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Kévin CANDELIER, Marie-France THÉVENON, Philippe GÉRARDIN, Stéphane DUMARÇAY, Robert COLLET - Anti-fungal and anti-termite activity of extractives compounds from thermally modified ash woods - MADERAS: Ciencia y Tecnología - Vol. 22, n°2, p.32 - 2020

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| | Maderas-Cienc Tecnol 22(2):2020 |
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| | Ahead of Print: Accepted Authors Version |
| 1 | DOI:10.4067/S0718-221X2020005XXXXXX |
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| 3 | ANTI-FUNGAL AND ANTI-TERMITE ACTIVITY OF EXTRACTIVES |
| 4 | COMPOUNDS FROM THERMALLY MODIFIED ASH WOODS |
| 5 | |
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| 15 | Accepted: February 11, 2020 |
| 17 | Posted online: February 12, 2020 |
| 18 | |
| 19 | ABSTRACT |
| 20 | Thermal modification of wood is a promising alternative to chemical and biocidal |
| 21 | modification processes, increasing the biological durability and dimensional stability of wood. |
| 22 | However, the wood-decay resistance properties of heat-treated wood are still not well known. |
| 23 | The main objective of this study was to determine the biological resistance of heat-treated ash |
| 24 | wood, and assess the antifungal and anti-termite activity of extractive compounds from heat- |
| 25 | treated ash woods, depending on the intensity of the modification process (2 hours at 170, |
| 26 | 200, 215, 228 (°C) - steam pressure). Untreated and heat-treated wood samples were extracted |
| 27 | with water or acetone. The extracts were then used to determine inhibition effectiveness |
| 28 | against white-rot (Trametes versicolor) and brown-rot (Rhodonia placenta) fungi. Whatman |
| 29 | papers impregnated with extractives were used to evaluate the inhibition of termite feeding. |
| 30 | Lastly, the extractives were analyzed by Gas Chromatography - Mass Spectrometry (GC-MS) |
| 31 | and compared for their level of anti-termite and antifungal activity. The results showed that |
| 32 | the degree of antifungal activity of these extracts depended on the solvent used during the |
| 33 | extraction process and varied depending on heat treatment intensity. The extracts were more |
| 34 | effective against brown-rot than white-rot fungi. However, the anti-termite activity of heat- |
| 35 | treated ash wood extracts was not really significant. A GC-MS analysis showed that the main |
| 36 | share of the extractives in untreated wood was removed. In addition, new chemical elements |
| 37 | were generated by the thermal degradation of wood polymers (lignin and hemicelluloses), |
| 38 | including aliphatic acids, monosaccharides and other products resulting from their |
| 39 | dehydration reaction. The most abundant element was syringaldehyde, from lignin derived |
| 40 | compounds, which might explain the antifungal activities of thermally treated ash wood |
| 41 | extracts. |
| 42 | Keywords: Antifungal activity, extractive compounds, <i>Fraxinus</i> spp., heat treatment, termite |
| 43 | resistance. |

44 **1. INTRODUCTION**

45

Ash (*Fraxinus* spp.) is a significant hardwood resource largely used in the European wood 46 47 industry. Given its very weak resistance to decay and insects, this timber is naturally in appropriate for class 3 use (EN 350 CEN 2016). Ash wood materials therefore need to be 48 treated with biocide additives for outdoor use (Candelier et al. 2017; Gérard et al. 2017). 49 Industrial wood protection methods usually focus on ways of impregnating active 50 formulations containing biocides. However, ash wood is only a moderately treatable timber 51 for both its sap and heartwood (EN350 CEN 2016; Gérard et al. 2017). In addition, due to 52 53 their impact on human health, these chemicals are a subject of environmental pressures, which will restrict their use in the near future (Schultz et al. 2007). The Implementation of EU 54 Environmental Legislation in recent years has resulted in the development of non-biocidal 55 alternatives, such as thermal modification (Sandberg et al. 2016), chemical impregnation 56 (Gérardin 2016; Mantanis 2017; Guo et al. 2018), or a combination of the two (Salman et al. 57 2016; Sandberg et al. 2017). Using antagonistic microorganisms, combined or not with 58 biochemicals, is also considered as a promising ecofriendly way of protecting wood: (i) 59 60 biocontrol microorganisms consume the available nutrients and produce biochemical compounds that are toxic and/or repellent to fungal decay and, (ii) the biochemical 61 compounds include cell wall-degrading enzymes, siderophores, chelating iron and other 62 antibiotics. This last wood preservation method is not vet industrialized and requires 63 additional studies, especially regarding the environmental impact of these formulations (Susi 64 *et al.* 2011). 65

66 On the other hand, thermal treatment of wood, using slow pyrolysis in an inert atmosphere, 67 has been widely studied and has now been industrially developed around the world, mainly to 68 increase wood durability against fungi in line with the process parameters (Calonego *et al.* 69 2010; Candelier *et al.* 2016). According to the literature (Kamdem *et al.* 2002; Weiland and

Guyonnet 2003), the improved resistance of heat-treated wood material to decay can be
explained by the following four assumptions, all derived from chemical modifications of
wood cell wall polymers due to their thermal degradation (Tjeerdsma and Militz 2005; Yildiz *et al.* 2006; Inari *et al.* 2007):

(i) The hydrophobic behavior of wood is increased, thus limiting water sorption within the
material and reducing any fungal growth. This improved hydrophobicity reduces the capillary
transfer of water into the thermally modified wood and thus limits fungal growth by removing
the optimum conditions for its development (due to lack of water). In addition, heat-treated
wood has a lower Fiber Saturation Point (FSP) than untreated wood, leading to better
resistance to rot fungi (Weiland and Guyonnet 2003; Hakkou *et al.* 2006).

(ii) Wood polymers are modified and the enzymes involved in the fungal degradation of wood 80 do not recognize it anymore as such. Firstly, the changes in lignin chemical composition 81 82 prevent fungal enzymes from recognizing and attacking specific target molecules (Vallet et al. 2001; Lekounougou et al. 2009). Secondly, changes in the ligneous polymer network also 83 seem to play a role in fungal inhibition. The improved resistance of modified wood to decay is 84 mainly due to possible crosslinking between lignin and certain molecules derived from wood 85 86 thermal degradation, such as furfural (Weiland and Guyonnet 2003). Such modifications lead to a substrate that is not recognized by the fungal enzyme system, resulting in an undecayed 87 material (Hakkou et al. 2006). Moreover, cellulose may undergo an esterification reaction, 88 89 due to the acetic acid generated during the thermal degradation of hemicelluloses (Tjeerdsma 90 and Militz 2005; Gao et al. 2016).

