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Article

Impact of High-Power Ultrasound for Barrel Regeneration on the Extraction of Wood Volatile and Non-Volatile Compounds

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Abstract: High-power ultrasound (HPU) is an innovative cleaning method used in wineries for oak barrel sanitation and regeneration. The process is associated with hot water (HPUhw) to ensure microbial stabilization and has been proved to be highly effective in recent years. This study thus examines the impact of different cleaning treatments on the subsequent extraction of wood compounds in wine and their impact on organoleptic properties. Red wines aging in barrels treated (HPUhw and steam) in different years (1, 2, and 3 years) were examined during the first 12 months for chemical exchange from wood to wine. Specific analyses were realized on ellagitannins, the physicochemical composition, and oak wood volatile compounds. Only a small increase in some wood volatile compounds occurred in the case of HPUhw, including furfural, 5-methylfurfural, trans-whisky lactone, vanillin, and syringaldehyde. The sensory analysis carried out by a panel of experts showed that the impact on the organoleptic properties of wines is similar with both processes (HPUhw and steam). However, since HPUhw treatment requires lower energy for the same efficiency, it could be an interesting alternative to steam treatment, given the promising prior microbial results.

Keywords: high-power ultrasound; aging; barrels; oak wood; sensory analysis; ellagitannins

1. Introduction

Oak wood releases several specific compounds into wine. Extraction of phenolic compounds from oak barrels depends on the quantity of extractable compounds; contact time; wine composition; and toasting, aging, and sanitation procedures. Conversely, when aging takes place in wooden barrels, bacteria and yeast can penetrate the wood to the same depth as the wine (close to 8 mm) [1]. Consequently, a frequent problem occurring in wines aged in barrels is the potential presence of ethylphenols and ethylguaiacol. Brettanomyces bruxellensis spoilage can lead to organoleptic deviations due to the production of several undesirable compounds, such as 4-Ethylphenol and 4-Ethylguaiacol [2], molecules that can deteriorate the wine quality [3]. Organoleptic deviations, known as ‘Brett flavors’, may appear, corresponding to ‘stable’, ‘leather’, ‘manure’, or ‘horse sweat’. To avoid such deviations, several treatments can be adopted in wineries. Many types of treatment are applied to sanitize barrels, such as the use of chemical agents (sulfur dioxide, ozone, sodium percarbonate, peracetic acid, and peroxide-based compounds) or physical agents (barrel rejuvenation, dry ice blasting, UV radiation, hot water, microwaves, and ultrasounds). These practices show varying levels of efficiency and may induce potential organoleptic modifications [4]. Steam treatments are still the most commonly used in wineries for two reasons: high efficiency and low impact on oak barrels. Solis (2018) found that steam was required for at least 10 min to achieve temperatures above 57.5 °C (the temperature required to kill Brettanomyces bruxellensis) [5] at a depth of 8 mm in a barrel.

Several studies have recently been published on the use of high-power ultrasound (HPU) on an industrial scale [6]. In this process, electrical energy is converted into ul-
Ultrasound (20 kHz–10 MHz) at frequencies higher than those audible to the human ear (16–20 kHz). High-power ultrasound has intensities over 1 W/cm² and frequencies between 20 and 100 kHz [7–9]. When emitted in a liquid, the process forms high-energy microbubbles (acoustic cavitation phenomenon) [10]. These cavitation bubbles generate very high temperatures locally (close to 5000 °C) and high pressure (above 50 MPa) [8]. The microcavitation bubbles (diameters around 1 µm) act homogeneously throughout the fluid (Pascal’s law) and can penetrate deep into the pores. Piyasena et al. (2003) identified a large number of pathogenic microorganisms inhibited by the cavitation phenomenon generated by HPU [11].

