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3	Supercritical CO ₂ extraction applied toward the production
4	of a functional beverage from wine
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25 Abstract

Supercritical CO_2 extraction has been proved to be a potential tool in the recovery of aroma compounds from different natural sources and in the removal of ethanol from aqueous solutions. In this work, both ideas are combined to develop a two-step process toward the production of a low-alcohol beverage from wine, but maintaining the aroma and the antioxidant activity similar to that of the original wine.

First, the recovery of aroma from wine was attained in a countercurrent packed column (white and red wines were investigated) using very low CO₂/wine ratios. Then, the aroma-free wine recovered from the bottom of the extraction column was dealcoholized by applying different extraction conditions.

The results obtained from these studies permit the design of a two-step countercurrent CO₂ extraction process at 9.5 MPa and 313 K, in which the different CO_2 /wine ratios employed in each step lead to the recovery of aroma or the removal of ethanol. The twostep process was applied to rose wine and the low-alcohol beverage obtained proved to have similar antioxidant activity and similar aroma profile to that of the original wine.

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44 **Keywords:** Supercritical CO₂ Extraction; Non-Alcoholic Beverages; Wine; Aroma.

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48 **1. Introduction**

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50 Several drinks with low ethanol content or without ethanol have been introduced on the 51 market in recent years. The increasing public consciousness about the abuse of alcohol 52 together with the severe control of alcohol consumption in drivers have led more people 53 to consume non-alcoholic drinks, and these drinks have gained significant sales 54 percentages in the beverage industry.

Wine is one of the most complex alcoholic beverages; more than 800 volatile organic compounds (acids, esters, alcohols, aldehydes, lactones, terpenes, etc.) present in very low amounts were identified [1], which all together are responsible of each particular bouquet. Therefore, the production of an alcohol-free wine by removing ethanol while preserving the organoleptic properties of wine is a very complex and challenging problem.

61 In recent years, carbon dioxide (CO_2) extraction has been suggested as a promising 62 alternative to the recovery of aroma compounds from natural matter [2-4]. On the other 63 side, the removal of ethanol from aqueous solutions using high-pressure carbon dioxide 64 has been comprehensively studied [5-7] and thus, supercritical fluid extraction has 65 appear as a promising alternative to other conventional dealcoholization of beverages 66 techniques [8-10], such as distillation [11, 12] or inverse osmosis [13-15]. All these 67 techniques have the disadvantage of eliminating the beverage aromas together with 68 ethanol, but still, among them, supercritical CO_2 extraction is particularly attractive 69 because water, salts, proteins and carbohydrates are not substantially removed or 70 denatured [9].

71 In a European patent for producing alcohol-free wine [16], a supercritical CO_2 72 extraction is at first employed to recover aroma compounds and then, the ethanol from

the raffinate is separated in a subsequent distillation column. Mixing the extracted aroma compounds into the bottom product of distillation, alcohol-free wine can be produced. Another European patent [17] describes a process in which the ethanol and aroma are removed in a first distillation step. Then, aroma compounds are extracted from the distillate using supercritical CO_2 and are recycled to the bottom product of the distillation to obtain an alcohol-free wine product.

- 79 In a previous contribution (Ruiz-Rodriguez et al., 2010) the authors developed a model
- 80 to simulate the countercurrent supercritical CO₂ removal of ethanol from alcoholic
- 81 beverages (brandy, wine, and cider) using the GC-EoS. The results obtained compared
- 82 good with experimental data from the literature and thus, the model was used to
- 83 estimate process conditions to achieve an ethanol content reduction from ca. 10 %wt to
- 84 values lower than 1 %wt.
- In this work, supercritical CO₂ technology was employed to produce a low-ethanol content beverage from wine by combining two different countercurrent extraction steps. In the first step, the extraction and recovery of aroma from the original wine was the target, while in the second step the extraction was driven towards the dealcoholization of the aroma-free product (obtained in the first step) up to ethanol content lower than 1 %wt. The key factor to attain these two different objectives was the selection of an adequate ratio between the flow rates of solvent and wine employed.
- 93 **2. Materials and methods**
- 94
- 95 2.1 Samples and Reagents
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97	The wines (white, red and rose) employed in this work were kindly supplied by a
98	Spanish wine seller company (Bodegas Torres S.A., Vilafranca del Penedès, Catalonia,
99	Spain). Ethanol content in wine was 9.5%, 10.5% and 11.3% v/v for white, red and rose
100	wines, respectively.
101	Ethanol (GC-assay, 99.5% purity) and MilliQ-water were obtained from Panreac
102	(Barcelona, Spain) and from Millipore (Millipore Iberica, Madrid, Spain), respectively.
103	CO ₂ , N48 (99.9998% purity), was supplied by AL Air Liquide España S.A. (Madrid,
104	Spain).
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106	2.2 Supercritical fluid extraction of ethanol
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108	The supercritical fluid extraction (SFE) device (Thar Technologies) comprises a
109	countercurrent packed column of 2.8 m height with two separator cells (S1 and S2),

110 where a cascade decompression takes place. The liquid sample can be introduced into

111 the column from two different points: the top (180 cm of effective packed height) and

medium (120 cm of effective packed height) feed points. The solvent (CO₂) is fed into the column through the bottom and is heated up to the extraction temperature before be introduced into the packed column.

