanical measurements could not be run in a climatic chamber, mass change was small: $-0.3\pm0.4\%$ for native state specimens, both during vibrational and DMA tests. For treated specimens, mass change was negligible ($-0.1\pm0.1\%$) during vibrational tests, and less than during DMA tests ($-0.6\pm0.7\%$).

116 Measurement of dynamic mechanical properties

Axial dynamic mechanical properties were measured both by vibrational tests and by DMA,while anisotropy was studied only by DMA.

119 Non-contact forced bending vibrations of free-free bars

120 A frequency scan was imposed through an electric magnet facing a tiny steel plate (15-20 mg, 121 a negligible additional weight) glued at one end of each specimen. A laser triangulation sensor measured the displacement. A program developed with LabView® software 122 (Brémaud²³) monitored vibration emission and detection. Specific elastic (storage) modulus 123 $(E_{\rm L}'/\gamma)$ was calculated from the first resonance frequency by the Euler-Bernoulli formula and 124 125 elastic modulus $(E_{\rm L}')$ was obtained by multiplying $E_{\rm L}'/\gamma$ by specific gravity (γ). Damping 126 coefficient was measured using both bandwidth at half power in the frequency domain (or 127 quality factor) and logarithmic decrement of amplitude in the time domain after stopping the vibration. Both measurements shall be equivalent to loss tangent $\tan \delta$, if $\tan \delta \ll 0.1$, which is 128 129 the case for air-dry wood in this temperature/frequency range. Three repetitions were made 130 for each specimen. Resonance frequencies were in the range of 200-400 Hz.

131 DMA (Dynamic mechanical analyzer)

132 The viscoelastic behavior of specimens was measured using a BOSE® ELF3230 DMA 133 equipped with tension/compression fatigue grips, a 22 N ($\pm 0.17\%$ maximum error) load cell 134 and a high-resolution displacement sensor (1 mm range with $\pm 0.26\%$ maximum error). The 135 specimens were fixed between two clamps with a working distance of 35 mm and tested in 136 tension along their longest direction. The tests were done in purely alternative 137 tension/compression loading controlled in displacement with an amplitude of ± 0.0175 mm 138 corresponding to a maximum tension/compression strain of $\pm 0.02\%$ (i.e., within the linear 139 viscoelastic region, see Sun et al.²⁴). Using BOSE WinTest® analysis software, E^* (norm of 140 the complex modulus), E' (storage modulus), E'' (loss modulus) and tan δ (loss tangent) were 141 calculated. Both moduli and $tan\delta$ were corrected for the stiffness of the whole apparatus 142 (around 1785 N/mm with a negligible viscous contribution), measured using a stiff steel 143 specimen. Frequency sweeps from 0.1 to 10 Hz were run in triplicate for each specimen, 144 native and then treated.

145 Extraction procedures

Extractions used solvents of increasing polarity: hexane (HX), dichloromethane (DM), acetone (AC), methanol (ME) and hot water (HW). Most used Soxhlet extractors. Groups of specimens, with similar ranges in properties, were defined after vibrational tests in native state and subjected to different treatments (Table 1).

For extractions on powder and on solid specimens for vibrational tests, extractions were run both in an independent (or parallel) and in a successive (or serial) way. Independent extraction means that a sample is submitted to one extraction in a given solvent, and its properties are measured before and after this single treatment. In this case, hot water extraction from vibrational specimens was run (for 8h) at 70°C (not using Soxhlet). Successive extractions means that a given group of specimens is submitted to extraction, first by the less polar solvent (HX), then oven-dried (brief process: weighted, air-dry stabilized, weighted, mechanically tested, and dried again for solid wood specimens), then extracted by the next solvent (DM), and so on. In this case, a "standard" Soxhlet extraction (<95°C) was run for water (complete description of conditions can be found in table 1).