Porter et al. (2011) recently studied the effects of HPU treatment on porous materials and cleaning efficiency in American oak wine barrels using X-ray tomography. The authors demonstrated that industrial-scale high-power ultrasonic (4 kW/20 kHz) applications provide an efficient way to remove tartrate sediment from the surface of staves up to a depth of 2 mm, with a relatively short treatment time (12 min) and at quite low temperatures (40–60 °C) [12]. Oak wood consists of macromolecules, such as cellulose and hemicellulose, and extractive compounds, such as ellagitannins, lignin, and aromatic precursors. However, only a few studies have investigated the impact of these treatments on the release of oak wood non-volatile compounds. Breniaux et al. (2019) proved that the combined effect of HPU and heat treatment had a significant impact on wood sanitation, wood wettability, and its specific surface and oxygen transfer kinetics [13]. The authors showed that HPUhw treatment removed B. bruxellensis cells up to a depth of 9 mm, with processing parameters set at 60 °C/6 min at 3.8 kW. While the study characterized the oak wood ultrastructure, it did not take the impact of HPUhw on volatile and non-volatile compounds into consideration. Indeed, oak wood aging changes the wine composition as non-volatile and volatile compounds are extracted from the wood. The extraction phenomena depend on several factors, such as the wood’s geographic origin, the type of seasoning, toasting, and the length of contact between wine and wood [14–16]. During the aging process, the wine’s phenol composition is modified with the release of phenols like ellagitannins, increasing the wine’s astringency [17]. Finally, various volatile compounds are released from the wood during aging, such as cis and trans-whisky lactones (coconut notes); volatile phenols, such as guaiacol, 4-methylguaiacol and eugenol (spicy and smoky notes); aldehyde phenols like vanillin or acetovanillone (vanilla notes); or furfural derivatives (smoky and toasty notes) [18]. Nevertheless, according to a recent study [19], the mix of volatile compounds from wood at concentrations reflecting various toasting levels could have a masking effect on the fruity pool. Thus, the main challenge during barrel cleaning is not to leach the volatile and non-volatile compounds from the wood into the washing water. The final goal is thus to reuse them once or twice without microbial contamination while continuing to benefit from the wood.

Our study investigated the effect of HPU associated with hot water (HPUhw) and steam treatment for barrel cleaning and regeneration. Steam treatment was retained, as it is widely employed in wineries and has high microbial stabilization efficiency, while the HPUhw process can ensure complete microbial stabilization and avoid wine spoilage. The aim of the study was to examine whether aqueous steam or HPUhw has a significant effect on wine composition and perceptions of aroma according to the barrel’s age (1, 2, or 3 years). Taking these processes into consideration, we investigated (i) the extraction of oak volatile compounds, (ii) non-volatile compounds after one year of wine aging, and (iii) the impact on wine organoleptic properties by sensory analysis (olfactory and gustatory).

2. Materials and Methods

2.1. Experimental Setup

The French oak barrels used were 1, 2, or 3 years old, and the experimental setup was divided into 4 stages (Figure 1). For each modality, two barrels were first cleaned and regenerated by steam treatment (Barriclean®, Brive la Gaillarde, France) or by HPUhw as described in Section 2.2. Red wine was then introduced from two estates: Estate A and
Estate B. Wine from Estate A was put in the 1- and 3-year-old barrels and wine from Estate B in the 2-year-old barrels. The wines were aged in the barrels for 2, 4, 8, and 12 months, and several physicochemical and sensory analyses were realized in the final stages and during the aging process. No filtration or fining agents were used in these cases.

![Figure 1. Experimental design of the study.](image)

### 2.2. Barrel Treatment: High-Power Ultrasound (HPU) and Aqueous Steam Treatment

The HPU\textsubscript{hw} treatment process consisted of filling the barrel with water (heated to 60 °C by an autonomous system) and then introducing the sonotrode before the treatment modality was applied (partially allowing the ultrasound to be emitted) via the bung hole (Figure 2), thereby causing pressurization (0.3 bars). The treatment was applied for 6 min (frequency 20 kHz, 3.8 kW). The operating parameters had been optimized in earlier research [15].

![Figure 2. HPU apparatus.](image)
The treatment by aqueous steam was carried out using an autonomous boiler (Barriclean®, 18 kW), which supplied pressurized hot water inside the barrel for 10 min (1 bar, 100 °C). Steam modalities were only used to compare the results with HPU treatment (a reference modality that can ensure complete microbiological stabilization). Total energy consumption was 0.38 kWh for HPU and 3 kWh for aqueous steam, which is 7.89 times lower.