Once the operating pressure and temperature were reached, the wine was pumped from the top of the column at a constant flow rate of 200 ml/h during 1 h. The temperature of the extraction column was kept at 313 K in all experimental assays. Extraction pressure was varied from 9.5 to 18 MPa and thus, CO_2 densities varied from 692.3 kg/m³ to 848.9 kg/m³, maintaining an appropriate density difference between the solvent and the liquid sample (> 100 kg/m³). 121 The CO_2 flow rate was varied from 1.8 to 6.0 kg/h in order to attain CO_2 /wine ratios in 122 the range of 9 - 30 kg/l. The extracted material was decompressed up to 5 MPa in the 123 first separator cell, while the second separator was maintained near ambient pressure. 124 The temperature in both separator units was kept at 308 K in all experimental trials. 125 Once the extraction was finished, CO_2 was pumped for another 20 min to extract the 126 remaining liquid sample that could have been left inside the countercurrent column. 127 Three products were collected from each extraction assay: two ethanol enriched extracts

128 were collected from S1 and S2, and a dealcoholized wine (raffinate) from the bottom of

the column. Typically, 8-13 mL of extract was collected in S1 and amounts lower than 2

- 130 mL in S2. The mass balance closed in all experiments with accuracy greater than 85%.
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132 **2.3 Supercritical fluid recovery of aroma**

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The SFE device employed is the same equipment utilized for the ethanol removal. In this case, the wine was injected into the column from the middle point to avoid dragging of the liquid sample, at a constant flow rate during 4-6 h. That is, a total amount of 1000-1500 mL of wine was feed to the extraction column in order to recover a significant amount of aroma in the separator cells. Extraction pressure was set to 9.5 MPa, the CO₂ flow employed was in the range 0.5-1.0 kg/h and the CO₂/wine ratio around 2-4 kg/l.

Again, temperature of the extraction column was kept at 313 K in all experiments. The extracted material was decompressed up to 5 MPa in the first separator cell, while the second separator was maintained near ambient pressure. Both separators were maintained at 308 K. Once the extraction is finished, CO₂ was pumped for another 20 145 minutes to help extracting the remaining liquid sample that could have been left inside146 the countercurrent column.

Three products were obtained from each extraction assay: around 10-30 mL of extract was collected in S1, 1-5 mL of extract in S2, and a liquid raffinate sample was recovered from the bottom of the extraction column. The mass balance closed in all experiments with accuracy greater than 95%.

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152 2.4 Aroma analysis

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154 Characterization of the wine extracts was carried out by a GC-2010 (Shimadzu, Japan), equipped with a split/splitless injector, electronic pressure control, AOC-20i auto 155 156 injector, GCMS-QP2010 Plus mass spectrometer detector, and a GCMS Solution 157 software. The column used was a CW-20M (Carbowax) capillary column, 30 m x 0.32 158 mm I.D. and 0.25 µm phase thickness. Helium, 99.996% was used as a carrier gas at a 159 flow of 58,2 mL/min. Oven temperature programming was as follows: 40 °C isothermal 160 for 1 min, increased to a final temperature of 150 °C (held for 2 min) at 2 °C/min. 161 Sample injections (1 μ L) were performed in split mode (1:30). Injector temperature was 162 of 210 °C and MS ion source and interface temperatures were 230 and 280 °C, 163 respectively. The mass spectrometer was used in TIC mode, and samples were scanned 164 from 40 to 500 amu. Compounds were identified by comparison with the mass spectra 165 from Wiley 229 library and by their linear retention indexes.

166

167 **2.5 Sensory evaluation**

169 The response used to evaluate the quality of the supercritical extracts was the 170 resemblance, based on a human olfaction test, of their aroma to that of their respective 171 starting wines. Aromatic extracts were evaluated with a panel of six experts panelist 172 (four females and two males, 25-50 year-old individuals) who judged the similarity of 173 the aromas. The scale used for sensorial evaluation was not structured [18] to mark the 174 similarity between the aroma of the extracts and that of the starting wines; that is, it only 175 had two extreme points, and the right end represented the aroma of the original wine. 176 Thus, the higher the score, the higher the similarity between the aroma of the 177 supercritical extracts and the aroma of the starting wines. The distance (in centimeters) 178 to the left end was considered for the statistical analysis of the data.

