

Methyl salicylate, a grape and wine chemical marker and sensory contributor in wines elaborated from grapes affected or not by cryptogamic diseases

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| 1 | Methyl salicylate, a grape and wine chemical marker and sensory |
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| 2 | contributor in wines elaborated from grapes affected or not by cryptogamic |
| 3 | diseases |
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28

29 ABSTRACT.

30 Methyl salicylate (MeSA) is a plant metabolite that induces plant defence resistance and an 31 odorous volatile compound presenting green nuances. This volatile compound was shown to 32 be present in wine samples, sometimes at concentrations above its olfactory detection 33 threshold. MeSA is localized in grapes, particularly in the skins and stems, and is extracted 34 during red wine vinification. It was detected at the highest concentrations in wines of several 35 grape varieties, made from grapes affected by cryptogamic diseases, namely downy mildew 36 caused by *Plasmopara viticola*, and black rot caused by *Guignardia bidwellii*. It has also been 37 detected in wines from vines affected by Esca, a Grapevine Trunk Disease. MeSA can also be 38 considered to be a chemical marker in grapes and wine indicative of the level of development 39 of several vine cryptogamic diseases.

40 KEYWORDS. Methyl salicylate, Green aromas, Red wines, Stem, Cryptogamic diseases,
41 Downy mildew, Grape black rot, Grapevine Trunk Disease (Esca)

43 **1. Introduction**

44 Methyl salicylate (MeSA) is a volatile odorous compound presenting green and mintlike flavour nuances. This compound is a secondary plant metabolite, and the main 45 46 constituent of essentials oils from the shrub genus Gaultheria, in particular, the oil of Gaultheria procumbens (98 %) (Gurung, 2007), which is known as "wintergreen". Its fragrant 47 48 properties are appreciated and commonly used as an aroma in chewing gum, sweets and 49 dental products. It is also known to possess analgesic and antipyretic properties (Chan, 1996). 50 Methyl salicylate is widespread in plants and vegetables. For example, previous studies have 51 shown it to be an odorous compound associated with green pepper and tomato aromas and 52 have demonstrated that it is synthesized from salicylic acid by salicylate methyl transferase 53 (Buttery et al., 1969; Buttery et al., 1990; Tieman et al., 2010).

54 Its involvement in plant defence phenomena has been highlighted. MeSA is an 55 herbivore-induced plant volatile (HIPV) (Gadino et al., 2012). This compound is released by 56 a number of plants (e.g., beans, hops, tomatoes, cucumbers, peppers) in response to herbivore 57 attack, as in the case of grapevines with leaves affected by spider mite (*Tetranychus urticae*) 58 infestation (Van Den Boom et al., 2004). The action of MeSA was shown to have (i) a direct 59 insecticidal effect, for example on spotted-wing drosophila (Drosophila suzukii) (Kim et al., 60 2016) and (ii) an indirect attracting effect for several natural predators of spider mites on hops (Chrysopa nigricornis, Hemerobius sp., Deraeocoris brevis, Stethorus punctum) (James & 61 62 Price, 2004). Similar effects were observed with natural enemies of aphids on soybeans, such 63 as seven-spotted ladybugs (Coccinella septempunctata) (Mallinger et al., 2011; Zhu & Park, 64 2005). In addition, MeSA plays a key role in the induction of systemic acquired resistance 65 (SAR) in plants (Tang et al., 2015). In particular, it was established that MeSA was 66 synthesized from salicylic acid by tobacco plants infected with tobacco mosaic virus (Park et 67 al., 2007). Its release into the air constitutes a volatile defence signal that activates the resistance of neighbouring tobacco plants (Shulaev et al., 1997). Karban et al. (2014), who combined the results of 48 studies, confirmed the existence of "chemical communication" between plants. MeSA is therefore an indicator of diseased plants (Jansen et al. 2015) and has been evidenced as biomarker of grapevine leaves infected with downy mildew caused by *Plasmopara viticola* (Chalal et al., 2015).

73 In grapes and wine, MeSA was initially identified as an odorous constituent of several 74 marc distillates from Muscat cultivar (marc distillate of White Muscat from Piedmont, 75 Catalan roxo cultivar marc distillate) and spirits (Cognac and Calva distillate) (Di Stefano, 76 1986, Versini et al., 1995; Ledauphin et al., 2004). It was also tentatively identified and 77 assayed in wines elaborated from grapes undergoing carbonic maceration (Dell'Oro & Di 78 Stefano, 1990), where its quantitative levels depended on the type of grape variety. It was 79 later assayed in white wines from Vitis vinifera Emir variety (Cabaroglu et al. 1997). It was 80 mentioned as a wine component based on various analytical approaches, including 81 Comprehensive GC (GCxGC) analysis (Robinson et al., 2011; Bordiga et al., 2014; Versini et 82 al. 2005; Carlin et al. 2019a). It was also analysed at higher concentrations in experimental 83 Cabernet Sauvignon wines elaborated with high proportions of petiole at harvest (Ward et al. 84 2015) and in some white Italian grape varieties (Verdicchio and Trebbiano di Lugana) (Carlin 85 et al. 2019a). In grapes and wine, MeSA was also observed in several varieties under a 86 glycosylated form, liable to be released by chemical and enzymatic means (Williams et al. 87 1989; Versini et al. 2005; Esti & Tamborra 2006; Ghaste et al. 2015; Carlin, 2019ab).