160 Specimens were dried before being extracted, so that less polar solvents could not enter cell 161 walls. Control specimens underwent only physical steps (drying and stabilization) and were 162 measured in the same time and condition as extracted ones. For determination of extractive 163 content, 3 g of powder were put in a cellulose cartridge, extracted for 8 h, and their oven-dry 164 (48 h at 60°C) weight loss was measured. For solid wood, as color hardly changed during extraction, duration of 12 h was chosen. Based on the yields of extractions in vibrational 165 166 specimens, ME and HW were applied as independent solvents on DMA specimens. For each 167 anisotropic direction, 4 specimens served as controls, 5 were extracted in ME and 5 in HW 168 (70°C, without Soxhlet). Although hemicelluloses may be partially depolymerized at moderately high temperatures for water-saturated wood (Placet et al^{25} ; Assor et al^{26}), 169 170 treatment temperatures and duration were sufficiently low to neglect such effect.

171 **Results and Discussion**

172 Properties of native mulberry wood

The specific gravity γ of the mulberry wood under study ranged from 0.45 to 0.61 (Fig. 2). This was significantly lighter than wood from different regions in a previous study (Se Golpayegani²⁰). The 1st batch (used for independent extractions) was significantly denser than the 2nd one (used for successive extractions), although they came from the same tree. The 3rd batch (used for DMA), which came from a different tree, covered a broad range, but its 178 average γ was not significantly different from the 2nd batch. Differences in density were not 179 clearly related to those in vibrational properties (Fig. 3).

180 The specific dynamic modulus in L direction of all studied batches of wood was rather low 181 (Fig. 3) and variable (11-18 GPa). Although dispersion was quite large, $\tan \delta_{\rm I}$ was negatively related to $E'_{\rm L}/\gamma$. However, all batches generally had a lower tan $\delta_{\rm L}$ than the "standard trend" 182 from Ono and Norimoto^{5, 6}, a statistical relationship obtained on 20 softwoods and 30 183 hardwoods that can be considered as a reference. In the 1st and 3rd batches tan δ_L was in 184 average 6% and 11% lower than "standard", but with a higher dispersion than in 2nd batch and 185 in wood from a previous study (Se Golpayegani²⁰), for which tan δ was always within the 186 187 lower range (21% and 23% lower than the standard).

188 $tan \delta_L$ measured with DMA method was higher than that measured with free-free vibration. 189 However, it exhibited a decreasing trend against frequency, so that the difference could be 190 attributed to the higher frequency of the free-free vibration. This decrease of $tan\delta$ with 191 frequency was also observed in other directions, as well as a slight increase of elastic moduli. 192 However, the anisotropic ratios remained constant in the observed frequency range. In the 193 following analysis, only the values measured at 10 Hz will be used. Ordering of the different $\tan\delta$ in the 3 principal directions was the same as previously reported, i.e., $\tan\delta_{\rm T} > \tan\delta_{\rm R} >$ 194 $tan\delta_L$ (Ono and Norimoto²⁷). However, damping anisotropies between the three main 195 directions were small: $R/L \approx 1.03$, $T/L \approx 1.34$, and $T/R \approx 1.30$. These values were lower than 196 those collected from several studies by Brémaud et al.³ in which the average ratios were of 2.7 197 198 (R/L), 2.9 (T/L) and 1.14 (T/R) for hardwoods. Similarly, the anisotropy of E' agrees with well-known relationships, i.e., $E'_L >> E'_R \ge E'_T$, but actual values of ratios (L/R \approx 4, L/T \approx 8 199 and R/T \approx 1.9) were in the lower range of anisotropy compared with literature reviews (e.g., 200 Guitard and El Amri²⁸, Nairn²). It can be noted that mulberry has a low longitudinal E'/γ and a 201

very reduced anisotropy when compared with resonance spruce that is used for top plates ofWestern string instruments, and is closer to maple (used for back and sides) in this respect.