179

180 **2.6 Ethanol analysis**

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A Perkin-Elmer Autosystem XL gas chromatograph (Perkin-Elmer, Norwalk CT) 182 183 equipped with a programmed split/splitless injector (PSS) and a flame ionization 184 detector (FID) was used to perform all the GC analysis. The system was coupled to a 185 Perkin-Elmer chromatography software system (Turbochrom). The column employed 186 was a 30 m x 0.25 mm i.d. fused silica capillary column (Quadrex Corp., New Haven, 187 CT) coated with a 0.25 µm layer of Carbowax 20M (polyethyleneglycol). To evaluate 188 the ethanol content of the raffinates obtained from red and white wines after 189 supercritical fluid extraction, a calibration curve was prepared using ethanol blank 190 solutions ranging from 1 to 20 % in ethanol content (v/v). The chromatographic conditions were as follows: injector temperature, 210 °C; detector temperature, 280 °C, 191 192 Helium at 15 psig was used as a carrier gas. The split ratio was 1:20 and the volume injected was 1 μ L. The oven temperature program was as follows: starting at 39 °C 193

(held for 3 min), and then heating to 65 °C (held for 1 min) at 5 °C/min, and then
heating to a final temperature of 200 °C (held for 1 min) at 40 °C/min.

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197 **2.7 Determination of antioxidant activity**

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199 **2.7.1. ABTS assay**

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201 The TEAC (Trolox Equivalent Antioxidant Capacity) assay described by Re et al. [19] 202 was used to measure the antioxidant activity of the wine samples. Briefly, ABTS[•] 203 radical cation was generated by reacting 7 mmol/l ABTS with 2.45 mmol/l potassium 204 persulfate after incubation at room temperature for 16 h in the dark. The ABTS[•] radical 205 solution was diluted with PBS (pH 7.4) to an absorbance of 0.70 - 0.20 at 734 nm. 10 µl 206 of wine (previously diluted) at five different concentrations extract was added to 0.990 ml of diluted ABTS[•] radical solution. The reaction was measured until the absorbance 207 208 reached a plateau. Trolox was used as reference standard, and results were expressed as 209 TEAC values (mmol Trolox/g extract). All analyses were done, at least, in triplicate.

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211 **2.7.2. DPPH[•] free radical-scavenging assay**

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The ability of wines to scavenge DPPH[•] free radicals was determined according to the method proposed by Brand-Williams et al. [20]. Briefly, 25 μ l of wine or standard (previously diluted) was added to 0.975 μ l of a 6 × 10⁻⁵ M solution of DPPH[•] in methanol. A control sample, containing the same volume of solvent in place of extact, was used to measure the maximum DPPH[•] absorbance. The reaction was allowed to take place in the dark until the reaction reach a plateau. Trolox was used as reference standard, and results were expressed as TEAC values (mmol Trolox/g extract). All
samples were assayed, at least, in triplicate.

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222 2.7.3. Oxygen radical absorbance activity (ORAC)

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224 The ORAC assay was performed essentially as described by Huang et al [21]. Briefly, 225 AAPH was dissolved in 10 ml of 75 mM phosphate buffer (pH 7.4) to a final concentration of 166 mM and made fresh daily. A fluorescein stock solution (8 \times 10⁻⁴ 226 227 mM) was made in 75 mM phosphate buffer and stored. The stock solution was diluted 228 1/10000 with phosphate buffer. To all experimental wells, 150 µl of working 229 fluorescein solution were added. In addition, blank wells received 25 µl of 75 mM 230 phosphate buffer, while standards received 25 μ l of trolox dilution and samples 25 μ l of 231 wine (previously diluted). Reactions were initiated by the addition of 25 μ l of AAPH 232 solution. Results were expressed as trolox equivalent antioxidant capacity.

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234 **2.8. Total phenolic content (TPC)**

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236 Total phenolic content of wines was determined with Folin-Ciocaltea reagent by the 237 Singleton et al. method [22] and the results were expressed as GAE (mg of gallic acid/L 238 of wine). Briefly, 3 mL of distilled water was mixed with 50 µL of sample or standard. 239 250 µL of Folin-Ciocalteu reagen was added and the content of the tube was mixed 240 thoroughly. After 3 min 0.75 mL of Na₂CO₃ (20% w/v) followed by 0.95 mL of water 241 was added and the mixture was allowed to stand for 2 h. The absorbance was measured 242 at 760 nm. The TPC of the wines was expressed as GAE (mg of gallic acid equivalent 243 per L of wine). All analyses were done in triplicate.