Based on previous research, the present study aimed to better characterize methyl salicylate in wines, study its range of concentrations, its sensory impact on wine aromas, its content during vinification, particularly in relation to the development of fungal diseases on *Vitis vinifera* grapevines (i.e., downy mildew, black rot, and Esca, a Grapevine Trunk Disease). 93

94 **2. Materials and Methods**

95 2.1. Chemicals

96 Dichloromethane (99.9%) was supplied by VWR Chemicals (Fontenay-sous-Bois, 97 France). Sodium chloride (NaCl) was supplied by Supelco (Bellefonte, PA, USA). Ultrapure 98 water was obtained from a Milli-Q Plus water system (Millipore, Saint-Quentin-en-Yvelines, 99 France). Volatiles chemicals [methyl salicylate (analytical purity, $\geq 99\%$), ethyl salicylate 100 (99%), 3-octanol (\geq 99%), 1,8-cineole (\geq 99%)], alcanes (C8–C20) in solution and 101 compounds for fermentative media as cited by Marullo et al. (2006) were provided, all in 102 reagant or ACS grade, by Sigma-Aldrich (Saint-Quentin-Fallavier, France). Volatiles stock 103 blended solutions of 100 mg/L were prepared in HPLC-grade absolute ethanol (99.9%) 104 (Merck, Fontenay-sous-Bois, France) and stored at + 4 °C.

105

106 2.2. Wines samples

French red wines (52 samples) from several origins (Bordeaux, Burgundy, Loire and
Rhône Valleys, South West area) and vintages (from 2002 to 2014) were used in this study.
Other samples (46) were from various Bordeaux vineyards (Table S1.).

110

111 2.3. Grapes sampling

112 2.3.1. Sampling of Merlot and Cabernet Sauvignon diseased and non-diseased grapes

In 2012, a Cabernet-Sauvignon vine experimental plot from Bordeaux area was selected as previously mentioned (Pons et al., 2018). In the plot, 40 vines received fungicide treatment (10 times to control the main diseases as downy mildew, powdery mildew and black rot) while 40 vines didn't at any time during the growing season. At optimum ripening stage, the grape bunches from each modality of the plot were harvested by hand, removingpossible foci of grey rot (*B. cinerea*).

119 In 2013 and 2014, 40 kg of Merlot grapes were harvested on 6 blocks of a Bordeaux 120 vineyard plot planted in 2011 at ISVV (Institut des Sciences de la Vigne et du Vin, Bordeaux) 121 within the ResIntBio experimental vineyard (low-input viticultural cropping system). The 122 visual estimates made it possible to place grape damage level by downy mildew (brown rot) 123 due to *Plasmopara viticola*. This damage was situated between 2.2 and 59.3% depending on 124 the block. In the same time, 40 kg of grapes from a downy mildew-resistance variety, Artaban 125 (reference IJ134) developed in the context of an INRAE project, were harvested on 3 126 vineyard blocks grown in the same place. These grapes were used for experimental 127 vinifications.

In 2014, on the same Merlot ISVV vine plot, various categories of grape berries were harvested on the same day: healthy berries, naturally shrivelled berries due to sun exposure, berries shrivelled under the action of fungal disease such as downy mildew due to *P. viticola* and grape black rot due to *Guignardia bidwellii*.

132 In 2015, healthy Cabernet Sauvignon grapes were sampled in the control vines (20 133 grape bunches harvested) and on the vines with strong foliar symptoms of grapevine trunk 134 disease (Esca) (20 grape bunches) at the optimal ripeness stage in the same Haut-Médoc, 135 Denomination of Appellation Origin vineyard plot (Bordeaux vineyard). In the laboratory, 136 from these selected grapes, 100 berries without pedicels were randomly sampled from these 137 grapes, frozen in liquid nitrogen, and introduced 2 min into a mechanical ball mill to be 138 ground into a fine powder. Grape stems were analysed according to the same protocol of 139 preparation. Each modality was performed in triplicate (n=3).

In 2016, 10 kg Cabernet Sauvignon grape bunches were harvested at optimal ripeness
stage in Côtes-de-Bordeaux, Denomination of Appellation Origin, (Bordeaux vineyard) from

142 control vines, vine plants affected by grapevine trunk disease (Esca) with moderate leaf143 symptoms (level 1), and with strong leaf symptoms (level 2).

144 2.3.2. Sampling of Furmint_grapes from diseased grapevine or not

145 In 2017, Vitis vinifera Furmint B grapes were sampled at two harvest periods in Tokaj 146 region (Hungary). The first harvest of healthy grapes, without infection of B. cinerea, was 147 done at optimal ripeness stage for the elaboration of dry white wine. Two categories of 148 harvest were considered, one on apparently healthy vine plants and the other on vine plants 149 affected by strong leaf symptoms of grapevine trunk disease (Esca). The second harvest was 150 done 1 month and half later (for elaboration of sweet wine) with berries affected by noble rot 151 due to *B. cinerea* considering the same categories of grapes. For each harvest, a percentage of 152 Esca diseased grape bunches were used to elaborate the wine in comparison with the control 153 (from healthy vines).

154

155 2.4. Experimental winemaking conditions

156 2.4.1. Red winemaking modalities

157 In 2012, red wines were elaborated with Cabernet-Sauvignon grapes naturally infected 158 by P. viticola (brown rot) and increasingly supplemented to control grapes (grapes from the 159 fungicide treatment modality without brown rot) (Pons et al., 2018). During the experiments, 160 incorporation was carried out in % of withered berries per cluster for the 5 modalities (0, 2, 5, 161 10, 20 %). Each modality contained 60 kg of grapes vinified in 100 L stainless steel tanks 162 after grapes destemming and crushing. Standard winemaking procedures were followed 163 including alcoholic and malolactic fermentation with commercial yeast and bacterial strains 164 (Laffort Œnologie, Floirac, France). At the end of malolactic fermentation, red wines were 165 sulfited at 50 mg/L (6% v/v; Laffort, Bordeaux, France). Before bottling, they were fined and filtered. Then, wines were filled into 750 mL glass Bordeaux bottles and closed using 166

standard commercial practices. Sulfur dioxide was adjusted at 30 mg/L before bottling.
Bottles were kept in a dark and in a temperature controlled room (18 °C) until required for
analysis.