91 (iii) Hemicelluloses are the main wood component degraded by thermal modification,
92 inducing a significant loss in potential nutrients for fungal growth (Hakkou *et al.* 2006; Altgen
93 *et al.* 2019). The chemical modification of polysaccharides is responsible for improving

94 resistance to wood decay, but cannot be the only factor, as lignin is also an important source

95 of nutrients for white-rot fungi (Lekounougou *et al.* 2009).

96 (iv) New extractive, fungicidal substances are generated by wood thermal degradation and can serve to prevent decay. In fact, some extracts from heat-treated beech, poplar and maritime 97 98 pine are already known to have antifungal activities (Kamdem et al. 2000; Peters et al. 2009; Lovaglio et al. 2017)). However, other studies showed that improvement in the resistance of 99 beech wood to decay does not appear to be due to new extractible compounds formed during 100 101 thermal treatment. Hakkou et al. (2006) reported that the new extractive compounds produced during heat treatment of beech wood, carried out between 200 °C and 280 °C, only had a 102 103 slight effect on the heat-treated wood against Trametes versicolor. Kamdem et al. (2002) found similar results in a study focusing on heat-treated pine and spruce wood samples 104 extracted with water and organic solvents. 105

The extractive content of heat-treated wood increased when treatment was carried out at low 106 temperatures and decreased with treatment carried out at higher temperatures (> 220 °C). 107 108 Most of the raw extractives disappeared and new compounds, such as anhydrosugars, mannosan, galactosan, levoglucosan and two C5 anhydrosugars, were generated. 109 110 Syringaldehyde, sinapaldehyde and syringic acid appeared to be the products formed in the largest amounts, all of which came from lignin degradation (Esteves et al. 2008). This 111 hypothesis regarding the formation of new extractives and their role in heat-treated wood 112 durability therefore remains to be confirmed. 113

The resistance of thermally modified wood to termites has also been studied. Contrary to fungal resistance, the influence of heat treatment on wood resistance to termites appeared to be more variable depending on the processing conditions and wood species. In many cases, resistance to termites was random and, in some cases, even reduced (Doi *et al.* 1998; Momohara *et al.* 2003; Nunes *et al.* 2004; Sivrikaya *et al.* 2015; Salman *et al.* 2016). Some

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studies revealed that thermally modified wood had lower termite resistance than untreated 119 wood (Sivrikaya et al. 2015; Salman et al. 2016, 2017). Generally, termite survival levels 120 121 reveal an effect of thermal treatment on the biology of these insects. Termite survival rates remained virtually constant for treatments carried out at a low temperature (180 °C to 210 °C) 122 123 and were lower for higher treatment temperatures. Treatments over 200 °C induced crucial modifications to the wood material, possibly reducing suitability for termites (Candelier et al. 124 2017). As heat-treated wood is consumed when exposed to termites, the toxicity of ingested 125 126 components may be the reason for the higher mortality rate observed (Surini et al. 2012) 127 when treatments are carried out at high intensities (temperature – duration). In addition, when 128 heat-treated and untreated control wood samples were exposed side-by-side (choice test), termites preferred to attack the untreated wood samples (Nunes et al. 2004). 129

In some cases, when the thermal modification process was carried out under optimum 130 conditions, heat treatment enabled low natural durability timbers to be used by making them 131 more resistant to decay, allowing them to be used in use classes 2 and 3 (use class 4 being 132 excluded due to the occurrence of soft rots) (EN 335 CEN 2013), thus up-grading their 133 economic value (Kamdem et al. 2002). These improved biological properties conferred to the 134 135 wood by chemical modifications have a negative impact on the mechanical resistance of heattreated wood (Dilik and Hiziroglu 2012; De Oliveira Araujo et al. 2016). Surface hardness 136 seems to be slightly enhanced, while other characteristics, such as bending and compression 137 strengths, stiffness and shear strength, are considerably weakened depending on the kind of 138 139 thermal processes and treatment intensities used (Boonstra et al. 2008; Hannouz et al. 2015).

The objective of this study was to investigate the antifungal and anti-termite activities of extracts from thermally modified ash wood depending on different treatment intensities. In order to ensure the reproducibility of the thermal treatment processes, as well as the quality of the treated wood products, the heat treatments were carried out under monitored conditions, using different heating temperatures (170, 200, 200, 215 and 228 (°C)) and a fixed residence

145 time (2 h).

146 2. MATERIALS AND METHODS

147 **2.1. Wood samples**

148 Untreated ash (Fraxinus excelsior L.) wood samples were obtained from a French wood company (Bois Durables de Bourgogne, 71120 Vendenesse-lès-Charolles, France). Twenty 149 boards measuring 4000 x 110 x 25 (mm³) [L x R x T] were selected. Special attention was 150 paid to ensure these planks only had small variations in density (around 650 kg/m³ \pm 10 %) 151 and a uniform width of annual rings. All the planks were sawn into two equal parts of 2 m in 152 153 length. Half of each plank was used as reference material and the other half was thermally treated at different treatment intensities. The planks were then dried at 103°C in an industrial 154 oven up to mass stabilization (m_0) . 155

156

157 2.2. Heat treatment protocols

The thermal treatment processes were carried out at the same company. These operations were carried out in a 20 cubic meter industrial oven (Jartek[®]), in convection heat-transfer mode and under a steam pressure process [ThermoWood® method, Finnish Thermowood Association (2003)].

162 The temperatures of the wood core and that of the oven atmosphere were dynamically 163 recorded and monitored throughout the treatment (Figure 1), in order to optimize the process 164 and the final wood qualities.

165 The oven temperature was firstly increased rapidly (0,5 °C/min) to 100 °C, then slowly (0,2

166 °C/min) to 150 °C. This temperature level was maintained for 1 h. The temperature was

- 167 finally increased (0,2 °C/min) to the desired level [from a low temperature (170 °C to 200 °C)
- to a high temperature (215 °C to 228 °C)] and kept at that constant level for 2 hours. Steam

- 169 was injected into the chamber to keep it oxygen-free and to influence chemical changes in the
- 170 wood. The heating system was then stopped and the wood samples were cooled to room
- temperature in an oxygen-free atmosphere with a water spray.
- 172 Each heat treatment was carried out simultaneously on four ash wood planks.