245 **3. Results and discussion**

246

- 247 **3.1 Ethanol extraction**
- 248

249 Table 1 shows the different extraction conditions (pressure and CO_2 /wine ratios) applied 250 at 313 K for the removal of ethanol from white (9.5 % v/v ethanol) and red (10.5 % v/v 251 ethanol) wines. Also given in the table are the corresponding ethanol content obtained 252 in the raffinates. Certainly, for the same CO_2 /wine ratio, CO_2 density defines the degree 253 of dealcoholization achieved: the higher CO₂ density the lower ethanol content in 254 raffinate (Exp. 1 and 4 in Table 1). Nevertheless, it can be clearly deduced from Table 1 255 that the significant variable in the dealcoholization process is the CO₂/wine ratio. This 256 was previously observed by several authors [9, 10].

- 257 According to the results obtained using the simulation GC-EoS model (Ruiz-Rodriguez
- el al., 2010) S/F ratios greater than 30 are necessary at 308 K to achieve an ethanol
- 259 reduction in wine from ca. 10 to 1 %wt. The same conclusion is driven from the

260 experimental assays: CO₂/wine ratios of ca. 30 ensured almost a complete

- 261 dealcoholization of the wines studied, under moderate temperature (313 K) and pressure
- 262 (9.5 MPa) conditions. Results obtained when combining the highest CO₂ density with
- low CO₂/wine ratios (Exp. 1) were not better than those obtained when using the lower
- 264 CO_2 density but high CO_2 /wine ratios (Exp. 3).
- 265
- 266 **3.2 Study of aroma recovery**

The same wines employed in the dealcoholization experiments (white and red wines) were employed to study the recovery of aroma from wine using supercritical CO_2 . The key idea to attain the target was utilizing a low CO_2 /wine ratio. Considering the facilities of the available experimental device, the CO_2 /wine ratio employed in this case was in the range 2-4 kg/l.

273 Certainly, low CO₂/wine ratios imply that the liquid sample is the continuous phase and 274 the supercritical solvent is the disperse phase. Thus, the solvent phase would be 275 saturated with the aroma compounds (which are present in wine in very low amounts) 276 while reduced amounts of ethanol should be extracted. On the contrary, during the 277 dealcoholization trials (CO₂/wine ratio = 9-30 kg/l), the supercritical CO₂ solvent is the 278 continuous phase and the wine is the disperse phase, and both aroma compounds and 279 ethanol a readily extracted.

280 Table 2 shows the results obtained in the recovery of aroma from white and red wines. 281 Ext. 1 and 2 in Table 2 are duplicates of the extraction accomplished for the white wine 282 at 313 K and 9.5 MPa. By comparison of the amounts (ml) of extract obtained in each 283 trial, it can be concluded that very good reproducibility is attained. Further, whilst the 284 raffinate was colored and absolutely odorless, the samples obtained in both S1 and S2 285 separators were completely transparent and very aromatic. This was assessed by 286 analyzing the scores given by the panelists to the different extracts obtained. It can 287 easily be seen that the extracts obtained in S1 and S2 corresponding to extracts 1, 2 and 288 4 obtained a high score. This means that they had a high resemblance to the original 289 aroma of the starting white and red wines. However, in the case of red wine, 290 significantly lower amounts of extract were obtained when applying the same CO₂/wine 291 ratio than in the case of white wine (Ext. 3 in Table 2). Additionally, the raffinate 292 obtained in this experiment somewhat preserved the characteristic wine odor. Thus, the 293 CO₂/wine ratio was slightly increased (Ext. 4 in Table 2) and then, also in this case, an
294 odorless raffinate was obtained.

295 According to Table 2, around 14 ml per liter of wine sample was obtained in the 296 separators (Ext. 1, 2 and 4); although in the case of white wine the amount of extract 297 recovered in S2 was larger than in the case of red wine. Moreover, the amounts of 298 extract recovered in these experiments are significantly lower than the amounts of 299 extract obtained in the dealcoholization assays (50-75 ml of extract per liter of wine). 300 The GC-MS chromatograms for extracts corresponding to the white wine are shown in 301 Figure 1. The figure shows a comparison between the chromatogram corresponding to 302 the original (white) wine, the extracts recovered in the separators and the raffinate 303 obtained from the bottom of the extraction column. As can be qualitatively observed 304 from the figures, the extracts are significantly concentrated in the aroma compounds 305 while the raffinates contain reduced amounts of aroma compounds in comparison to the

306 original wine. In the case of red wine the chromatograms followed the same pattern.