204 Yield of extraction

In wood powder (Fig. 4a), approximately the same cumulated amount of extractives was removed when applying solvents independently or successively. In solid wood (Fig. 4b), on the contrary, cumulated weight losses differed between these two extraction procedures, suggesting some structural effect on the accessibility to various solvents. This was supported by different extraction yields from axial, radial and tangential DMA specimens: 9, 14 and 18% respectively for ME.

211 Less polar solvent HX and DM, which should reach only the lumen, removed small and 212 comparable amounts (1.6% and 1.7%) when used independently on powder. Their yields were 213 much smaller in solid wood. When DM was used after HX, its yield was very low, suggesting 214 that HX had already removed most of extractives accessible to apolar solvents. Similarly, AC 215 had a smaller yield when used after HX and DM, suggesting that when used independently, it 216 also removed compounds from lumens. Although AC and ME are thought to be able to 217 solubilize similar types of compounds, ME caused the most exhaustive extraction from solid 218 wood: 6.2% when used independently. Its efficiency was even increased when used after AC 219 (7.14% from solid wood). On the contrary, in powder, HW had the highest yield (8.31%) 220 when used independently, while it had a much smaller yield on solid. This suggests that in 221 solid wood, polar extractives are more easily removed by ME than by HW, as the final 222 cumulated yields were however similar between powder and solid.

223 Extraction effects on E_L'/γ

As extractions change the mass and specific gravity (Table 2, Table 3) of wood specimens, $E_{\rm L}'/\gamma$ values should be corrected for the contribution of extractives to γ :

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$$\left(\frac{E_{L}^{'}}{\gamma}\right)_{c} = \left(\frac{E_{L}^{'}}{\gamma}\right)_{ap} \times \frac{\gamma_{ap}}{\gamma_{ef}}$$
(1)

where $(E_L'/\gamma)_{ap}$ and γ_{ap} are the apparent measured values, and $(E_L'/\gamma)_c$ is the corrected value taking into account the specific gravity γ_{ef} after the most effective extraction (ME). $(E_L'/\gamma)_c$ should be more representative of an "extractives-free cell wall" modulus, and appeared 0.01 to 1.2 GPa higher than the apparent one (Table 2, Table 3).

Corrected E_L'/γ gradually decreased due to extractions. HX, DM, AC, ME and HW modified E_L'/γ by 0.2%, -0.6%, -5%, -11% and -1.8% respectively after independent extractions (Fig. 5a). Cumulated changes (i.e. relative to native state) after successive extractions followed a similar trend but with a bigger amplitude (Fig. 5b). E_L' shows a similar gradual decrease. Decrease in E_L'/γ appears nearly proportional to weight loss for different solvents (Fig. 5b), which suggests that extractives removed by AC, ME and HW bulked the cell wall matrix, and affected elasticity independently of compounds removed by these various solvents.

238 Changes in $tan\delta_L$ related to weight loss

Figure 6 shows relative changes in $\tan \delta_L$ caused by each solvent either used individually or after others (successively). Apolar solvents (HX and DM) removed small amounts (approx. 1%) of extractives, probably from the lumens, resulting in small changes in $\tan \delta_L$. In successive extraction, however, DM increased $\tan \delta_L$ by 14%, which might be linked to previous exposition to HX. On the other hand, in another species (Padauk), DM was able to slightly attain cell wall and increase $\tan \delta_{\rm L}$ of approx. 20% for approx. 3% extracts removed (Brémaud et al.¹¹).