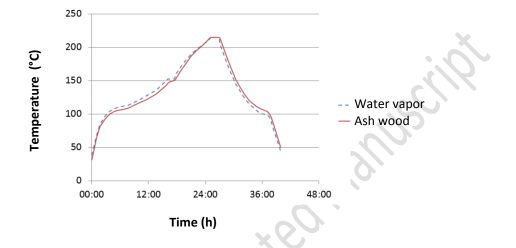


Figure 1: Temperature kinetics of ash wood and water vapor during heat treatment carried
out at 215°C for 2 hours.

175

176 2.3. Mass Loss (ML) due to wood thermal degradation

Each raw ash wood plank measuring 2000 x 110 x 25 (mm³) [L x R x T] was dried at 103 °C up to mass stabilization and its anhydrous mass was measured (m₀). After a thermal modification process each treated ash wood plank was again dried at 103 °C and its anhydrous mass was re-measured (m₁). The Mass Loss (ML %) due to wood thermal degradation depending on treatment conditions was determined by the following equation (Eq.1).

183

184
$$ML(\%) = [(m_0 - m_1) \div m_0] \times 100$$
 (1)

185

186 **2.4. Biological resistance tests on wood block samples**

187 2.4.1. Decay resistance

Decay and termite resistance tests were carried out according to adaptations of XP CEN/TS 189 15083-1 (CEN 2006) and EN 117 (CEN 2012) standard criteria, respectively, in one of our 190 previous studies (Candelier *et al.* 2017). The protocols used for these durability tests are 191 detailed in that previous work.

192

193 **2.5. Extraction protocols**

All the experimental procedures used to determine extractive contents were adapted, with minor modifications, from procedures found in the scientific literature (Rowell *et al.* 2005).

A mix of several pieces (One 20-cm long piece, per wood board) from untreated and heattreated planks was crushed and sieved to obtain particle sizes from to 0,2 and 0,5 mm, for each temperature level. After a drying step at 103 °C up to mass stabilization (m_2), the sawdusts (10 g) were separately extracted in a Soxhlet with acetone [Sigma Aldrich, 32201-M] (6 h), or with hot water (2 x 6 h), and dried at 103 °C for 48 h to obtain the anhydrous mass (m_3).

For untreated and heat-treated ash wood under varying temperature conditions, extractive contents were determined by the following equation (Eq.2):

204

$$Ext.(\%) = [(m_2 - m_3) \div m_2] \times 100$$
(2)

205

206 **2.6. Antifungal activity test**

Antifungal activities against the growth of *Trametes versicolor* Quélet (TV) [Linnaeus, CTB
863 A] and *Rhodonia placenta* Coocke sensu J. Erikson (PP) [Fries, FPRL 280] were tested as
per Salem *et al.* (2014) with some variations.

Each Petri dish (9 cm diameter) was filled with 10 mL of malt-agar medium containing

211 1200 μL of the diluted extracts in water or acetone (Merck company, Darmstadt, Germany)

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[C= 2.5% m/m] and left to solidify. These dilutions were chosen according to past studies 212 (Esteves et al. 2011; Pal et al. 2011). A 1-cm square section of a seven-day-old culture of 213 214 Rhodonia placenta or Trametes versicolor was placed in the center of the Petri dish and stored in a climatic chamber regulated at 22 °C \pm 2 °C and 70 % \pm 5 % relative humidity (RH) for 215 216 seven days' incubation. Three replicates were carried out per diluted wood extract sample and per fungus. Three media free of extractives and without solvent were used as a control. In 217 addition, three media free of extractives and with solvent (water or acetone alone) were used 218 to check that the water and acetone solvents did not have any impact on the activity of each 219 220 fungus. Following the fungal exposure period, the mycelium growth diameter was measured 221 in millimeters, for the control medium (M_c) and the extract-supplemented medium (M_t) . The percentage of mycelium growth inhibition was determined by equation 3 (Eq.3): 222

% inhibition =
$$[(M_c - M_t) \div M_c] \times 100$$
 (3)
mite activity test

224

225 2.7. Anti-termite activity test

226 The anti-termite activities of each heat-treated ash wood extract were tested by screening tests. Anhydrous cellulose paper measuring 2,5 cm in diameter was weighed (m₄) and then 227 impregnated with 70 μ L of the diluted extracts in water or acetone [C= 2,5% m/m], air dried 228 $(20 \text{ }^{\circ}\text{C} \pm 2 \text{ }^{\circ}\text{C} \text{ and } 65 \text{ }^{\circ} \pm 5 \text{ }^{\circ}\text{)}$ for 2 hours (m₄), then placed in the center of a Petri dish (5.5 229 cm diameter). Fifteen grams of wet sand (4 vol. sand/1 vol. water) was placed evenly around 230 231 the paper and 20 termite workers (*Reticulitermes flavipes*, ex. santonensis) were added to each 232 test set-up. Three replicates were carried out for each diluted wood extract. Three papers 233 impregnated with water or acetone alone were tested to estimate the impact of the solvent on 234 termite activity. Tests with water only were considered as a virulence control. For each test 235 set-up, the paper samples were placed on a plastic grid. Lastly, three diet control set-ups 236 containing only 15 g of wet sand and 20 termites were used to check termite survival without any feeding possibilities/without any trophic sources. All test set-ups were kept at 27 °C and >
75 % RH (Figure 2).

Every two days, each test set-up was observed to check sand humidity, add water if needed and keep track of termite behavior and activity. When all the termites contained in the diet control set-ups had died, the test was stopped. The termite survival rate was then determined, the anhydrous mass of the cellulose papers was measured (m_5) and the Weight Losses (WL_{term.} %) due to termite degradation were calculated by the following equation 4 (Eq. 4) :

 $WL_{term.}(\%) = [(m_4 - m_5) \div m_4] \times 100$ (4)

245



246

247

Figure 2: Termite screening test set-ups at the start of termite exposure.

248

249 **2.8. GC–MS analysis**

The wood extracts (acetone and water) were analyzed on a Perkin Elmer Clarus 680 Gas Chromatograph (GC) with a fused silica DB-5MS [(diméthyl-/diphényl-polysiloxane, 95:5]

column (30 m, 0,25 mm, 0,25 μm), coupled with a Perkin Elmer Clarus SQ8 Mass
Spectrometer (MS) and monitored by Turbo Mass v.6.1 software.