Figures 2 and 3 show the peak identification of the chromatograms corresponding to S1 extracts of experiments reported in Table 2. Figure 2 corresponds to the S1 extract recovered in Ext. 1 (white wine) while Figure 3 refers to the S1 extract of Ext. 4 (red wine). In qualitative terms, both extracts showed very similar chromatographic profile, being compounds such as 3-methyl-1-butanol, ethyl lactate, acetic acid, 2,3-butanediol and phenylethyl alcohol the ones who presented the highest chromatographic peak areas.

Further, Table 3 shows a comparison between the peak areas obtained for the different compounds identified in the original red wine and the corresponding extract (Ext. 4 in Table 2). All the injections were carried out following the same chromatographic method and conditions (see Materials and Methods section). Thus, peak areas in Table 3

318 were employed to estimate concentration factors (peak area in extract / peak area in 319 original wine) of some aroma compounds observed in the samples. Concentration 320 factors up to 50 could be calculated from the results of the GC-MS analysis. 321 Nevertheless, it should be pointed out that several compounds that are present in very 322 low concentration in the original red wine could only be identified in the extract. For 323 example, several alcohols (n-butanol, 3-methyl-1-pentanol, 1-hexanol, 3-ethoxy-1-324 propanol, 3-hexen-1-ol, 3-methyl thiol propanol), acids (3-OH-ethyl ester -butanoic 325 acid, 2-methyl-propanoic acid, isovaleric acid, 2-OH-ethyl-3-phenylpropionate, 326 diethylhydroxybutanedioate, caprylic acid, 2-OH-diethyl-pentanedioate), esters 327 (isoamyl acetate, ethyl hexanoate, ethyl octanoate), aldehides (2-furancarboxaldehyde), 328 and ethers (1-methoxy-3-methyl-butane) could only be detected in S1 extract and thus, 329 it is expected that very high concentration factors (> 50) were attained for these 330 substances.

331

332 **3.3 Production of a non-alcoholic functional beverage from rose wine**

333

On the basis of previous studies the manufacture of a non-alcoholic beverage from rose wine (11.3% v/v of ethanol) was accomplished. Two CO_2 -SFE steps were carried out, both at 313 K and 9.5 MPa, but employing different CO_2 /wine ratios in order to achieve (Step 1) the recovery of aroma and then (Step 2) the dealcoholization of the raffinate obtained in the first step. S1 separator was maintained at 5 MPa whereas in S2 the extract was depressurized up to 1 MPa. Temperature in both separators was kept at 308 K.

341 *Step 1: recovery of aroma from rose wine.* CO_2 flow rate was 0.9 kg/h and wine flow 342 rate was 0.25 l/h (CO_2 /wine ratio = 3.6). A total of 12 liters of wine were fed to the extraction column. Top and bottom products were collected during the continuous
operation; 220 ml of extract were recovered in S1 and considerably lower amounts (30
ml) in S2 separator. The mass balance closed with accuracy greater than 97%.

346 The extract obtained in S1 (18.3 ml per liter of rose wine) was completely transparent 347 and highly aromatic; the chromatogram obtained by GC-MS is shown in Figure 4. 348 Additionally, Table 4 shows the chromatographic areas of the aromatic compounds 349 identified in the original rose wine and in the S1 extract obtained. Again, high 350 concentration factors could be calculated for some aromatic compounds, such as 14 for 351 ethyl acetate, 36 for ethyl lactate, 47 for 3-methyl-1-butanol and 53 for phenyl ethyl 352 alcohol, and higher concentration factors would be expected for those compounds which 353 could not be detected in the original red wine (2-methyl-1-propanol, isoamyl acetate, 354 hexanoic acid, etc.).

355 The odorless raffinate obtained from the bottom of the extraction column contained 356 8.8% v/v of the ethanol.

Step 2: removal of ethanol from the raffinate obtained in step 1. The liquid sample collected from the bottom of the extraction column in Step 1 was utilized to completely remove the remained ethanol. In this case, the CO_2 flow rate was 4.8 kg/h and the liquid sample flow rate was 0.20 l/h (CO_2 /liquid ratio = 24). The concentration of ethanol in the raffinate obtained in this case (850 ml per liter of original rose wine) was lower than 1%.

363 *The non-alcoholic functional beverage from rose wine.* 850 ml of the raffinate 364 obtained from Step 2 (ethanol content < 1% v/v) was mixed with 18.3 ml of the extract 365 produced in Step 1. This beverage (1.1% v/v ethanol) produced from rose wine 366 contained several of the aromatic compounds detected in the original wine, as can be 367 deduced from the GC-MS analysis given in Table 4. Some substances are present almost in the same concentration (3-methyl-1-butanol, acetic acid, 2,3-butanediol, 2methyl-propanoic acid) although some other substances that were detected in the original wine, could not be detected in the non-alcoholic beverage (ethyl acetate, 3hydroxy-2-butanoate, ethyl lactate, cis-5-hydroxy-2-methyl-1,3-dioxane).