2.3. Physicochemical Parameters

The anthocyanin, tannin, colour intensity (CI), modified colour intensity (MCI), and total polyphenol index (TPI) values were determined by spectrophotometry methods using a Lambda 25 UV/Vis Spectrophotometer (Perkin Elmer, Waltham, MA, USA). TPI was measured at 280 nm after wine dilution (1/100), CI is the sum of optical density at 420 and 520 nm, and MCI is the sum of optical density at 420, 520, and 620 nm. The other physicochemical parameters of wines, such as pH, ethanol content (% vol.), and total and volatile acidity, were determined by Fourier transform infrared spectroscopy using an OenoFoss™ (Foss electric, Hilleroed, Denmark) that was previously calibrated with wine samples in accordance with OIV methods. Free SO₂ was measured with a Y15 enzymatic auto analyzer (Biosystems, S.A., Barcelona, Spain). The Y15 equipment was calibrated using the external standards provided in every kit by Biosystems. Free SO₂ was performed using the appropriate kits from Biosystems S.A. (Barcelona, Spain).

2.4. Search for Brettanomyces in Wine

Serial dilutions of wine samples were plated on solid YPG (yeast extract 10 g/L; bacto-peptone 10 g/L; glucose 20 g/L; agar 20 g/L; adjusted to pH 5), supplemented with antibiotics to limit the growth of bacteria, molds, and yeast of the Saccharomyces genus (0.1 g/L chloramphenicol; 0.15 g/L biphenyl; 0.5 g/L cycloheximide). Brettanomyces yeast colonies were counted after 7 days of incubation at 30 °C. All of the assays were performed in triplicate.

2.5. Determination of Volatile Wood Compounds

A quantitative determination of the volatile compounds extracted from oak wood was based on an adaptation of the method developed by Barbe and Bertrand (1962) [20]. We spiked 50 mL wines with a solution of dodecan-1-ol (200 µL) as an internal standard. Then, 3 liquid/liquid extractions were performed using 4, 2, and 2 mL of dichloromethane. The organic fractions were dried using sodium sulfate anhydrous and then concentrated to 250 µL under a nitrogen flush. All of the samples were analyzed in duplicate.

An Agilent HP 7890 GC (Les Ulis, France) was coupled with a mass spectrometer (HP 5975, electronic impact 70 eV, eMV = 2 kV). The quantitative determination was made in SIM mode, selecting ions of \( m/z = 99 \) for trans-3-methyloctano-4-lactone and cis-3-methyloctano-4-lactone (oak lactone), \( m/z = 96 \) for furfural, \( m/z = 110 \) for 5-methylfurfural, \( m/z = 109 \) for guaiacol, \( m/z = 164 \) for eugenol and isoeugenol, \( m/z = 154 \) for syringol, \( m/z = 194 \) for allylsyringol, \( m/z = 152 \) for vanillin, \( m/z = 166 \) for acetovinone, \( m/z = 182 \) for syringaldehyde, \( m/z = 196 \) for acetosyringol, and \( m/z = 83 \) for the internal standard (dodecan-1-ol). The column was BP21 (SGE) (50 m, 0.32 mm, 0.25 mm); the carrier gas was helium 5.5 (Linde, France, pressure: 70 kPa); temperatures were injector, 280 °C; detector, 280 °C; oven, 60 °C for 1 min programmed at 3 °C/min to 220 °C. The final step lasted 40 min; the split-less time was 30 s with a split flow of 30 mL/min.

2.6. Determination of Ellagitannins

Determination of total ellagitannins was realized as previously noted [21]. Each sample was analyzed in triplicate, and each reaction mixture was subjected to HPLC-UV using a Lichrospher100 RP 18 column, 250 × 4.6 mm, 5 µm. The apparatus used for the HPLC analysis was composed of a Finnigan Surveyor UV–Vis detector (UV–Vis 200), a Finnigan autosampler, and a Finnigan ternary pump. The mobile phases employed were solvent A (water/formic acid (99.9/0.1)) and solvent B (methanol/formic acid (99.9/0.1)),
and the gradient elution was 0–35% of B in 5 min, 35–4% of B in 25 min, and 45–100% of B in 5 min. The flow rate was established at 1 mL/min, with detection set at 370 and 280 nm.