254 Before being injected into the GC-MS analyzer, the extractive substance samples were derivatized in order to improve the detection of all chemical compounds. To perform this 255 256 silvlation derivatization process, 2 mg of extract was solubilized in a glass tube with 50 μ L of 257 BSTFA + 1 % TMCS solution (Bistrimethylsilvltrifluoroacetamide + Trimethylchlorosilane) [Acros Organics]. The glass tube was sealed and dried in an oven at 70 °C for 120 minutes, 258 259 then opened to evaporate the BSTFA. The derivatized wood extract was dissolved in 1 mL of C₄H₈O₂ solution (ethyl acetate [Acros Organics]). One µL of this solution was injected into 260 261 the Gas Chromatograph at a 250 °C inlet temperature in splitless mode. Helium was used as the carrier gas. The temperature program was: 80 °C (2 min), 10 °C/min to 190 °C 15 262 °C/min⁻¹ to 280 °C - maintained for 10 minutes, 10 °C/min to 300 °C 15 °C/min⁻¹ -263 264 maintained for 14 minutes. A helium flow of 1 mL/min was used as the mobile phase. After this separation step, compounds were transferred to the Mass Spectrometer by a transfer line 265 heated at 250 °C and ionization was achieved by the Electron Impact method (70 eV 266 ionization energy). 267

The recognition of each component was achieved by comparing its mass spectrum with the NIST Library 2005 using NIST MS Search 2.0 (2011) software. The identification was deemed to be relevant when the comparison coefficient was higher than 900 (The comparison coefficient corresponds to the match factor between the mass spectrum obtained by a GC-MS analysis and the mass spectrum from the NIST MS Search 2.0 (2011). The match factors ranged from 0 to 1000, with 0 meaning no match and 1000 meaning a total match).

274

275 **2.9. Determination of pH Value**

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pH measurements were conducted according to Wang *et al.* (2008). Untreated and heattreated samples were ground and passed through 40-60 mesh screens. Three grams of ovendried sawdust samples was soaked in 30 mL of distilled water and then stirred for 5 min, allowed to stand for 15 min, stirred for another 5 min, and then left to stand for another 20 min. After this procedure, the pH values of the liquid were determined using a pH meter (PH/mVmeter Knick 911 ATEX; Knick ElektronischeTM, Berlin, Germany).

282

283 **2.10. Statistical Analysis**

284 The impact of heat treatment intensity compared to the untreated ash wood on (i) the termite 285 and fungal resistance of the wood samples, (ii) the content and (iii) the effects of water and 286 acetone extractives on their antifungal and anti-termite efficiency were evaluated using an ANOVA (one-way analysis of variance) and Duncan's comparison test. These statistical 287 analyses were carried out by the JMP 10.0.2 program (SAS 2012) by applying the Fisher test. 288 The results were then ranked into several categories, from "a" to "e" for the water-extractives 289 and from "A" to "E" for the acetone-extractives. The impact of a parameter on a system not 290 291 connected by the same letter was considered as non-significant at the 5 % level.

292

293 3. RESULTS AND DISCUSSION

3.1. Biological resistance tests on wood block samples

Decay and termite resistance tests were previously undertaken in Candelier *et al.* (2017). In order to put into perspective, the biological durability of untreated and heat-treated solid wood with the following antifungal and anti-termite activities of the respective wood extractive fractions, this section focuses solely on the main results obtained by Candelier *et al.* (2017).

3.1.1. Decay resistance

300 Similar results were found for both of the rots tested. However, *Rhodonia placenta* was the 301 most aggressive rot on beech control samples (WL 49,9 %), whereas *Trametes versicolor* was

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the most degrading rot on ash control samples (WL 48 %). Thermal modification increased
the durability of all wood materials, which was in agreement with previous studies (Kamdem *et al.* 2002; Esteves and Pereira 2009). These results were expected as Rousset *et al.* (2004)
and Metsä-Kortelainen *et al.* (2005) also found that the thermal treatment of wood at high
temperatures increases the resistance of wood to decay.
According to Table 1, the thermal treatment carried out at temperatures over 200 °C conferred
the "very durable" durability class 1 to the heat-treated wood materials, according to the

classification method of XP CEN/TS 15083-1 (CEN 2006).

310

Table 1: Resistance of untreated and heat-treated ash wood block samples to *Trametes versicolor* and *Rhodonia placenta* fungi and to *Reticulitermes flavipes* termite species [taken
from Candelier *et al.* (2017)].

| | Temperature | De | ecay resistance | | Termite resistance (Reticulitermes flavipes) | | | | | |
|-----------------|----------------------------|------------------------|----------------------|------------|--|-------------------|------------------|--|--|--|
| Wood species | of thermal modification | Trametes versicolor | Rhodonia placenta | Durability | ML (%) | Survival rate (%) | Visual rating * | | | |
| | process (°C) | WL (%) | WL (%) | class | | | | | | |
| Beech | Control | 23,0 (0,13)a | 49,9 (0,11)a | 5 | 8,87 (1,98)abc | 69,50 (13,10)ab | 4 | | | |
| Pine | Control | n.c | n.c | n.c | 11,12 (1,57)a | 64,50 (9,43)bc | 4 | | | |
| | Control | 48,0 (0,10)b | 39,7 (0,09)ab | 5 | 4,58 (0,57)d | 69,67 (8,33)ab | 4 | | | |
| Ash | 170 | 5,02 (0,05)c | 6,2 (0,12)c | 2 | 11,80 (2,60)a | 78,67 (4,16)a | 4 | | | |
| ASII | 200 | 2,8 (0,13)d | 2,2 (0,09)d | 1 | 11,60 (2,12)a | 71,33 (6,11)ab | 4 | | | |
| | 215 | 1,5 (0,19)e | 0,9 (0,07)e | 1 | 10,86 (1,51)a | 56,67 (5,03)bcd | 4 | | | |
| | 228 | 1,1 (0,10)f | 0,7 (0,05)e | 1 | 9,23 (1,97)ab | 51,33 (13,61)bcd | 4 (67%); 3 (33%) | | | |

WL% - Median values

ML% - Average values

* "0" for no attack "1" for attempted attack, "2" for slight attack, "3" for average attack, "4" for a strong attack

** According to a one-way analysis of variance, systems not connected by the same letter are largely different at the 5 % level.