As it is shown in Table 5 aroma removal from wine only caused slight modifications in its antioxidant activity and polyphenols content. ABTS and DPPH assays shown a very small increase in the antioxidant capacity according to the TPC increment. However ORAC value was slightly smaller in this odorless raffinate, maybe to the different mechanism of action of these methods. The non-alcoholic functional beverage had similar DPPH and ORAC values than original wine, together with similar TPC. Only a smaller ABTS value was detected.

379

380 Conclusion

Supercritical fluid CO_2 extraction was employed in a two-step process to produce a novel beverage from rose wine. Several aroma compounds were determined to be present both in the original rose wine and in the low-alcoholic beverage. Further, the new beverage maintains the antioxidant capacity of the original wine; it contains around 1% v/v ethanol, and thus might be potentially commercialized with a functional claim.

386

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394 **References**

- 395 [1] P. Schreier, Flavour composition of wines: a review, CRC Food Science and
 396 Nutrition 12 (1979) 59.
- 397 [2] P.K. Rout, Development of process for extraction of floral fragrances by
 398 subcritical carbon dioxide, PhD Thesis, Indian Institute of Technology (2008).
- M. Perrut, M. Nunes da Ponte, Liquid-fluid fractionation: the extraction of aromas
 from fermented and distilled beverages, Proceedings of the 4th International
 Symposium on Supercritical Fluids, Sendai, Japan, vol. C, 1997, p. 845.
- 402 [4] F. J. Señoráns, A. Ruiz Rodríguez, E. Ibáñez, J. Tabera, G. Reglero, Isolation of
 403 brandy aroma by countercurrent supercritical fluid extraction, Journal of
 404 Supercritical Fluids 26 (2003) 129-135.
- 405 [5] E. A. Brignole, P. M. Andersen, A. Fredenslund, Supercritical fluid extraction of
 406 alcohols from water, Industrial & Engineering Chemistry Research 26 (1987) 254407 261.
- 408 [6] G. Bunzenberger, R. Marr. Countercurrent high pressure extraction in aqueous
 409 systems, in: M. Perrut (Ed.), Proceedings of the 5th International Symposium on
 410 Supercritical Fluids, vol. 2, 1988, pp. 613-618.
- 411 [7] G. Brunner, K. Kreim, Separation of ethanol from aqueous solutions by gas
 412 extraction, German Chemical Engineering 9 (1986) 246-250.
- 413 [8] U. Schobinger, Nonalcoholic wine manufacturing processes and sensory aspects,
- 414 Mitt. Gebiete Lebensm. Hyg. 77 (1) (1986) 23.
- 415 [9] I. Medina, J. L. Martínez, Dealcoholization of Cider by Supercritical Extraction
- 416 with Carbon Dioxide, Journal of Chemical Technology & Biotechnology 68 (1997)
- 417 14-18.

- 418 [10] T. Gamse, I. Rogler, R. Marr, Supercritical CO₂ extraction for utilization of excess
- 419 wine of poor quality, Journal of Supercritical Fluids 14 (1999) 123-128.
- 420 [11] H. Kieninger, J. Haimerl, Manufacture of alcohol-reduced beer by vacuum
 421 distillation, Brauwelt 121 (17) (1981) 574.
- 422 [12] R. Pérez, M.D. Salvador, R. Melero, M.I. Nadal, F. Gasque, Desalcoholización de
 423 vino mediante destilación en columna: Ensayos previos, Revista de Agroquímica y
- 424 Tecnología de Alimentos 29 (1) (1989) 124.
- 425 [13] K. Bui, R. Dick, G. Moulin, P. Glazy, A reverse osmosis for the production of low
- 426 ethanol content wine, American Journal of Enology and Viticulture 37 (4) (1986)
 427 297.
- 428 [14] G.W. Von Hodenberg, Production of alcoholfree beers using reverse osmosis,
 429 Brauwelt Int. 2 (1991) 145.
- 430 [15] H. Goldstein, C.L. Cronan, E. Chicoye, Preparation of low alcohol beverages
 431 byreverse osmosis, US Patent 4612196 (1986).
- [16] A. Wiesenberger, R. Marr, E. Kolb, J. Schildmann, R. Weisrock. Process for
 producing alcohol-reduced or alcohol-free beverages made by natural
 fermentation. European Patent No. 0 228 572 B1.
- [17] H. Seidlitz, E. Lack, H. Lackner. Process to lower the alcohol content of alcoholic
 beverages. European Patent No. 0 397 642 A1.
- 437 [18] M.A. Amerine, R.M. Pangborn, E.B. Roessler, Principles of Sensory Evaluation of
 438 Foods. Academic Press: New York, 1965.
- 439 [19] R. Re, N. Pellegrini, A. Proteggente, A. Pannala, M. Yang, C. Rice-Evans,
- Antioxidant activity applying an improved ABTS radical cation decolorization
 assay, Free Radical Biology & Medicine 26 (1999) 1231-1237.