246 More polar solvents (AC, ME and HW) are expected to remove higher proportions of cell wall extractives, more susceptible to alter $tan \delta_L$. However, the highest change in $tan \delta_L$ 247 resulted from AC extraction (\approx +20% independently or used after DM, \approx +38% for cumulated 248 249 successive effect), although its weight loss was moderate. ME and HW, despite their higher 250 yields, had smaller effects on $\tan \delta_L$ both in independent and in successive-cumulated order. 251 When used after AC, ME even reduced tan $\delta_{\rm L}$ lower than the value for the previous state (AC-252 extracted, Fig. 6b). On the contrary, in other species, methanol extractives often have a high "anti-damping" effect in wood (Yano¹⁰; Minato et al.¹⁴; Brémaud et al.¹¹). Acetone has also 253 been used recently in this kind of study, on two tropical species (Brémaud²³; Brémaud et 254 al.¹¹). It was more efficient than ME in one species, and less in the other. The contrasted 255 256 effects observed in mulberry could be explained by different hypotheses: (i) AC reached all 257 extractives able to alter damping, leaving only "inactive" compounds for subsequent ME or HW extractions. As drying cycles could reduce $\tan \delta_L$ by $\leq 8\%$, changes in $\tan \delta_L$ after ME and 258 259 HW in successive order could partly reflect drying history of specimens. However, a smaller 260 effect on tan $\delta_{\rm L}$ of ME and HW than that of AC was also observed for independent extractions, 261 without cumulated drying cycles. This brings us to the second hypothesis; (ii) Two kinds of 262 extractives compounds would co-exist in cell walls, some decreasing, and some increasing, 263 $\tan \delta_{\rm L}$ in native wood. AC would have removed the first ones, so that extraction of remaining, "tan $\delta_{\rm L}$ raising", compounds by ME and HW would decrease again the tan $\delta_{\rm L}$ of solid wood. 264

In various woods, methanol is reported to take out non-structural carbohydrates and phenolics (e.g. Rowe and Conner²⁹). White mulberry has quite important content of phenolic extractives (De Rosso et al.³⁰). In some woods, polyphenols can form part of the fiber cell wall matrix

(Kleist and Bauch³¹), while in some other woods, they may be either preferentially in vessels, 268 parenchyma walls (Dünisch et al.³²), or too much condensed to enter the cell wall substance 269 (Koch and Kleist³³). In the later hypothesis, ME might have removed some lumen deposits, 270 resulting in a lower ratio between changes in $\tan \delta_{\rm L}$ and extraction yield. However, decreases 271 in $E'_{\rm L}/\gamma$ after ME or HW suggested cell wall location. Thus, ME could have removed 272 compounds such as sugars or simple phenols, which presence will increase $\tan \delta_{\rm L}$ (Obayata et 273 al.¹⁵; Sakai et al.³⁴). A decrease in tan δ_L is also observed after ME extraction of *Thuja plicata* 274 sapwood – which should contain non-structural carbohydrates – whereas a completely 275 opposite effect is observed in the heartwood of the same species (Yano¹⁰). In mulberry, 276 extraction in HW, following ME, did not bring additional changes in tan δ_{L} , although it further 277 278 removed $\approx 4\%$ extractives. In *Caesalpinia echinata*, water soluble extractives decrease tan $\delta_{\rm L}$ 279 in wood, which was first ascribed to their ability to form hydrogen bonds with cell walls components (Matsunaga et al.¹²). But other extractives, without hydroxyl groups and/or 280 insoluble in water, also reduce $\tan \delta_{L}$ (Minato et al.¹⁴; Brémaud et al.¹¹). Finally, differences in 281 $\tan \delta_1$ might be related to changes in wood moisture content (Dunlop³⁵; Obavata et al.¹⁹; 282 Inokuchi et al.³⁶). However, changes in EMC due to independent extractions were small 283 284 (Table 2). In successive extractions (Table 3), with higher cumulated yields, EMC was more 285 significantly increased after ME and HW. This, however, should lead to an increase in tan $\delta_{\rm L}$, 286 instead of the observed decrease for successive use of ME and HW.