170 For vintages 2013, 2014, 2016, red wines were vinified at ISVV facility. All hand-171 harvested Merlot, Cabernet, Artaban grape bunches (20 kg each modality) were destemmed 172 mechanically (mechanical destemmer, Bellot, Gradignan, France), manually crushed and 173 vinified in stainless-steel tanks. Grape juices were inoculated with a commercial strain of 174 Saccharomyces cerevisiae (FX10; Biolaffort Enologie, Bordeaux, France) at 20 g/hL. In 175 order to reproduce the winemaking conditions prevalent in the Bordeaux region, maceration 176 lasted three weeks. During alcoholic fermentation (around 7 days), the cap was punched down 177 twice per day. After spontaneous malolactic fermentation in 3 L bottles maintained at 20°C, 178 wine were supplemented with 50 mg/L sulfur dioxide solution (6% v/v; Laffort, Bordeaux, 179 France), bottled and kept at 18°C until analysis.

180 2.4.2 White winemaking modalities

181 In 2017, dry and sweet Furmint white wines were vinified in Tokaj experimental 182 winery. Vinification modalities were organized at each harvesting stage with incorporation of 183 25%, 50% or even 75% of Furmint grape bunches from strongly Esca diseased vines. In each 184 situation, grapes were crushed and the juices were put in 5L bottles per modality with 3 185 replicates. Then, grape juices were inoculated with commercial Saccharomyces cerevisiae 186 yeast strains Uniferm 228 (Uniferm, Werne, Germany) for dry wine and Actiflore B0213 187 (Laffort Œnologie, Bordeaux, France) for sweet wine at 20 g/hL. The dry white wines were 188 sulfited at 70 mg/L at the end of alcoholic fermentation (less than 2 g/L residual sugars) then 189 kept in bottles at 18°C until analysis. The sweet wines were sulfited at 140 mg/L when 190 alcohol strength reached 12% vol. then kept in bottles until analysis.

192 2.5. Laboratory scale microvinifications with incorporation of diseased berries

With Merlot grape berries from 2014 vintage (healthy, naturally shrivelled berries due to sun exposure or shrivelled under the action downy mildew and grape black rot), 20 selected berries of each type were fermented in a 100 mL-bottle containing 60 mL of a model must solution with a composition described by Marullo et al., 2006. After alcoholic fermentation, lees were removed by centrifugation (5 min, 6000 r.p.m.), samples were sulfited (50 mg/L) and stored at 4 °C before analysis experiments. Each modality was performed in triplicate (n=3).

200

201 2.6. Sample preparation for Methyl salicylate (MeSA) evidence in wine

202 2.6.1. Wine extraction and semi-preparative HPLC

203 As described by Poitou et al. (2017), a 750 mL Cabernet wine sample was extracted 204 using 80, 50, and 50 mL dichloromethane for 10 min each with magnetic stirring (700 r.p.m.) 205 and separated in a funnel. The organic phases were collected, frozen overnight at -20°C to 206 remove the emulsion, and then dried over sodium sulfate and concentrated to around 20 mL 207 using a Buchi R-114 rotary evaporator (Buchi, Rungis, France). The extract was filtered using 208 a 0.45 µm membrane syringe and concentrated under a nitrogen flow (100 mL/min) in a 209 graduated glass tube (Jean Premont, Bordeaux) to obtain 500 µL wine extract. Then, the 210 extract was injected on to Reverse-phase (RP) HPLC system using water and ethanol as 211 solvents. the Ultimate 3000 semipreparative HPLC system (Dionex, Courtaboeuf, France) and 212 a Novapak C18 column (300×7.8 mm internal diameter (i.d.), 6 μ m, Waters, Saint Quentin, 213 France) with a guard column of the same phase.

Chromatographic conditions included a 250 µL injection volume with a flow rate of 1
mL/min. The linear program gradient was as follows: phase A, water; phase B, ethanol; 0–2

216 min, 100 % A, 0–50 min, linearly programmed until 100 %. Fifty fractions, containing each 1
217 mL effluent, were collected and evaluated for their sensory properties.

Selected fractions were mixed with the next or previous fraction on the basis of their odour to form a 2 mL fraction group. Each was then diluted with ultrapure water to obtain 12 % ethanol (v/v) and then re-extracted three times with 10 % (v/v), 5 % (v/v), and 5 % (v/v) dichloromethane, respectively, for 10 min each time. The organic phases were combined and concentrated to 100 µL under nitrogen flow before analysis.

223

224 2.6.2. Identification of MeSA by Heart-cut Gas Chromatography coupled to Olfactometry and
225 Mass Spectrometry (MDGC-O-MS)

Heart-Cut Multidimensional gas chromatography separation was performed on two Agilent 7890 GC (Agilent Technologies, Palo Alto, CA, USA), connected via a heated transfer line at 230°C as previously mentionned (Poitou et al. 2017).

229

230 2.7. Wine sample treatments for studying MeSA release from bound forms

Red wine samples were acidified at pH 3.0 with addition of 5M H_2SO_4 and aliquoted in 20-mL headspace amber vials before closure with a PTFE-faced silicone septum/aluminum crimp cap. They were then kept in the dark at 60°C prior to analysis or at 20°C for control.

234

235 2.8. Quantitation of MeSA, ethyl salicylate (EtSA) and 1,8-cineole.

236 Concentrations of MeSA, EtSA and 1,8-cineole were determined by solid-phase 237 microextraction coupled to gas chromatography–mass spectrometry (SPME-GC-MS). MeSA 238 concentrations were also determined in comparison, by solid-phase-extraction coupled to gas 239 chromatography–mass spectrometry (SPE-GC-MS) analysis on selected samples. For each 240 modality, analyses were performed using a Combi PAL sampler (CTC Analytics, Zwingen, Switzerland) on an Agilent 6890N gas chromatograph (Agilent, Palo Alto, CA, USA),
coupled to an Agilent HP 5973N mass spectrometer (electron impact mode at 70 eV).