442	[20] W.	Brand-Willi	ans, I	M.E.	Cuvelier,	C.	Berset,	Use	of	а	free
443	radic	al method t	o evalu	ate an	tioxidant	activity,	Lebens	mittel-	Wissc	hens	chaft
444	und	Fechnology 2	8 (1995)) 25-30							

- [21] D.H. Huang, B. Ou, M. Hampsch-Woodill, J.A. Flanagan, R.L. Prior, HighThroughput Assay of Oxygen Radical Absorbance Capacity (ORAC)
 Using a multichannel Liquid Handling System Coupled with a Microplate
 Fluorescence Reader in 96-Well Format, Journal of Agricultural and Food
 Chemistry 50 (2002) 4437-4444.
- 450 [22] V.L. Singleton, R. Orthofer, R.M. Lamuela-Raventós, Analysis of
 451 total phenols and other oxidations substrates and antioxidants by means of
 452 Folin-Ciocalteu reagent, Methods in enzymology 299 (1999) 152-178.

Exp.	P (MPa)	CO ₂ density (g/cm ³)	CO ₂ /wine ratio (kg/l)	% wt ethanol in raffinate
white wine				
1	18	0.820	9	3.5
2	13	0.742	12	2.1
3	9.5	0.516	29	< 1
4	9.5	0.516	9	5.5
red wine				
5	9.5	0.516	11	3.5
6	9.5	0.516	30	< 1

Table 1. CO₂-SFE for the removal of ethanol from red and white wines at 313 K.

Table 2. CO₂-SFE for the recovery of aroma from red and white wines at 313 K and 9.5

461 MPa. Total extraction time = 4 h. Total amount of wine feed to the extraction column =

462 1000 ml.

Ext. 1	Ext. 2	Ext. 3	Ext. 4
white wine	white wine	red wine	red wine
0.23	0.23	0.23	0.23
0.60	0.60	0.60	0.90
2.6	2.6	2.6	3.8
11.0	10.8	5.2	13.5
15.0	15.5	3.1	16.0
0.7	1.4	1.0	0.8
4.3	4.0	0.5	1.0
17.3	19.1	2.4	17.0
0.7	0.7	0.8	1.4
	Ext. 1 white wine 0.23 0.60 2.6 11.0 15.0 0.7 4.3 17.3 0.7	Ext. 1Ext. 2white winewhite wine0.230.230.600.602.62.611.010.815.015.50.71.44.34.017.319.10.70.7	Ext. 1Ext. 2Ext. 3white winewhite winered wine0.230.230.230.600.600.602.62.62.611.010.85.215.015.53.10.71.41.04.34.00.517.319.12.40.70.70.8

464 ^a Standard Deviation

469 Table 3. Chromatographic areas obtained in the original red wine, S1 extract and

470 raffinate (Ext. 4 in Table 2). NI: non identified compound.

4	7	1
		•

compound	original red wine	S1 extract	concentration factor
Ethyl acetate		14467940	100001
2-methyl-1-propanol	975555	28864100	29.6
Isoamyl acetate		266518	
n-butanol		597800	
3-methyl-1-butanol	6561474	193130059	29.4
Ethyl hexanoate		210295	
2-butanone,3-hydroxy	139081	1782147	12.8
2-OH-propanoic acid, methyl ester		113465	
1-pentanol,3-methyl-		70898	
2-OH-isobutyric acid, methyl ester		106333	
Ethyl lactate	2632592	(*)	
1-hexanol		1159865	
3-ethoxy-1-propanol		141465	
3-hexen-1-ol		68231	
Ethyl octanoate		241426	
Tert-butoxymethoxy, methane		46473	
2-furancarboxaldehyde		52418	
Acetic acid	3957189	11090461	2.8
Butanoic acid,3-OH-ethyl ester		287263	
2,3 butanediol	7363015	7351706	1.0
Butane,1-methoxy-3-methyl		412724	
Ethanol,2-methoxyethanol	1990796	1210931	0.6
Propanoic acid,2methyl-		435945	
2(3H)-furanone,dihydro-	213612	2277658	10.7
NI-I		169072	
Butanedioic acid, diethyl ester	310726	15553593	50.1
Isovaleric acid		518754	
3-methyl thiol propanol		759264	
NI-II		624306	
N-(-3-methylbutyl)acetamide		774003	
NI-III		890390	
Phenylethyl alcohol	1339270	50154470	37.4
2-OH-ethyl-3-phenylpropionate		461626	
Diethylhydroxybutanedioate		289933	
Caprylic acid		1466425	
2-OH-diethyl-pentanedioate		1035159	

472 (*) Chromatographic area too high leading a saturated detector response.