314

315 **3.1.2. Termite resistance**

316 Termite survival revealed an effect on termite biology. Although the survival rate was similar

for treatments at 170 °C and 200 °C, it was lower above those temperatures (Table 1). Heat

treatments at temperatures over 200 °C caused critical changes to the wood and consequently

its durability could be improved. As the wood was nonetheless degraded, the toxicity of the

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320 components consumed may explain the higher termite mortality rate (Surini et al. 2012) 321 depending on the increase in heat treatment intensity. However, it was not particularly 322 effective over a short time scale, as 51,33 % of the termites were still alive after the test, with no significant differences between treated and untreated wood. The current results are in 323 324 agreement with those of Nunes et al. (2004), who studied the resistance to termites of the 325 species Reticulitermes grassei with wood treated by the German method (OHT) and concluded that, in spite of the slightly higher mortality of termites in treated samples and a 326 327 smaller weight loss, the differences were not significant.

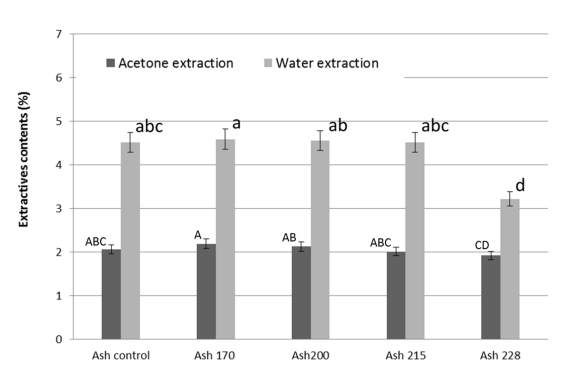
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329 **3.2. Extractive content**

For thermal treatment carried out at a low temperature (< 215 °C), the proportions of 330 heat-treated wood extractives increased in comparison with those of untreated ash wood, 331 whatever the solvent used during the extraction process (Figure 3). This result tallied with 332 past studies, indicating an increase in extractive rate resulting from the generation of different 333 degradation products (Poncsak et al. 2009; Esteves et al. 2008). A past study by Esteves et al. 334 (2008), carried out on eucalypt wood, showed that thermal degradation of lignin and 335 336 hemicelluloses led to an increase in extractive content. In fact, the formation of new extractive compounds arising from polysaccharide degradation at around 160 °C may be one of the 337 reasons for that phenomenon. At higher temperatures (>215 $^{\circ}$ C), these new compounds are 338 339 also generated but, under the heat effect, they are then converted into volatile products that 340 leave the structure of the wood, thus causing a loss of matter, leading to a decrease in 341 extractive content (Wang et al. 2016).

The relative percentage contents of extractives (Figure 3) increased at low treatment temperatures, reaching maximum values of 4,59 % and 2,19 % for thermal treatment carried out at 170°C and for water and acetone extractions, respectively. Beyond that temperature the

345 relative extractive percentage decreased.



Thermal treatment intensities (°C)

Figure 3: Extractive contents (%) of untreated and heat-treated ash woods at different
temperature levels (170 °C to 228°C), depending on the extraction solvent used.

349

350 **3.3. Antifungal activity test**

Fungal activity was inhibited by the heat-treated and untreated ash wood extracts. Table 2 and 351 Figure 4 shows how fungal development was hindered depending on the extractive 352 353 compounds. However, for both of the fungus species tested, the effectiveness of the extracts varied depending on the heat treatment temperature and the solvent used during the extraction 354 355 process. It clearly appeared that untreated and heat-treated ash wood extracts were more 356 efficient against brown-rot (*Rhodonia placenta*) than against white-rot (*Trametes versicolor*) 357 growth, whatever the extraction solvent used. In both cases, the efficiency of the extracts 358 seemed to be correlated to the heat treatment temperature.

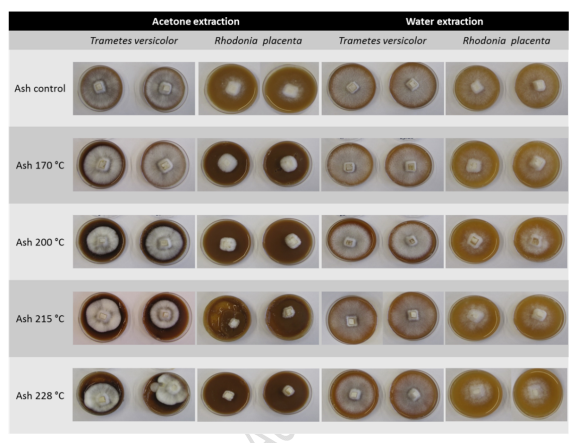


Figure 4: *Trametes versicolor* and *Rhodonia placenta* growth depending on the inhibition of
 untreated and heat-treated ash wood extracts (1200 μL), after seven days.

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- 372

Table 2 indicates that heat-treated ash wood extracts were more efficient against *Trametes versicolor* and *Rhodonia placenta* after 3,5 days than after 7 days, except for the modified ash wood extracts extracted with acetone against *Rhodonia placenta*. It turned out that, 7 days after fungal exposure, acetone-extracts from ash wood modified at a temperature of 228 °C delayed the growth of *Trametes versicolor* and *Rhodonia placenta* most.

Maderas-Cienc Tecnol 22(2):2020 Ahead of Print: Accepted Authors Version

| 373 Table 2: I | Effectiveness | of untreated | and | heat-treated | ash | wood | extracts | (1200) | μL) | against |
|-----------------------|---------------|--------------|-----|--------------|-----|------|----------|--------|-----|---------|
|-----------------------|---------------|--------------|-----|--------------|-----|------|----------|--------|-----|---------|