243

244 2.8.1. Quantitation of MeSA, EtSA and 1,8-cineole by SPME procedure

A 7 mL sample diluted in deionized water containing 5 mL grape juice, wine or 3 g grape-berry powder was transferred into a 20-mL headspace amber vial containing 3 g sodium chloride (NaCl). A 10 μ L sample of internal standard solution 3-octanol at 100 mg/L in ethanol was supplemented before closure with a PTFE-faced silicone septum/aluminum crimp cap and homogenized manually.

250 Α 2 cm, 50/30 μm Divinylbenzene-Carboxen-Polydimethylsiloxane 251 (DVB/CAR/PDMS) SPME fiber purchased from Supelco Inc. (Bellefonte, PA, USA) was 252 used. The sample was pre-incubated for 5 min at 40°C. Adsorption lasted 30 min, at the same 253 temperature, with stirring at 500 r.p.m. (3 seconds on, 2 seconds off). Then, desorption took 254 place in the injector in splitless mode (3 min) at 240 °C for a duration of 10 min, with a purge 255 flow rate of 50 mL/min. The fiber was then reconditioned for 10 min at 250 °C.

256

257 2.8.2. Quantitation of MeSA by SPE procedure

258 The isolation of MeSA from wine was done by its percolation on cartridge sorbents 259 using an automatic liquid handler GX-274 ASPEC (Gilson, Middleton, WI). Initially, the 260 cartridge sorbents [SPE Chromabond HRX, 500 mg sorbent, 6 mL cartridge volume from 261 Macherey-Nagel (Bethlehem, PA, USA)] were conditioned with 7 mL of methanol at a 6 262 mL/min flow, followed by a 3 mL volume of ultrapure water/ethanol (90:10, v/v) mix at the 263 same flow. Then, to a 20 mL of wine sample were supplemented 50µL of 3-octanol at 100 264 mg/L in ethanol. 19640 μ L of this mixed wine sample were percolated on the cartridge at a 265 flow of 3 mL/min. After loading, the cartridge was washed with 2 mL of ultrapure water, then 266 compounds were eluted with 3 mL of dichloromethane/pentane (50:50, v/v) and 3 mL of 267 dichloromethane/methanol (95:5, v/v). Percolation speed was fixed at 10 mL/min. The 268 organic phases were combined, dried with sodium sulfate, concentrated under nitrogen flow 269 (at approximately 100 mL/min) to 150 μ L and kept -20 °C before analysis.

270

271 2.8.3. GC-MS analysis.

272 The carrier gas was Helium N60 (Air Liquide) with a flow rate of 1 mL/min. A 273 Carbowax 20 M type fused capillary column was used: BP20, 50 m, 0.25 mm i.d., 0.22 µm 274 film thickness (SGE, Ringwood, Australia). Temperature program was as follows: 45 °C for 5 275 min, increasing temperature by 3 °C/min to 180 °C then 20 °C/min to 240 °C with an 276 isotherm at the final temperature for 10 min. The mass spectrometer, operating in electron 277 impact (EI) mode (70 eV), was connected to the GC with a heated transfer line at 230 °C. The 278 compounds were quantitated using selected ion monitoring mode (SIM) on MSD Chemstation 279 software (v B.04.03) from Agilent. Selected ions for internal standards were m/z 83, 59 (3-280 octanol), and m/z 83 for quantitation. MeSA was detected using m/z 152, 120, 92 ions, and 281 quantitated using m/z 152 ion. EtSA was detected using m/z 166, 120, 92 ions, and quantitated 282 using m/z 120 ion. 1,8-cineole was detected using m/z 154, 139, 111, 108 ions, and 283 quantitated using m/z 108 ion.

284

285 2.8.4. Method validation for MeSA, EtSA and 1,8-cineole quantitative analysis

286 2.8.4.1. SPME-GC-MS analysis

Linearity (n=3), estimated by a spike at 6 increasing concentration levels of MeSA and EtSA ($5/10/25/50/100/200 \mu g/L$), were 0.998 and 0.993 respectively. *Precision* (n=5) estimated as the relative standard deviation (% RSD) were 2.87 % and 5.84% respectively. The *recovery* values (% RSD) estimated by adding 30 µg/L to a reference wine were 104.5 % and 98.7 % respectively. *Limit of Quantitation (LOQ, S/N 10)* were estimated at 1.05 μ g/L and 0.41 μ g/L for MeSA and EtSA respectively. *Limit of Detection (LOD, S/N 3)* were estimated at 0.35 μ g/L and 0.13 μ g/L for MeSA and EtSA respectively. Method validation for 1,8-cineole quantitative analysis was described elsewhere (Poitou et al. 2017).

Validation for berry analysis was performed in deionized water following the same procedure. Values were equivalent for the various parameters calculated. Results were reported on a per weight basis in micrograms per kilogram of fresh weight for berries and per litre for wines.

299 2.8.4.2. SPE-GC-MS analysis

Linearity (n=3) estimated by a spike at 5 increasing concentration levels of MeSA $(1,25/2,5/5/10/20 \ \mu g/L)$ was 0.997. *Precision* (n=3) estimated as the relative standard 302 deviation (% RSD) was 3.44 %. *LOQ* and *LOD* were estimated respectivley at 0.03 $\mu g/L$ and $0.01 \ \mu g/L$.

2.8.4.3. Comparative accuracy of methyl salicylate assay by SPE-GC-MS and SPME-GC-MS
The determination of methyl salicylate in wines by the SPE GC-MS method and by the SPME
GC-MS method were compared on the accuracy parameter using the protocol defined in the
Compendium of international methods of wine and must analysis (2020), which refers to ISO
13528 (2015).