2.7. Sensory Analysis

The impact of both processes on the olfactory and gustatory level was evaluated with a trained panel by sensory analysis. The samples were evaluated in individual booths in an appropriate sensory analysis room [22] with controlled temperature, lighting, and humidity, using black glasses [23] containing 50 mL of liquid, coded with random three-digit numbers, and with repetitions of tasting combinations. Before the tastings, the wine bottles were taken out of the room at 10 °C and left to rest to reach optimal room temperature. The sensory panels consisted of at least 24 trained individuals. All the panelists were research laboratory staff at ISVV, University of Bordeaux, selected for their experience. Descriptive sensory analyses were conducted of the red wine aged for 12 months in barrels of 1, 2, and 3 years treated with HPU or steam, employing scale descriptors graduating from 0 (low intensity) to 7 (high intensity). The intensity level of each descriptor was then expressed as the mean value of all the panelists.

2.8. Statistical Analysis

The statistical significance of the differences was determined via the Student test ($p = 0.05$). The XLSTAT program was used for data processing.

The results of all the triangular tests were statistically analyzed according to tables derived from the literature based on the binomial law corresponding to the distribution of answers in the test [24].

3. Results

3.1. Oenological Parameters

We analyzed two different wines aged in barrels of 1, 2, and 3 years treated with steam or HPU to evaluate the potential changes in the physicochemical and microbial properties of these wines before and after 12 months.

Comparisons of the impact of HPU$_{hw}$/steam and aging realized before and after treatment for both processes showed no evident changes in terms of colour, free SO$_2$, pH, total acidity, volatile acidity, or micro-organism content (Table 1). However, a significant increase in TPI was observed after one year of aging for both treatments and all modalities (1-, 2- and 3-year-old barrels for both estates). The increase in TPI before aging in oak barrels and after one year was the same in the case of steam and HPU$_{hw}$ treatments. The release of phenolic compounds was very similar for HPU$_{hw}$ and steam treatment in both cases (Estate A and Estate B). In addition, there were no significant differences in terms of classical chromatic characteristics (MCI and CI) between either modality during aging. Titratable acidity, volatile acidity, pH, and ethanol content did not change significantly following the two treatments. These results were expected, as neither barrel aging process affects such components in wine. However, oak volatile extraction is affected by pH and ethanol, and it was essential to measure these parameters. No significant differences were found, and the extraction of phenolic oak wood compounds appeared similar and led to no differences in terms of color. The results thus confirmed that the wine was homogenous upon filling, and that differences in wine composition should not have had a significant effect on the extraction of oak-related compounds in the different experiments. Finally, we noted no _Brettanomyces_ contamination before or after aging in either modality.

3.2. Oak Wood Non-Volatile Compounds: Ellagitannins

The increase in ellagitannin content during the wine aging process in oak barrels can be very interesting with regard to improving wine quality. In this study, HPU$_{hw}$ was used to step up the mass transfer of phenolics and was then compared to steam treatment. The comparison is interesting given the positive effect of temperature on the release of phenolics. This is in accordance with thermodynamic law, since the rising temperature can
increase the diffusivity and solubility coefficient [25]. Heat is also employed for HPU \(_{hw}\), but the temperature is lower (60 °C), and time treatment is shorter (6 min). Here, a comparative study was realized by examining total ellagitannin content to evaluate if the acoustic cavitation phenomenon could have an influence on extraction of oak wood compounds. Our results show that the total of ellagitannins released for HPU \(_{hw}\) was comparable to that obtained with steam treatment (Figure 3), apart from a significant improvement for HPU \(_{hw}\) treatment in a barrel of 1 year after 2 months of aging and in a barrel of 2 years after 4 months of aging. Otherwise, while there is a tendency towards higher extraction in the case of HPU \(_{hw}\), the statistical analysis did not reveal a significant difference. Furthermore, with regard to the mechanism of mass transfer of phenolics in wine following ultrasonic field and steam treatment, the release of total phenolics was very similar. We can assume that alcohol contact time is mainly responsible for ellagitannin desorption in both cases. Thus, the accessibility and diffusivity of phenolic compounds remain the same in these treatments after 12 months of aging.