Therefore, the present results suggest the co-existence of some "tan δ – lowering" and of some "tan δ – raising" compounds in the heartwood of white mulberry. The first ones are extractable by ME but also with AC and HW, consequently independent extraction with those solvents had always resulted in an increase in tan δ_L . The latter compounds, being probably hydrophilic, are not extractable by HX, DM or AC. Thus a successive extraction using ME and HW had reduced $\tan \delta_{\rm L}$, as the " $\tan \delta$ – lowering" had been already removed by previous solvents and there were only the " $\tan \delta$ – raising" components left. The methodology using both independent and successive series of extraction might reveal similar trends in other species.

296 Changes in anisotropic properties due to extractions

Figure 7 shows the variations in E'/γ and $\tan \delta$ in the three principal directions of orthotropy after extractions with HW and ME or 2 drying cycles (controls). Variations in E'/γ after ME extraction were similar in all three directions: -18% to -20%, although weight losses were different (9%, 14% and 18% for L, R and T specimens respectively). Hot water caused both lower weight losses (6-8% in L, R and T), and smaller decrease in E'/γ along L and R directions, while it was similar to ME in T direction.

303 Concerning tan δ , uncertainty was high in L direction, as clamping of specimens may crush 304 wood in the softer transverse direction. Even controls showed important variations, which 305 cannot be ascribed to physico-chemical changes. However, after ME extraction, change in 306 radial tan δ was at least twice more important than in L (DMA estimations and changes in 307 vibrational tests) and T direction (Fig. 7).

The quasi-isotropic effect of extractions on E'/γ is quite surprising. In previous works, extractives stiffened the transverse moduli (Yano et al.³⁷; Minato et al.¹⁴), but not the axial one, in which the influence of the microfibrillar reinforcement predominates (Ono and Norimoto ^{5, 6}; Obataya et al.⁷). However, mulberry wood has a low longitudinal E'/γ , suggesting high microfibril angle, and is characterized by a very low anisotropy in native state. Effects of extractions on tan δ were clearly different in radial and tangential directions. This is interesting, as there are relatively few works on the tan $\delta_{\rm R}/tan\delta_{\rm T}$ anisotropy and sources of variations. Backman and Lindberg¹ stated that, for softwood, in tensile test latewood contributes mostly to the response in tangential direction while most of the strain occurs in early wood in radial direction. In mulberry, a ring-porous hardwood, the response in radial tension tests would more involve vessels and rays, whose cell walls are thinner, but are also susceptible to contain higher proportions of extractives (Koch³⁸; Kleist and Bauch³¹; Dünisch et al.³²). Thus, the very high change in tan δ_R after ME extraction (\approx 50%) could express a bigger modification of rays and vessels, than of fibers.

322 Conclusion

323 Dynamic mechanical properties E'/γ , tan δ , and their anisotropic ratios in the 3 principal axes 324 were measured on white mulberry wood (*Morus alba* L.), a representative raw material for 325 musical instruments in Iran. They were compared before and after extractions by different 326 solvents in order to investigate extractives effects. Results could be synthesized as follows:

- 327 Mulberry wood has a quite low $E'_{\rm L}/\gamma$ but its damping factor $(\tan \delta_{\rm L})$ is lower than 328 expected. The anisotropy of these two properties is also much lower than average.
- Combining two methods of extractions -using each solvent as an individual agent and
 using solvents in a successive order on a single sampling- could suggest the
 importance of extractives nature compared to their amount.
- The highest changes in $\tan \delta_{\rm L}$ were not due to the most polar solvents, with highest extraction yields, but to acetone. Nevertheless, acetone did not alter $E'_{\rm L}/\gamma$ with the same intensity, suggesting that the extracted amounts or compounds were not essential for wood stiffness.

The different effects of methanol extraction on $\tan \delta_L$, between individual extraction and when used successively after acetone, suggest the existence of two types of $\tan \delta$ altering compounds in white mulberry. Some would be able to increase, some other would be able to reduce the damping, and both types would be accessible by acetone and methanol independently.