309

310 2.9. Sensory Analysis

311 2.9.1. General conditions.

Sensory analyses were conducted by 2 panels from the research unit in Enology (enologists, researchers, or students) with a good experience of wine tasting (usual tasting wine several times a week). The panelists (n = 27 for first session, n=18 for second and third sessions) were aged between 22 and 56 years old. 77% and 61% respectively, depending on 316 the session, were having less than 35 years old (average age 29 years old for the first session, 317 and 33 years old for the second and third sessions). The percentage of female panelists was 67 318 % and 72 % respectively. Orthonasal sensory evaluations took place in a temperature-319 controlled room (ISO 8589:2007) maintained at $20 \pm 1^{\circ}$ C, equipped with individual boxes. 320 Samples were presented in random order, in black tasting glasses, coded by three-digit 321 numbers, and covered with plastic caps. All samples were exclusively smelled by orthonasal 322 evaluation. Generally, sensory evaluations were conducted during a unique session of 3 h 323 with a renewal of the samples at mid-session. In the context of the COVID-19 epidemic 324 (second and third sessions), the olfactory detection thresholds were conducted all a day, by 325 changing the glasses and refilling them, after each experimenter had reached twice the 326 olfactory detection threshold, with a time volume of 15 minutes between each. Between each 327 repetition, the order of the glasses and the coding were changed.

328

329 2.9.2. Determination of MeSA olfactory detection threshold.

330 Olfactory detection threshold was determined in model wine [L(+)-tartaric acid 5 g/L, 331 12 % vol., pH 3.5]. Samples were presented as a series of triangular tests with ascending 332 order of methyl salicylate concentration: 11.8, 17.7, 26.6, 40, 60, 90, 135, 202.5 µg/L 333 respectively. Value was calculated using the BET (Best Estimate Threshold) method (NF ISO 334 13301: 2002). For each panelist, the best estimate threshold was the geometric mean of the 335 highest concentration missed and the next higher concentration tested. BET for the group is 336 the result of the geometric mean of the individual BETs. The odor active value (OAV) which 337 represents the measured concentration of a volatile present in a sample, divided by its 338 measured sensory detection threshold in a similar matrix, was also determined on selected 339 wine samples (Patton and Josephson, 1957).

341 2.9.3. Evaluation by a free vocabulary task.

342 26 panelists from the first panel (17 women, 9 men; average age 28 years old) 343 expressed their olfactory perception with their own terms during a single session. A unique 344 modality containing 200 μ g/L of methyl salicylate, was evaluated in a model wine [L(+)-345 tartaric acid 5 g/L, 12 % vol., pH 3.5] and in a non-barrel-aged, commercial Bordeaux wine. 346 Terms with only one citation or non-adapted (hedonic, specific of taste) were not considered. 347 Several were also considered together on the basis of their analogy.

348

349 2.10 Statistical analysis.

Analysis of variance (ANOVA) was performed on normalized scores (judges), homogeneity of variance was evaluated with Levene's test, and the normality of residuals was assessed using the Shapiro–Wilk test. Statistical significance was set at 5 % (p <0.05). Statistics were calculated using R software (v 3.1.1) except Spearman correlation test with XLStat-Premium software (Addinsoft).

355

356 3. Results and discussion

357 3.1. Methyl salicylate as an odorous volatile compound in experimental and commercial red
358 wines

In a past publication, while performing GC-O analysis of an HPLC fraction from an extract of an unripe Bordeaux Cabernet-Sauvignon experimental wine, several odorous zones were detected that corresponded to several impact volatile compounds, such as 2-methoxy-3alkylpyrazines, 1,8-cineole, 1-(2,3,6-trimethylphenyl)buta-1,3-diene, and several C6, C8, C9 saturated and unsaturated aldehydes (Poitou et al. 2017). When this approach was pursued using other extracts, an odorous zone marked by fresh, camphor and medicinal notes was detected, with linear retention indices of 1775 and 1202 on BP20 and BP1 respectively, 366 corresponded to the retention time of methyl salicylate (MeSA). Identification was confirmed
367 by mass spectrum by comparison to NIST library and by injection of the pure commercial
368 standard.

MeSA was then quantified by SPME-GC-MS method, in a series of 98 French red wines, including 52 commercial wines from different wine-producing regions (Bordeaux, Loire and Rhône valleys, Burgundy, etc.). The remaining wines were experimental wines made from various clones of Cabernet Franc and Merlot grapes. The numerical quantitative data was graphically depicted by a boxplot (**Fig. 1**).

374 The quantitative results showed that the methyl salicylate concentrations in the 375 analysed wines mainly ranged between 5 and 25 µg/L, with about 15% of the wines ranging 376 between 30 and 40 μ g/L. The mean and the median of the 98 wines were 18.7 μ g/L and 13.6 377 µg/L, respectively, which is similar to the values observed by Carlin et al. (2019a) in 378 Verdicchio wines. However, proportionally high levels were observed in some wines (n = 5), 379 with concentrations ranging from 69.2 to 131.8 µg/L (Table 1). These 5 wines corresponded 380 to 3 commercial wines from the 2014 vintage (Côtes-du-Rhône, Irouleguy and Bordeaux 381 Denominations of Appellation Origin) and two experimental Merlot wines obtained from 382 different soils (gravel and sand) from the 2006 vintage. The accuracy of the MeSA 383 quantitative measurements led by SPME-GC-MS was confirmed by assays done on selected 384 wine samples using an alternative method by SPE-GC-MS (Table S2.).