| | | nal ent ature Inhibition (%) Standard deviation (%) Inhibition (%) Stan dev (%) 24,01 ^D $1,40$ $31,49$ ^C 1 24,01 ^D $1,40$ $31,49$ ^C 1 37,83 ^{BC} $6,98$ $32,93$ ^{BC} 5 44,74 ^B $2,79$ $42,31$ ^B 2 48,68 ^{AB} $8,37$ $46,63$ ^{AB} 8 3 $59,54$ ^A $6,98$ $51,68$ ^A 7 14,14 ^e $1,40$ $31,49$ ^c 1 20,07 ^d $1,40$ $32,21$ ^{abc} 6 | | | | Rhodonia placenta | | | | | |
|----------------------|-------------------------------------|---|-----------|----------------------|------------------------------|---------------------|------------------------------|---------------------|------------------------------|--|--|
| | Ash wood | 3,5 0 | days | 7 d | ays | 3,5 0 | days | 7 d | ays | | |
| Extraction method | Thermal treatment temperature | | deviation | | Standard deviation (%) | Inhibition (%) | Standard deviation (%) | Inhibition (%) | Standard deviation (%) | | |
| | 0 | 24.01 ^D | 1,40 | 31.49 ^C | 1,02 | 42,05 ^D | 0,00 | 44,90 ^E | 0,00 | | |
| | 170 | 37,83 ^{BC} | | 32,93 ^{BC} | 5,10 | 50,57 ^C | 2,41 | 61,22 ^D | 2,89 | | |
| Acetone | 200 | 44,74 ^B | 2,79 | 42,31 ^B | 2,04 | 42,05 ^{CD} | 9,64 | 68,37 ^C | 1,44 | | |
| rectone | 215 | 48,68 AB | 8,37 | 46,63 AB | 8,37 | 60,80 ^{AB} | 2,41 | 76,53 ^{AB} | 1,44 | | |
| | 228 | 59,54 ^A | 6,98 | 51,68 ^A | 7,14 | 64,20 ^A | 2,41 | 79,63 ^A | 1,44 | | |
| | | | | | | | | | | | |
| | 0 | 14,14 ^e | 1,40 | 31,49 ^c | 1,02 | 47,16 ^a | 2,41 | 14,29 ^d | 2,89 | | |
| | 170 | 20,07 ^d | 1,40 | 32,21 ^{abc} | 6,12 | 42,05 ^b | 0,00 | 24,49 ^{bc} | 2,89 | | |
| water | 200 | 38,82 ^{bc} | 2,79 | 35,82 ^b | 1,02 | 45,45 ^a | 0,00 | 32,65 ^a | 5,77 | | |
| | 215 | 35,86 ^{ab} | 1,40 | 36,54 ^{ab} | 0,00 | 48,86 ^a | 0,00 | 35,71 ^{ab} | 1,44 | | |
| | 228 | 41,78 ^a | 1,40 | 40,14 ^a | 3,06 | 43,75 ^{ab} | 2,41 | 37,76 ^{°a} | 1,44 | | |
| Water cont. | / | 0 | / | 0 | | 0 | / | 0 | / | | |
| Acetone cont. | / | 0 | / | 0 | | 0 | / | 0 | / | | |

374 *Trametes versicolor* and *Rhodonia placenta*.

375

The different degrees of fungal development for untreated and thermally modified wood reflecting the contrasts in extractive chemical composition are displayed in Figures 6 and 7. Although some effects of water and acetone extracts from heat-treated ash could be seen on fungal growth depending on the treatment intensity, this phenomenon was not significant. It appeared that the fungi adapted themselves to the new medium and regained normal growth after a few days of incubation. In other words, heat-treated wood extracts seem to have short-term fungistatic action.

383

384 **3.4. Anti-termite activity test**

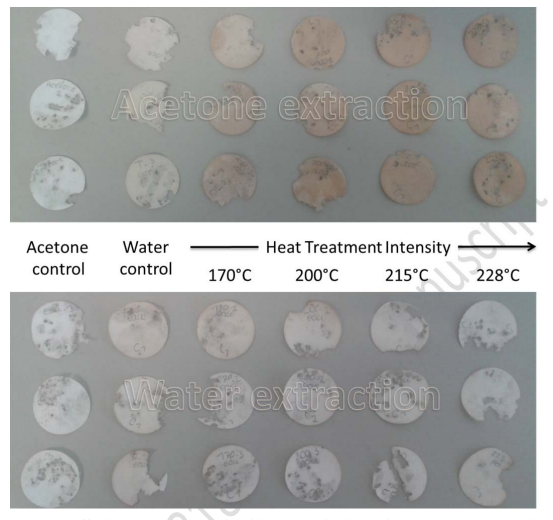
Table 3 and Figure 5 show that the heat-treated ash extract yields and their anti-termite activities were largely linked to the polar or nonpolar nature of the solvent used during the extraction process. In fact, anti-termite compounds can have many varied chemical 388 characteristics and different polarity levels, which may or may not be soluble in a specific

- solvent (Kadir *et al.* 2015).
- 390 The test was stopped after 21 days, when the termites in the diet control set-ups had died. The
- results in Table 2 show that the anti-termite activity of the heat-treated ash wood extracts was
- 392 not significant.
- 393 Table 3: Effectiveness of untreated and heat-treated ash wood extracts (1200 µL) against
- 394 *Reticulitermes flavipes.*

| Extraction solvent | Ash wood Thermal treatment temperature | Weight loss (%) | Standard deviation (%) | Survival rate (%) | Standard deviation (%) |
|-----------------------|---|------------------------|------------------------------|-----------------------|------------------------------|
| | 0 | 45,25 ^A | 9,31 | 85,00 ^A | 5,00 |
| Acetone | 170 | 34,86 ^{AB} | 1,52 | 86,67 ^A | 7,43 |
| C=2,5% [m/m] | 200 | 34,49 ^{AB} | 4,39 | 81,67 ^{AB} | 2,89 |
| | 215 | 32,17 ^{AB} | 4,02 | 76,67 ^{ABC} | 5,77 |
| | 228 | 31,26 ^B | 1,84 | 86,67 ^A | 2,89 |
| | 0 | 54,62 ^{abc} | 6,78 | 90,00 ^a | 5,00 |
| Water | 170 | 0 | | 73,33 ^{cd} | 2,89 |
| C=2,5% | 200 | 53,25 ^c | 3,36 | 80,00 ^c | 0,00 |
| [m/m] | 215 | 59,55 ^{ab} | 2,52 | 76,67 ^{bc} | 5,77 |
| | 228 | 61,12 ^a | 3,05 | 90,00 ^a | 5,00 |
| | Acetone | 32,77 ^{AB/d} | 6,63 | 86,67 ^{A/ab} | 2,89 |
| Control | Water | 31,98 ABC/de | 6,29 | 88,33 ^{A/ab} | 5,77 |

395

For untreated and treated ash wood under different treatment intensities, the mass loss due to termite degradation indicated that the relative extracts obtained with water were statistically more attractive for termites and the termite survival rate was quite similar to that of the control sample. This effect was related to the final temperature level of the thermal treatment. On the other hand, the same extractives obtained with acetone seemed to have no significant effect on termites compared to the water control samples (Figure 5).