**Table 1.** Physicochemical and microbial properties of wines detected before and after 12 months of aging.

<table>
<thead>
<tr>
<th>Wine before Aging</th>
<th>Estate A</th>
<th>Estate B</th>
<th>Steam</th>
<th>HPU (_{hw})</th>
<th>Steam</th>
<th>HPU (_{hw})</th>
<th>Steam</th>
<th>HPU (_{hw})</th>
</tr>
</thead>
<tbody>
<tr>
<td>Estate A</td>
<td>50.9 ± 0.6</td>
<td>46.5 ± 1.0</td>
<td>74.1 ± 0.5 **</td>
<td>74.4 ± 0.5 **</td>
<td>69.9 ± 0.7 **</td>
<td>69.5 ± 0.4 **</td>
<td>71.1 ± 0.2 **</td>
<td>71.9 ± 0.4 **</td>
</tr>
<tr>
<td>CI</td>
<td>0.84 ± 0.00</td>
<td>0.67 ± 0.00</td>
<td>0.83 ± 0.00</td>
<td>0.86 ± 0.01</td>
<td>0.69 ± 0.01</td>
<td>0.67 ± 0.01</td>
<td>0.77 ± 0.01</td>
<td>0.83 ± 0.03</td>
</tr>
<tr>
<td>MCI</td>
<td>0.96 ± 0.01</td>
<td>0.76 ± 0.01</td>
<td>0.94 ± 0.01</td>
<td>0.97 ± 0.01</td>
<td>0.79 ± 0.01</td>
<td>0.77 ± 0.01</td>
<td>0.87 ± 0.02</td>
<td>0.93 ± 0.03</td>
</tr>
<tr>
<td>Ethanol (% vol)</td>
<td>12.2 ± 0.1</td>
<td>13.6 ± 0.4</td>
<td>11.9 ± 0.2</td>
<td>12.4 ± 0.3</td>
<td>13.7 ± 0.1</td>
<td>13.6 ± 0.2</td>
<td>12.1 ± 0.1</td>
<td>12.3 ± 0.1</td>
</tr>
<tr>
<td>AT (g/L)</td>
<td>3.18 ± 0.02</td>
<td>2.69 ± 0.01</td>
<td>3.23 ± 0.02</td>
<td>3.24 ± 0.01</td>
<td>2.66 ± 0.00</td>
<td>2.68 ± 0.05</td>
<td>3.20 ± 0.02</td>
<td>3.20 ± 0.02</td>
</tr>
<tr>
<td>AV (g/L)</td>
<td>0.22 ± 0.02</td>
<td>0.22 ± 0.02</td>
<td>0.29 ± 0.02</td>
<td>0.29 ± 0.02</td>
<td>0.26 ± 0.00</td>
<td>0.26 ± 0.02</td>
<td>0.29 ± 0.01</td>
<td>0.29 ± 0.01</td>
</tr>
<tr>
<td>pH</td>
<td>3.6 ± 0.0</td>
<td>3.7 ± 0.0</td>
<td>3.5 ± 0.1</td>
<td>3.5 ± 0.1</td>
<td>3.7 ± 0.0</td>
<td>3.7 ± 0.0</td>
<td>3.5 ± 0.0</td>
<td>3.5 ± 0.0</td>
</tr>
<tr>
<td>Free SO(_2) (mg/L)</td>
<td>34.0 ± 0.6</td>
<td>29.0 ± 1.0</td>
<td>30.3 ± 0.92</td>
<td>29.0 ± 1.1</td>
<td>24.5 ± 0.7</td>
<td>29.3 ± 1.2</td>
<td>32.0 ± 0.0</td>
<td>28.7 ± 0.6</td>
</tr>
<tr>
<td>Brettanomyces (CFU/mL)</td>
<td>&lt;DL</td>
<td>&lt;DL</td>
<td>&lt;DL</td>
<td>&lt;DL</td>
<td>&lt;DL</td>
<td>&lt;DL</td>
<td>&lt;DL</td>
<td>&lt;DL</td>
</tr>
</tbody>
</table>

<DL: detection limit (10 CFU/mL); (**) significant at \( p < 0.01 \).

**Figure 3.** Total ellagitannins concentration of red wine aging for 2, 4, and 8 months in barrels of 1, 2, or 3 years cleaned with HPU \(_{hw}\) or steam. (*) significant at \( p < 0.05 \); (**) significant at \( p < 0.01 \).

These findings are in line with previous studies, confirming that the treatment induced no surface modification [13]. Furthermore, we can note that in the case of 2-year barrels, similar results were obtained in terms of TPI, CI, and MCI for HPU and steam treatment for Estate B.
3.3. Oak Wood Volatile Compounds

Oak barrels contain a high concentration of volatile compounds that positively impact a wine’s aroma. The main volatile compounds susceptible to desorb from oak wood to wine are the cis and trans-whisky lactones; volatile phenols, such as eugenol, guaiacol, furfural, and its derived compounds; and phenolic aldehydes, such as vanillin and syringaldehyde.