Analysis of ethyl salicylate (EtSA), the structural analogue with a similar odour, revealed its presence in these wines, but at much lower concentration levels, ranging from 0.3 to 9.8 μ g/L with a mean of 0.91 μ g/L. Furthermore, 1,8-cineole, another odorous marker of green eucalyptus notes in red wines, presented concentrations that did not exceed 2.24 μ g/L, which were not related to those of MeSA and EtSA (**Table 1**). Concerning the assay of MeSA in wines from Cabernet Franc clones grown on the same plot, the results showed variability

between clones from the same vintage (10-40%), and even greater variability in different
vintages, with variations in average concentrations ranging from 1.5 to 4.8 fold (Fig. S1.).

393

394 3.2. Sensory impact of methyl salicylate in red wines evaluated through psychophysical tests
395 and sensory reconstitution

396 The olfactory detection threshold of methyl salicylate was determined three times in a 397 model wine solution. Value estimation (BET method) revealed an olfactory detection 398 threshold of 75 \pm 25 µg/L in the model wine solution. For each olfactory session, the 399 minimum detection threshold value by certain panel members was 9.6 µg/L. These values 400 indicated that MeSA could have a sensory contribution at the highest concentration levels 401 assayed in red wines (Table 1). Synergistic effects were observed between MeSA and 402 guaiacol, an odorous oak wood volatile compound (Yodder et al. 2012). Furthermore, MeSA 403 could be involved in perceptual interaction phenomena with other wine volatile compounds.

In order to qualitatively characterize the odour change caused by an increase of MeSA concentration in the wine, a panel of 26 wine experts conducted analysis using a free vocabulary technique. An addition of 200 μ g/L of this molecule to red wine introduced an aromatic freshness expressed by an increase in the number of citations of the terms "fresh" or "fresh fruit". Specific fresh green nuances were also noted in the model wine as "chemical", "dentist", "pharmaceutical", "menthol", and "camphor" (**Table S3.**).

410

411 3.3. Location of MeSA in healthy grapes (berries, stem), must and extraction kinetics during
412 red wine vinification.

To determine the distribution of methyl salicylate in the different parts of the grape,
quantitation was performed after separating the berries and stems and on the pulp and skins.
Analysis carried out on Cabernet Sauvignon grapes harvested on healthy plants at optimal

416 ripeness (2015 vintage) revealed that concentrations of MeSA in stems (122 µg/kg fresh 417 weight) were significantly higher than concentrations in berries (0.5 µg/kg fresh weight). 418 These observations were consistent with those of Ward et al. (2015), which showed an 419 increase in methyl and ethyl salicylate concentrations, as well as several terpenes, with an 420 incorporation of petioles in the must. Further analysis revealed that methyl salicylate was 421 more abundant in the skins (68%) than in the pulp (32%). After harvesting, quantitation of 422 MeSA was performed on must from healthy Cabernet Sauvignon cultivar grapes. MeSA was 423 progressively extracted during alcoholic fermentation and post-fermentation maceration. The 424 assay of MeSA in the must therefore showed a gradual increase in its concentrations during 425 vinification (Fig. 2.).

426 The maximum concentration was reached at the end of fermentation, after approximately ten 427 days. Furthermore, since glycosylated forms of MeSA had already been observed in grapes 428 and wine (Williams et al. 1989; Esti et al. 2006; Ghaste et al. 2015; Carlin et al. 2019b), the 429 increase in MeSA observed during alcoholic fermentation may result from a combined effect 430 of extraction (skin) and release through the enzymatic action of yeast, as observed with 431 monoterpene glycosides (Bisotto et al., 2015). Moreover, an acidic treatment, at pH 3, 432 performed on several red wine samples stored at 20°C (control) and at 60°C led to increased MeSA concentrations in the wines (Table S4.). This observation supports the existence of 433 434 "bound" forms of this compound in these wines, presumably related to glycosides, as 435 previously mentioned.

436 Moreover, since MeSA levels appear to be much higher in the stems, red wine 437 vinification procedures, with and without destemming, were performed according to 438 Bordeaux red winemaking protocol, with 100% destemmed grapes, or the incorporation of 439 non-destemmed grapes (20% in volume). The results showed an increase in MeSA levels in

440 wines (82% of increase in free run wines) elaborated with non-destemmed grapes (Table
441 S5.).

442

3.4. Detection and quantitation of methyl salicylate in grapes, musts and wines from diseased
vines infected with cryptogamic fungi

445 Considering the possible origins of MeSA, as stated in the introduction, the hypothesis
446 of a defence reaction to infection by various grapevine pathogens (downy mildew; grapevine
447 trunk disease, ESCA) was naturally considered to interpret the high content measured in
448 several wines.

449 *3.4.1. Impact of downy mildew and other grape pathogens*

450 Two series of wines were considered for MeSA analysis. The first series was produced 451 from Merlot grapes affected by downy mildew (P. viticola), with estimated proportion 452 ranging from 2.2 to 59.3% from grape bunches with visual damage (10 wines). The second 453 series was produced from the newly created Vitis sp. variety Artaban, which is resistant to 454 downy mildew (4 wines). All of the grapes had been vinified under the same conditions. 455 MeSA content was assayed in these experimental wines in 2 vintages (2013-2014). MeSA 456 was therefore assayed in Merlot wines from grapes infected with downy mildew with 457 concentrations of 120.7 μ g/L, and a maximum value of 157.9 ± 6.7 μ g/L, while 458 concentrations were below 29.8 µg/L in wines from resistant varieties, without any downy 459 mildew symptoms (Fig. S2.).