403

404 Figure 5: Effectiveness of untreated and heat-treated ash wood extracts (1200 μL) against
 405 *Reticulitermes flavipes.*

The termite survival rate revealed no significant effect on the biology of these insects. The survival rate was not significantly dissimilar for the different treatments compared to the control samples. The effects of untreated wood extracts on the termite survival rate were very similar to those for heat-treated wood, confirming the innocuous property of these treated wood extracts in terms of termite toxicity.

411

412 **3.5. GC-MS analysis**

413 **3.5.1. Water extractives**

The typical chromatograms from GC-MS analyses of the chemical composition of untreated 414 415 and thermally modified ash wood extracts by the water extraction process are presented in Figure 6. 416

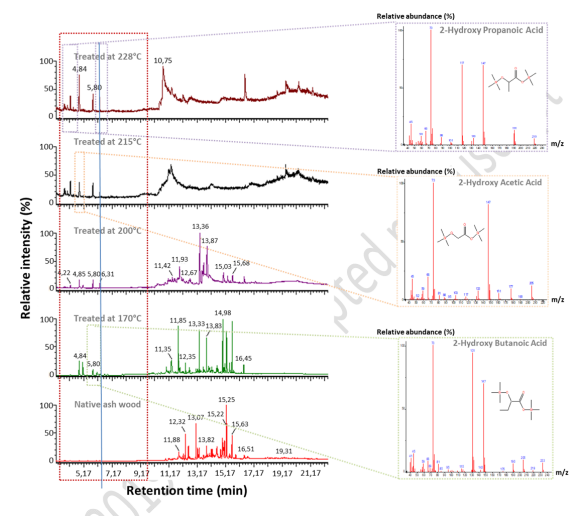


Figure 6: Untreated and heat-treated ash wood water-extractive composition identified by 418 419

420

417

GC-MS, depending on the thermal treatment temperature.

For untreated ash wood, the GC-MS analysis highlighted that almost all of the original water 421 extractives resulted from polysaccharides or their silvlated fragments. Only the weak peak 422 11,88 min corresponded to the 423 observed antioxidant Tyrosol (4at rt = 424 hydroxyphenylethanol).

Maderas-Cienc Tecnol 22(2):2020

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For the ash wood treated at 170 °C, it was found that the respective water extracts mainly 425 comprised low molecular weight organic 2-hydroxy-acids [propanoic acid (rt = 4,84 min and 426 6,09 min), acetic acid (rt = 5,10 min), butanoic acid (rt = 5,80 min)] (Table 5) and derivated 427 phenylpropanoid compounds (rt = 11,00 min and 11,35 min). The presence of these acid 428 429 compounds, decreasing the pH of the wood, may have played a role in fungal inhibition. Indeed, according to the results presented in Table 4, the pH values of ash wood decreased 430 depending on the temperature used during the heat treatment process. The pH values varied 431 432 from 5,16 for untreated wood to 3,92 for wood thermally modified at 228 °C. Similar trends 433 were observed on softwood and hardwood species by Niemz et al. (2008), showing that the 434 pH decreases with increasing heat treatment intensity.

| 435 | Table 4: | pH values of | untreated a | and heat-treated | ash wood. |
|-----|----------|--------------|-------------|------------------|-----------|
|-----|----------|--------------|-------------|------------------|-----------|

| Wood species | Thermal treatment temperature (°C) | рН |
|--------------|---------------------------------------|----------------------------|
| | 0 | 5,16 (0,15) ^a |
| | 170 | $4,51(0,16)^{b}$ |
| Ash wood | 200 | $4,12(0,17)^{c}$ |
| | 215 | 4,03 (0,16) ^{cd} |
| | 228 | 3,92 (0,14) ^{cde} |

436

Indeed, Yalcin and Sahin (2015) reported that heat treatment conferred on narrow-leaved ash 437 wood a pH that was less conducive to fungal growth, with the untreated wood pH of 5,7 438 falling to 3,9 in the modified wood, while the pH for optimum fungal growth is around 5-6 439 440 (Bozkurt et al. 1993). In addition, such a decrease in wood pH could disturb the enzymatic digestion of termites during their wood destruction (Lima et al. 2014). In fact, pH widely 441 442 affects the activity of fungal enzymes. The pH dependence is usually due to the side groups of the amino acids. A diminution in pH changes the protonation pattern resulting in protein 443 denaturation (in shape, charge and location within the substrate) (Purich 2010). 444

Maderas-Cienc Tecnol 22(2):2020

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For ash woods treated at higher temperatures, from 200 °C to 228 °C, only traces of the 445 previously mentioned low molecular weight organic acids were observed. In addition, a large 446 peak located in the polysaccharide area, from rt = 10 min and rt = 13 min, was observed but 447 remained unidentified. These last peaks may have been due to the condensation of several 448 chemical compounds produced by wood thermal degradation and might also partially explain 449 450 the greater fungal resistance of wood thermally treated at high temperatures (Hakkou et al. 2006). 451

452

3.5.2. Acetone extractives 453

The typical chromatograms from GC-MS analyses for the chemical composition of untreated 454

455 and thermally modified ash wood extracts by the acetone extraction process are presented in

456 Figure 7.

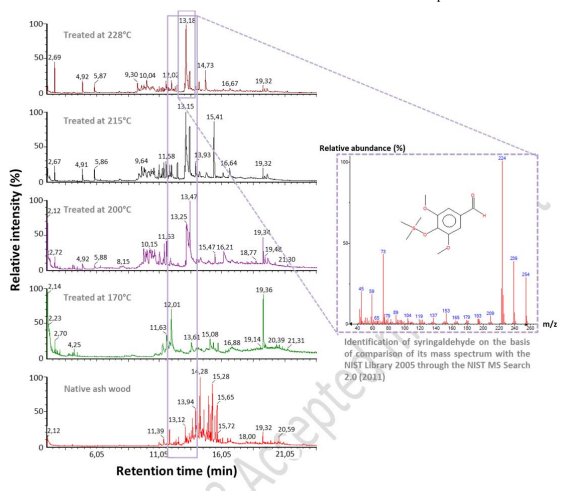


Figure 7: Untreated and heat-treated ash wood acetone-extractive composition identified by
 GC-MS, depending on the thermal treatment temperature.