In our case, the results show that there are some significant differences between the volatiles from the oak wood composition of wines aged in barrels treated with HPU\textsubscript{hw} compared to those treated with steam (Figure 4). Indeed, the concentration of furfural was higher in the case of HPU\textsubscript{hw} treatment at 8 months of aging (barrels of 1 year) and 12 months of aging (barrels of 2 and 3 years), with the concentration increasing between 18.8 and 92.6% compared to steam. For the 5-methylfurfural, the concentration was significantly higher for HPU\textsubscript{hw} treatment for barrels of 1 year (12 months of aging) and 2 years (2, 8, and 12 months of aging) with an increase between 20.5 and 97%. Regarding the whisky lactone diastereoisomers, the trans-whisky lactone was significantly higher for barrels of 3 years treated with HPU\textsubscript{hw} after 12 months of aging with a concentration at 75.2 ± 5.6 µg/L, resulting in an increase of 46.9% compared to steam. This appears to also be a trend for barrels of 1 and 2 years. On the other hand, for cis-whisky lactone, there was no significant difference between steam and HPU\textsubscript{hw} treatments.

For syringaldehyde, the concentration was significantly higher for barrels of 1 and 2 years (12 months of aging) treated with HPU\textsubscript{hw}, with an increase of between 32.9 and 194.9% compared to steam. For barrels of 3 years (8 months of aging), we noticed a slight decrease in syringaldehyde concentration in the case of HPU\textsubscript{hw} treatment (8.7%) compared to steam. Finally, the concentration of vanillin was higher for barrels of 2 years treated with HPU\textsubscript{hw} after 8 and 12 months of aging, with an increase of 42.2 and 34%, respectively. There were no significant differences for eugenol, syringol, iso eugenol, 4-Allylsyringol, acetovanillione, acetosyringone, and guaiacol compounds.

Nevertheless, we noted that for all significant differences in volatile compounds, the concentrations were below the detection thresholds reported in Table 2. However, despite their high detection thresholds, they can contribute to the ‘toasty’ notes of wines and also increase the overall intensity of the woody character [18]. Moreover, Cameleyre et al. (2019) recently described the indirect impact of some volatile oak compounds at infra-threshold levels on the perception of panels [19]. Omission tests revealed the important role of vanillin and coniferaldehyde as, when omitted alone or with other compounds from the same family, their absence was significantly perceived, even under the detection threshold. The results are compelling since these compounds are widely sought by winemakers and can improve the organoleptic properties of wine. Increasing the extraction of these positive compounds by HPU\textsubscript{hw} treatment is very interesting and may be explained by the preservation of the wood structure following treatment [13].

3.4. Sensory Evaluation of Wines

Spider web diagrams obtained from average values of olfactory and gustative descriptors from a sensory analysis of wines after 12 months of aging for each modality are presented in Figure 5. All of the treated wines were characterized, with no significant difference (\( p < 0.05 \)) found for any of the olfactory descriptors (aromatic clarity and intensity, complexity, roasted/woody, vanilla and plank). The olfactory profile of all the wines was very similar. The focus on the roasted/woody descriptor revealed no significant difference, even though some volatile compounds from wood were extracted to a greater degree in the case of HPU\textsubscript{hw} treatment. Increased extraction of furfural and 5-methylfurfural as well as syringaldehyde by HPU\textsubscript{hw} was not detected by the panel, as they were below the detection threshold (Table 2). We noted the same trend in the case of the vanilla descriptor, despite finding a significant increase in vanillin for barrels of 2 years. The concentration differences of volatile compounds from wood between HPU\textsubscript{hw} and steam were insufficient to be detected by the panel.
Figure 4. Oak wood volatile composition of red wines aging for 2, 4, 8, and 12 months in barrels of 1, 2, or 3 years cleaned with HPUhw or steam. (*) significant at \( p < 0.05; (**) \) significant at \( p < 0.01; (***) \) significant at \( p < 0.001 \).

Table 2. Volatile compounds present in oak wood and their detection threshold [19,26–29].