341 - Changes in E'/γ were nearly isotropic between the 3 principal axes of orthotropy, may 342 be due to the very low anisotropy of mulberry in native state. On the contrary, $\tan \delta$ 343 was much more modified in radial than in tangential direction, suggesting different 344 responses to extractions of rays and vessels walls as compared to fibers ones.

345 It should be noted that even though extractives are found to definitely affect vibrational 346 properties of *Morus alba* L., it would be enlightening to identify the most important 347 compounds present in those extractives and how they cause changes in vibrational properties.

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458 **Tables**

Number of specimens	Solvent	Drying and stabilization condition	Number of cycles of drying and stabilization
12	Hexane (HX)		2
12	Dichloromethane (DM)		2
12	Acetone (AC)		2
12	Methanol (ME)		2
5	Hot Water (HW, 70°C)	60°C for 48 h	2
12	HX/DM/AC/ME/HW (100°C)	followed by	6
12	Controls	20±2°C & 65±5% RH for 3weeks	6
15 DMA(5L, 5R, 5T)	5 DMA(5L, 5R, 5T) ME		2
15 DMA(5L, 5R, 5T)	HW (70°C)		2
12 DMA(4L, 4R, 4T)	Controls		2

459 Table 1: Specimens number and treatments for both independent and successive methods of extraction

460

- 474 Table 2: Basic statistics for differences in properties between groups submitted to independent extractions and to
- 475 one cycle of drying/re-stabilization (=control).

Treatment		$\left[C\left(\%\right)^{d}\right] $	speen	c gravity γ		rected (GPa)	i i i i i i i i i i i i i i i i i i i	_L (10 ⁻³)	
	Corrected				$E'_{\rm L}$ / γ Apparent (GPa)				
	Ар	parent							
	Native	Treated	Native	Treated	Native	Treated	Native	Treated ^f	
Control ^e	9.2±0.5	9.2±0.2 ns	0.556 ± 0.027	0.553±0.030 ns	15.0±2.2	15.2±2.4 ns	8.6±0.3	7.9±0.5 **	
	8.2±0.4	8.2±0.2	(a)	(ab)	14.0 ± 1.6	14.2 ± 1.6	(a)	(a)	
	(a)	(ab)			(b)	(a)			
Extracted									
HX	9.1±0.2	8.9±0.4 ns	0.570±0.031	0.566±0.030 **	13.5±2.0	13.7±2.0 **	10.2 ± 0.6	9.6±0.6 **	
	8.1±0.2	7.9±0.4	(a)	(b)	12.3±1.4	12.5±1.4	(b)	(bc)	
	(a)	(a)			(a)	(a)			
DM	9.0±0.3	9.6±0.4 **	0.560 ± 0.031	0.557±0.031 **	14.3±2.2	14.3±2.2 ns	9.6±1.0	9.3±0.8 *	
	8.0±0.3	8.6±0.3	(a)	(b)	13.2±1.6	13.3±1.6	(b)	(bc)	
	(a)	(c)			(ab)	(a)			
AC	9.1±0.4	9.4±0.4 *	0.567 ± 0.026	0.550±0.025 **	14.6±1.6	13.9±1.6 **	8.7±0.4	9.8±0.6 **	
	8.1±0.3	8.6±0.3	(a)	(ab)	13.3±1.1	13.2 ± 1.1	(a)	(c)	
	(a)	(c)			(ab)	(a)			
ME	9.2 ± 0.4	8.7±0.2 **	0.554 ± 0.026	0.520±0.025 **	14.4 ± 2.2	12.9±2.0**	8.9±0.5	9.5±0.5 **	
	8.2±0.4	8.3±0.3	(a)	(a)	13.5±1.4	12.9±1.4	(a)	(bc)	
	(a)	(bc)			(ab)	(a)			
HW	9.2±0.3	9.0±0.2 <i>ns</i>	0.550±0.015	0.530±0.017 **	14.1±1.7	14.0±1.9 ns	8.5±0.5	8.6±0.6 ns	
	8.2±0.3	8.3±0.1	(a)	(ab)	13.3±1.3	13.7±1.7	(a)	(ab)	
	(a)	(bc)	values; "Corr		(ab)	(a)			

477 (for EMC) and specific gravity (for E'/γ).