The concentration of MeSA was the assayed in experimental Cabernet-Sauvignon wines from 2012 vintage, elaborated with the incorporation of various proportions of grapes infected by downy mildew. This analysis had previously been performed by Pons et al. (2018) for other volatile compounds (lactones, 2-methoxy-3-isobutylpyrazine, 3-methyl-2,4nonanedione) on samples kept in a wine cellar. In this case, while no significant differences were observed in the usual analysis between the modalities, a proportional increase of MeSA content was observed when berries infected with brown rot were incorporated (**Table 2**, **Table S6.**). The results showed MeSA content ranging from 8.46 μ g/L in wines obtained from must without any addition of diseased berries to 32.14 μ g/L in wines obtained with an incorporation of 20% diseased berries.

470 In addition, the impact of other grapevine pathogens on MeSA content in wines was 471 considered through fermentation in model must supplemented with selected berries of each 472 type: healthy, shrivelled by the sun, shrivelled by the effects of downy mildew (*Plasmopara* 473 viticola) and infected grape black rot (Guignardia bidwellii). The same number of Merlot 474 grape berries (20 berries), harvested in 2014, were incorporated into the model must and 475 fermented. It is important to note that, unlike berries affected by downy mildew, the solutions 476 obtained after the alcoholic fermentation and maceration of berries infected with grape black 477 rot were colourless. Analysis of the solutions revealed a slight increase in the concentrations 478 of MeSA linked to the shrivelling phenomenon in the berries. However, variations between 479 the samples remained low, particularly in comparison to those generated by the two grapevine 480 pathogens, mildew and grape black rot (Fig. 3.).

481 Therefore, all the experiments confirmed the relationship between MeSA content in 482 wine and downy mildew, as suggested in the vine V. vinifera (Chalal et al., 2015). Moreover, 483 it was demonstrated that other fungi, such as grape black rot, can also induce significantly 484 higher concentrations of this compound in berries and in wine. Furthermore, we observed that 485 fermentation of the model must solution with 1 mg/L of salicylic acid did not induce any 486 distinct increase in the concentrations of methyl salicylate (results not reported). This shows 487 that it cannot have originated from the metabolism of salicylic acid by yeast, which is 488 possibly present in greater quantities in the infected plants and grapes.

490 *3.4.2. Esca, grapevine trunk disease impact*

491 The impact of Esca (one Grapevine Trunk Disease) on methyl salicylate 492 concentrations in wines was also considered. For this purpose, Cabernet Sauvignon grapes 493 from healthy plants and Esca-affected plants with moderate or strong leaf symptoms were 494 harvested (2016 vintage) and vinified under the same conditions. Classical wine analysis 495 evidenced a slightly lower alcoholic strength of wines from Esca-affected vines, in 496 accordance with previous studies (Lorrain et al., 2012) (Table S7.). The quantitation of 497 MeSA was carried out on samples collected immediately after alcoholic fermentation was 498 complete, then one week later in the wines in contact with the grape pomace, and a year later 499 for the final wines. Quantitation of MeSA showed a significant increase of its concentrations 500 in wines linked to the Esca infected vine plants (**Table 3**). Moderate damage as expressed by 501 leaf symptoms was sufficient to double the MeSA concentrations compared to the control. 502 Severe Esca symptoms in the plant led to an even greater increase in concentrations compared 503 to the control. Quantitative results one year later in the aged wine confirmed the results 504 obtained immediately following the end of alcoholic fermentation. This experimentation 505 therefore confirmed the real influence of Esca grapevine disease on the increase of MeSA 506 content in red wines. However, under these conditions, the concentrations remained low 507 compared to the previously determined olfactory detection threshold of the compound. 508 Indeed, concentrations of MeSA in wines are also dependent on other factors such as those 509 stated above, and perhaps on the vine's ability to synthesize it. On the other hand, a 510 comparative analysis of MeSA content in berries and stems from Cabernet Sauvignon grapes 511 (Haut-Medoc, 2015), harvested from grapevines both severely affected and unaffected by 512 Esca, showed increases in MeSA concentrations in stems from diseased grapes that were 513 similar to those observed in the berries (Table S8.).

514 Another study conducted an analysis of Tokaj white wines made with increasing 515 proportions of grape from *Vitis vinifera* of the Furmint variety, harvested from Esca-infected 516 vine plants (25%, 50%, 75%) in comparison with the control. While the alcohol content was 517 slightly lower in the wines containing a higher proportion of Esca, the quantities of MeSA 518 were also higher in Esca wines than in the control dry white wine (Table S9.). However, for 519 the study carried out on the same vineyard with Furmint grapes affected by noble rot caused 520 by Botrytis cinerea for the elaboration of sweet wines, the level of MeSA were similar in all 521 modalities (Table S9.).

522

523 Conclusion

524 Methyl salicylate (MeSA), as a volatile odorous compound, can be present in red wines at 525 sometimes relatively high levels (70 to 130 μ g/L), that is, concentrations near or above its 526 olfactory detection threshold as determined in a model solution at 76.2 \pm 25.5 μ g/L. Based on 527 sensory experiences with supplementation at 200 µg/L, this compound could contribute to 528 strengthening the expression of fresh green aromatic nuances in red wines, e.g. 529 "pharmaceutical", "camphor" or "menthol" aromas. Furthermore, the significant concentration 530 of MeSA in stems provides evidence that their supplementation during red wine vinification 531 may increase the abundancy of this compound in wines. In addition, the relation between 532 MeSA and several vine pathogens (downy mildew, grape black rot) or grapevine trunk 533 disease (Esca) was demonstrated in grapes and in red and white wines. The vinification of 534 such diseased grapes may affect the wine aroma quality, particularly when stems are 535 incorporated. It appears that this change was induced by a host defence mechanism against 536 fungal infection. Therefore, methyl salicylate may constitute a good volatile indicator of the 537 vineyard's state of infection, revealing the physiological state of vine plants under fungal 538 pressure.