<table>
<thead>
<tr>
<th>Family</th>
<th>Compound</th>
<th>Odorants</th>
<th>Detection Threshold</th>
</tr>
</thead>
<tbody>
<tr>
<td>Lactones</td>
<td>trans-whisky lactones</td>
<td>Coconut</td>
<td>130 µg/L (MS)</td>
</tr>
<tr>
<td></td>
<td>cis-whisky lactones</td>
<td>Coconut</td>
<td>20 µg/L (MS)</td>
</tr>
<tr>
<td>Furanic Aldehyde</td>
<td>Furfural</td>
<td>Toasty, caramel</td>
<td>15 mg/L (MS) 65 mg/L (W)</td>
</tr>
<tr>
<td></td>
<td>5-methylfurfural</td>
<td>Toasty, bitter almond</td>
<td>16 mg/L (W) 52 mg/L (W)</td>
</tr>
<tr>
<td>Phenols</td>
<td>Guaiacol</td>
<td>Smoky</td>
<td>9.5 µg/L (MS) 95 µg/L (W)</td>
</tr>
<tr>
<td></td>
<td>Eugenol</td>
<td>Clove</td>
<td>5 µg/L (MS)</td>
</tr>
<tr>
<td></td>
<td>Isoeugenol</td>
<td>Clove, spicy, woody</td>
<td>6 µg/L (MS)</td>
</tr>
<tr>
<td></td>
<td>Syringol</td>
<td>Smoky, spicy</td>
<td>/</td>
</tr>
<tr>
<td></td>
<td>4-allylsyringol</td>
<td>/</td>
<td>/</td>
</tr>
<tr>
<td>Phenolic Aldehydes and</td>
<td>Vanillin</td>
<td>Vanilla</td>
<td>1.1 mg/L (MS)</td>
</tr>
<tr>
<td>Ketones</td>
<td>Acetovanilline</td>
<td>Vanilla</td>
<td>1 mg/L (MS)</td>
</tr>
<tr>
<td></td>
<td>Syringaldehyde</td>
<td>Woody</td>
<td>5 mg/L (MS)</td>
</tr>
<tr>
<td></td>
<td>Acetosyringone</td>
<td>/</td>
<td>/</td>
</tr>
</tbody>
</table>

(MS): measured in a model solution (alcoholic solution, 12%; v/v); (W): measured in wine.
In addition, for gustatory attributes, we only noted a significant increase ($p < 0.05$) in astringency for barrels of one year treated with steam. The other attributes were not impacted. Thus, it appears that the HPU$_{\text{hw}}$ and steam treatments have a similar impact on the organoleptic properties of wine.

**Figure 5.** Sensory evaluation (olfactory and gustatory) of red wine aging for 12 months in barrels of 1 $(\text{a,b})$, 2 $(\text{c,d})$, or 3 years old $(\text{e,f})$ cleaned with HPU$_{\text{hw}}$ or steam. (*) significant at $p < 0.05$.

### 4. Conclusions and Perspectives

The study investigated the impact of using high-power ultrasounds associated with hot water or steam treatment on barrels containing a red wine aging in a barrel of different years with respect to (i) the chemical composition of the wine, (ii) the extraction of volatile compounds from oak, and (iii) ellagitannins over the course of one year, evaluated by sensory analysis (olfactory and gustatory).

In winemaking, the traditional steam treatment is widely acknowledged to provide complete sanitation, even though removing tartrate deposits is not complete at the porous surface. Application of HPU with hot water (60 °C) ensures complete removal of tartrate deposits and can also remove microorganisms in the depths of the wood. In this study on whole oak barrels, specific analyses proved that these treatments do not change the accessibility or diffusivity of phenolic compounds.

Few effects were observed on oak volatile compounds extraction in wine stored in HPU$_{\text{hw}}$-treated barrels. The concentration of furfural/5-methylfurfural, trans-whisky lactone, syringaldehyde, and vanillin could be extracted to a greater degree in the case of HPU$_{\text{hw}}$ treatment for barrels of 2 and 3 years. In the case of a younger barrel (one
year), syringaldehyde could also be extracted to a greater degree as well as furfural and 5-methylfurfural. However, differences between the steam and the HPU_{hw} wines were not significant in the sensorial analyses and could be explained by the high detection threshold of volatile compounds from wood. Further investigation on various wines, especially white wines, is required to complete these initial findings.

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