460

The GC-MS analysis revealed that almost all of the untreated acetone-extractives disappeared after heat treatment, but on the other hand the thermal degradation of hemicelluloses and lignin generated new compounds. Monosaccharides and some products derived from their dehydration reactions, along with syringaldehyde (13,18 min, Figure 7), were the most abundant products generated by the heat treatment. Syringaldehyde appeared to be the product formed in the largest amounts, arising from lignin degradation, and was generally the major detected component according to the TIC chromatograms, Table 5.

- 468
- 469
- 470

| | Thermal | Propanoic acid | | | | Acetic acid | | But | anoic acid | Syringaldehyde | |
|-----------------|----------------------------------|----------------|--------------------|-------------|-------------------|-------------|------------------|-------------|------------------|----------------|------------------|
| Wood species | treatment temperature (°C) | Rt * (min) | Intensity** (%) | Rt (min) | Intensity (%)* | Rt (min) | Intensity (%) | Rt (min) | Intensity (%) | Rt (min) | Intensity (%) |
| | Control | 4,21 | 1 | 6,31 | 1 | 4,84 | 5 | 5,8 | 2 | 13,12 | 28 |
| | 170 | 4,21 | 3 | 6,31 | 4 | 4,84 | 22 | 5,8 | 6 | 13,11 | 12 |
| Ash | 200 | 4,22 | 9 | 6,31 | 12 | 4,85 | 18 | 5,8 | 19 | 13,25 | 67 |
| | 215 | 4,23 | 37 | 6,31 | 21 | 4,85 | 39 | 5,8 | 38 | 13,15 | 100 |
| | 228 | 4,23 | 39 | 6,32 | 37 | 4,84 | 78 | 5,8 | 41 | 13,18 | 100 |

| 471 | Table 5: | Retention | time and | Peak | Intensities | from | GC-MS | analyses | of | untreated a | and h | neat- |
|-----|----------|-----------|----------|------|-------------|------|-------|----------|----|-------------|-------|-------|
| | | | | | | | | | | | | |

* Rt = Retention time (in minutes) shows the time taken for the analytes to pass through the column and reach the mass spectrometer detector.

** Intensity (as a %) represents a reflection of the amount of a specific analyte that was present, compared to the predominant one (e.g. Intensity = 100%)

472 treated ash wood extracts.

473

Similar results for syringaldehyde extraction, by water and ethanol, from heat-treated wood 474 were found by Ibrahim et al. (2012). The syringaldehyde content increased depending on the 475 thermal treatment intensity, more particularly for wood treated at temperatures over 200 °C 476 (Figure 7), and might explain the fungal inhibition activities of the heat-treated wood 477 extractives. Indeed, previous studies highlighted the antifungal and anti-microbial activities of 478 the phenolic compound syringaldehyde. Murugesan et al. (2009) also showed that 479 syringaldehyde inhibited the growth of bacteria and of the fungus Ganoderma lucidum in 480 syringaldehyde-treated malachite green samples. De Souza et al. (2005) also reported that 481 syringaldehyde expounded anti-fungal activity against *Leucoagaricus gongylophorus*. 482 According to Ibrahim et al. (2012), syringaldehyde (also called 3,5-dimethoxy-4-483 484 hydroxybenzaldehyde), has the same kind of structure as vanillin and is a unique natural compound that has bioactive properties (antioxidant, antifungal, anti-microbial and anti-485 tumorigenesis), which belongs to the phenolic aldehyde family. In a past study, 486 487 syringaldehyde displayed a successful role in inhibiting fungal growth rates (Kelly et al. 488 2008). Its fungicidal effects were mainly due to aldehyde moiety and the hydroxyl substituent

present in syringaldehyde (De Souza et al. 2005). Total hydroxyl groups from lignin also had 489 490 a large impact on cellulase adsorption and enzymatic hydrolysis occurring during fungal 491 degradation (Yu et al. 2014), and more particularly acetosyringone, vanillin and syringaldehyde (Qin et al. 2016). Ximenes et al. (2011) confirmed that syringaldehyde and 492 493 vanillin inhibit cellulose hydrolysis. Although syringaldehyde displayed insecticidal properties against Acanthoscelides obectus beetles in a past study (Regnault-Roger et al. 494 2003), the concentrations tested in this work did not make it possible to confirm its 495 496 insecticidal behavior on termite activity.

497

498 **4. CONCLUSIONS**

Heat treatment improved wood durability, clearly increasing resistance to brown and white rots, but it had only a slight effect in improving resistance to termites. The process carried out at 228 °C was found to give ash wood the best biological resistance properties. However, it can be expected that such treatment will greatly degrade the mechanical properties of the wood.

In addition, the results presented in this work confirmed that thermal treatment generates new 504 extractive substances within the modified wood material, depending on the treatment 505 temperature level, which could act as fungistatics and prevent fungus from growing, 506 improving the resistance of the modified wood to decay. The antifungal activity of heat-507 508 treated ash wood extracts varied depending on heat treatment intensity and the solvent used during the extraction process, but these effects were not efficient over the long term and were 509 510 not significant. It also clearly appeared that extracts from heat-treated ash wood were more 511 efficient against brown-rot growth than against white-rot growth. The analysis of these extractive compounds showed that the most abundant was syringaldehyde. In addition, the 512 513 syringaldehyde content increased in line with the thermal treatment intensity, whatever the 514 extraction process used. Although syringaldehyde has shown insecticidal properties in past

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studies, it could not be shown in our study that it has an insecticidal role against termites. The 515 generation of low molecular weight organic acids might decrease the pH of wood, also 516 impacting fungal inhibition and the enzymatic digestive system of termites. Lastly, the 517 condensation of several chemical compounds from wood thermal degradation might also 518 partially explain the better decay resistance of wood treated at high temperature. However, 519 taken separately, the extractive compounds might act differently from the same extracts 520 present within the wood. The increased durability of heat-treated ash wood is due to thermal 521 522 degradation of wood cell wall polymers involving a combination of modifications: 523 hydrophobic behavior, lignin modification, hemicellulose degradation and also extractive 524 generation. To conclude, the results obtained in this study showed that the extractive compounds of heat-treated wood have a slight impact on the better durability of thermally 525 modified wood, but are certainly not the only reason. 526

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