539

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543

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- 547

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| | Volatile compounds ^{<i>a</i>} | | | |
|--|--|-----------------------------|----------------------------|-----------------------|
| | Vintage | Methyl salicylate (µg/L) | Ethyl salicylate (µg/L) | 1,8-Cineole (µg/L) |
| Bordeaux (Merlot, on sand) | 2006 | 131.80 (19) | 0.79 (0.1) | 0.07 (0.01) |
| Côtes du Rhône (Shiraz) | 2014 | 94.04 (0.5) | 2.82 (0.01) | 0.2 (0.01) |
| Bordeaux (CS ^{<i>b</i>} , Merlot blend) | 2014 | 80.36 (6.0) | 7.61 (0.5) | 0.16 (0.02) |
| Bordeaux (Merlot, on gravels) | 2006 | 72.94 (3.3) | 0.74 (0.1) | 0.05 (0.01) |
| Bordeaux (hard press wine) | 2014 | 69.20 (1.0) | 4.76 (0.1) | 0.1 (0.01) |
| Irouleguy (Cabernet franc) | 2013 | 43.30 (0.1) | 1.94 (0.1) | 0.08 (0.01) |
| Bordeaux (Merlot, on clay) | 2006 | 38.71 (1.5) | 0.73 (0.2) | 0.15 (0.01) |
| Chinon (Cabernet franc) | 2014 | 38.62 (0.5) | 1.06 (0.1) | 0.17 (0.03) |
| Bordeaux (Merlot, on sand) | 2007 | 37.54 (0.5) | 1.01 (0.1) | 0.21 (0.02) |
| Bergerac (Merlot, organic wine) | 2013 | 37.15 (1.5) | 0.29 (0.1) | 0.23 (0.02) |
| Girondas (Grenache) | 2012 | 35.24 (2.5) | 1.36 (0.1) | 0.39 (0.01) |
| Chateauneuf du Pape (Grenache) | 2014 | 34.90 (4.0) | 1.61 (0.1) | 0.28 (0.01) |
| Fronton | 2007 | 34.50 (0.5) | 2.19 (0.1) | 0.85 (0.01) |
| Bordeaux (Merlot, on clay) | 2007 | 30.60 (0.5) | 1.01 (0.1) | 0.38 (0.01) |
| Valréas (Grenache, organic wine) | 2014 | 30.36 (1.5) | 0.98 (0.1) | 0.26 (0.01) |
| Saint Estèphe (CS b , Merlot blend |) 2014 | 26.32 (1.5) | 5.37 (0.3) | 0.13 (0.01) |
| Pauillac (CS ^b , Merlot blend) | 2014 | 25.80 (4.0) | 8.08 (2.0) | $1.04(0.1)^{c}$ |
| Pauillac (CS, Merlot blend) | 2009 | 25.67 (0.1) | 9.81 (0.1) | 2.24 (0.03) |
| Pauillac (CS, Merlot blend) | 2013 | 23.29 (4.5) | 0.63 (0.7) | 0.29 (0.05) |
| Chinon (Cabernet franc) | 2004 | 18.66 (0.6) | 0.51 (0.1) | 0.26 (0.04) |
| Margaux (CS, Merlot blend) | 2004 | 17.32 (0.1) | 1.10 (0.1) | 0.31 (0.01) |

Table 1. Methyl salicylate concentrations in various French red wines in comparison with ethyl salicylate and 1,8-cineole content (n=2)

^{*a*} For each compound, concentrations in bold corresponded to OAV (Odor Activity Value) >1. ^{*b*} CS : Cabernet Sauvignon ^{*c*} Poitou et al. (2017)

| | diseased berries $(\%)^{b}$ | | | | RC ^c | |
|-------------------|-----------------------------|---------|--------|--------|-----------------|------------------|
| | 0 | 2 | 5 | 10 | 20 | |
| Methyl salicylate | 8.46 | 19.39 | 24.36 | 29.42 | 32.14 | <i>p</i> = 0.017 |
| (µg/L) | (0.9) | (1.4) | (0.05) | (2.5) | (1.65) | |
| Ethyl salicylate | 1.42 | 1.36 | 1.13 | 1.39 | 1.11 | n.s. |
| (µg/L) | (0.18) | (0.15) | (0.04) | (0.3) | (0.06) | |
| 1,8-cineole | 0.10 | 0.09 | 0.10 | 0.09 | 0.10 | n.s. |
| (µg/L) | (0.005) | (0.005) | (0.01) | (0.01) | (0.01) | |

Table 2. Incidence of increasing content of diseased berries infected by Plamospara viticola on volatile compounds in Cabernet Sauvignon wines. ^{*a*} (n=2)

 ^a Analysis done 6 years after bottling.
 ^b Results concerning wines made with healthy grapes and grapes infected with increasing % of berries infected by brown rot (*P. viticola*)

^c RC : Regression coefficient with Spearman correlation test; n.s. = non significant

| Grape variety | Modality | Methyl salicylate (µg/L) | | |
|-------------------------------------|-----------------------|--|--|--|
| | Control | 0.47 (0.25) ^{<i>a</i>} / 0.37 (0.1) ^{<i>b</i>} | | |
| Cabernet Sauvignon (red wine, 2016) | Esca level 1 | 1.25 (0.17) / 0.68 (0.05) | | |
| | Esca level 2 | 5.34 (0.93) / 1.48 (0.23) | | |
| | Control | 0.06 (0.02) ^b | | |
| Furmint (dry white wine, 2017) | Incorporation level 1 | 0.16 (0.01) | | |
| | Incorporation level 2 | 0.42 (0.06) | | |
| | Incorporation level 3 | 0.62 (0.02) | | |

Table 3. Concentration of methyl salicylate in wines from healthy vines (control) or elaborated from grapes harvested on vine affected by grapevine trunk disease (Esca) (n=2)

^{*a*} Concentration determined after 1 year ageing or ^{*b*} just after the end of alcoholic fermentation