478 (a, b, c): homogenous subsets in one-way ANOVA at a level α :0.05 (=comparison between groups in a given

479 column, based on measured values).

480 (ns, *, **): differences between untreated and treated properties of a given group (based on corrected values

481 whenever applicable) in *t*-test for paired samples; **: significant at α: 0.01, *: α: 0.05, *ns*: not significant.

d) EMC relative to 103°C oven-drying. "Apparent" are measured values; "Corrected" is calculated by taking into

483 account the total extractive content (cumulated % on powder) from 1st batch of wood (see Hernandez 2007).

484 e) "Treated" state for controls stands for values measured after one cycle of oven-drying followed by air-dry re-

485 stabilization.

486 f) $\tan \delta_{\rm L}$ are raw values, whereas $\tan \delta_{\rm L}$ variations in Fig. 6 are corrected for controls.

487

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488

490	Table 3: Basic statistics for the successive ranges in properties for the group of specimens submitted to
491	successive extractions.

Treatment	EMC (%) ^a Corrected Apparent	Specific gravity γ	$E'_{\rm L}/\gamma$ Corrected (GPa) $E'_{\rm L}/\gamma$ Apparent (GPa)	$\tan\delta_{\rm L} (10^{-3})$		
Untreated	7.0 ± 0.3	0.515 ± 0.011	13.72 ± 1.47	8.2± 0.5		
	6.1 ± 0.3		12.99 ± 1.25			
HX	$7.1 \pm 0.3 \ ns$	0.512 ± 0.016 ns	13.46± 1.68 ns	7.9±0.4 **		
	6.2 ± 0.2		12.82 ± 1.35			
DM	9.2± 0.2 **	0.515 ± 0.012 ns	13.51± 1.26 ns	9.3± 0.6 **		
	8.0 ± 0.2		12.78 ± 1.07			
AC	$9.0 \pm 0.4 \ ns$	0.515 ± 0.016 ns	13.53± 1.64 ns	11.3± 0.8 **		
	8.0 ± 0.3		12.79 ± 1.30			
ME	8.6± 0.1 **	0.505±0.014 **	12.69±1.38 **	10.6± 0.6 **		
	8.2 ± 0.1		12.24±1.13			
HW	8.8± 0.1 **	0.488± 0.011 **	11.85±1.23 **	10.2± 0.5 *		
	88 + 01		11.84 ± 1.05			

 $[\]frac{8.8 \pm 0.1}{11.84 \pm 1.05}$ 492 "Apparent" stand for measured values; "Corrected" take into account the contribution of extractives to weight

493 (for EMC) and specific gravity (for E'/γ).

494 a) EMC relative to 103°C oven-drying. "Apparent" are measured values; "Corrected" is calculated by taking into

495 account the total extractive content (cumulated % on solid specimens) from 2nd batch of wood (see Hernandez

496 2007).

497 (*ns*, *, **): differences between each successive extracted state in relation to the previous phase of ttreatment

498 (based on corrected values whenever applicable) in *t*-test for paired samples; **: significant at α: 0.01, *: α: 0.05,

- *ns*: not significant.

509 Figure legends

Fig. 1. Cutting plan for vibrational and DMA specimens. Variations in length of DMA specimens were due tolimitations in raw material.

512

513 Fig. 2. Box plot of specimens specific gravity (γ) for the three batches of wood under study, and for previous 514 data on wood from different origins (Se Golpayegani²⁰). (a, b, c) significantly different groups (one-way 515 ANOVA).

516

517 Fig. 3. Relationship between $E'_{\rm L}/\gamma$ and $\tan \delta_{\rm L}$ for 4 batches of native mulberry wood (N specimens: see Fig. 2).

518

519 Fig. 4: Comparison of weight loss due to independent and successive extraction in (a) powder, and (b) solid 520 (vibrational) specimens. "Successive" weight losses are relative to the previous step; "independent" and 521 "successive-cumulated" are relative to native oven-dry weight.

522

Fig. 5. Relationship between weight loss (%) and relative change of $E'_{\rm L}/\gamma$ (%) after (a): independent extractions and (b): successive extractions where changes are relative to initial state of native specimens (i.e. cumulated changes). $E'_{\rm L}/\gamma$ was corrected for contribution of extractives to γ .

526

527 Fig. 6. Relative changes (%) in $\tan \delta_{L}$ plotted against weight loss (%) for (a) independent and (b) successive 528 extractions. Changes are relative to initial state of native specimens (independent and successive-cumulated). 529 Arrows: order of successive extractions. Values of $\tan \delta_{L}$ in independent extractions are corrected for controls.

- 531 Fig. 7. Variations in mechanical properties after extractions measured by DMA at 10 Hz. Values for controls
- 532 were not deduced from treated ones.

534 Figures

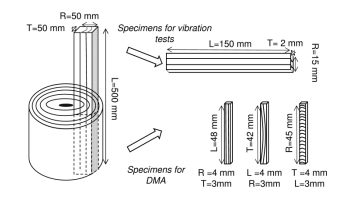






Fig. 1.

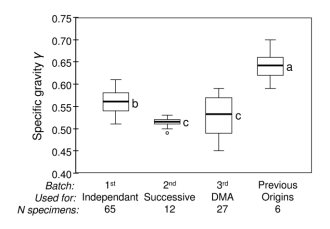




Fig. 2.

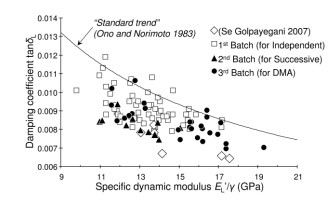


Fig. 3.





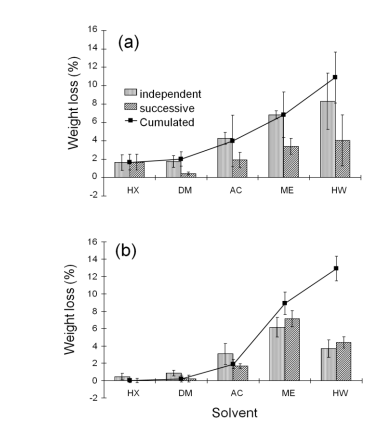


Fig. 4.

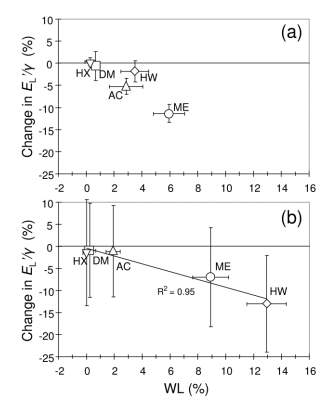


Fig. 5.

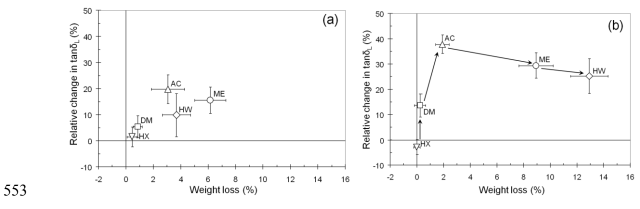


Fig. 6.

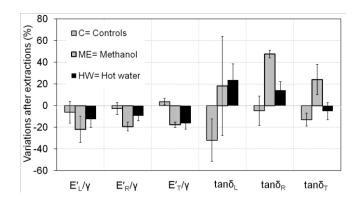


Fig. 7.