473

- **Table 4.** Chromatographic areas obtained in the original rose wine, S1 extract obtained
- 476 from Step 1, raffinate obtained from Step 2 (dealcoholized wine) and non-alcoholic
- 477 beverage produced. NI: non identified compound.

	original rose wine	S1 extract	dealcoholized wine	non-alcoholic beverage
Acetaldehyde		119166		
Ethyl acetate	194430	2894893		
2-methyl-1-propanol		2144850		
Isoamyl acetate		257327		
n-butanol		145410		
3-methyl-1-butanol	749848	34944236		674623
Ethyl hexanoate		172957		
3-hydroxy-2-butanoate	47548	561970		
Ethyl lactate	56900	2053307		
1-hexanol		474860		
Ethyl octanoate		203616		
2-furfural	309200		249722	210090
Acetic acid	1520309	7690182	1152546	1163573
Cis-5-hydroxy-2-methyl- 1,3-dioxane	47770	132720	35001	
2,3-butanediol	3206841	4511741	3580614	3493937
5-methyl furfural			134611	
2-methyl-propanoic acid	964189	826606	1157857	1152847
1,2-propanediol			276019	245267
2-(3H)-dihydrofuranone	102998	288085	97772	64033
Butyric acid		322514		
NI-I			25156	
NI-II			84553	
Diethyl ester butanedioic acid		510897		
Hexanoic acid		3325559		
Phenyl ethyl alcohol	168806	9062757		106534
NI-III				505895
2-furancarboxaldehyde- 5(hydroxymethyl)-				
NI-IV				2301994
Diethyl hydroxybutanedioate		804047		
Caprylic acid		6615062		
TOTAL	7090559	78062762	6793851	9918793

Table 5. Antioxidant activity of rose wine, raffinate and non-alcoholic beverage.

	ABTS ^b	DPPH ^b	ORAC ^b	TPC
Original wine	$8.751\pm0.055^{\text{b}}$	1.499 ± 0.020^{b}	17.290 ± 0.593^{a}	$429.860 \pm 14.801^{\text{b}}$
Raffinate	9.313 ± 0.181^a	1.666 ± 0.140^a	15.611 ± 0.550^{b}	$444,513 \pm 11.841^{a}$
Non-alcoholic beverage	$8.148 \pm 0.046^{\circ}$	1.542 ± 0.042^{b}	16.653 ± 0.834^{a}	423, 587 ± 12. 617 ^b

^aDifferent superscript letters denotes statistically significant differences (p < 0.05) among data in the same

484 column

485 ^bAntioxidant activity was expressed as TEAC mmol of Trolox/g of extract.

486 ^cTotal phenolic compounds was expressed as mg GAE/l)



490 Figure 1. Aroma recovery from white wine (Ext. 1 in Table 2): comparison between the
491 GC-MS chromatograms obtained for (a) the original wine; (b) S1 extract; (c) S2 extract;
492 (d) raffinate.



494

496 Figure 2. Chromatogram corresponding to the extract recovered from white wine in S1497 separator (Ext. 1 in Table 2).

498 1) ethyl acetate, 2) 2-methyl-1-propanol, 3) isoamyl acetate, 4) n-butanol, 5) 3-methyl,1-butanol, 6) ethyl 499 hexanoate, 7) hexyl acetate, 8) 2-butanone,3-hydroxy-, 9) 2-hydroxy-isobutyric acid,methyl ester, 10) 500 ethyl lactate, 11) 1-hexanol, 12) 3 ethoxy-1-propanol, 13) 3-hexen-1-ol, 14) ethyl octanoate, 15) acetic 501 acid, 16) butanoic acid, 3-hydroxy-ethyl ester, 17) 2,3-butanediol, 18) linalool, 19) etanol, 2-502 methoxyethanol, 20) 1.2 propanediol, 21) 2(3H)-furanone, dihydro-, 22) Ho-trienol, 23) NI-I, 24) 503 butanoic acid, 25) butanedioic acid, dietil ester, 26) isovaleric acid, 27) 3-methyl thiol propanol, 28) 1,3 504 propanediol, diacetate, 29) Acetic acid, 2-phenylethyl ester, 30) NI-II, 31) Nerol, 32) N-(3-505 methylbutyl)acetamide, 33) phenylethyl alcohol, 34) ethyl-2-hydroxy-3-phenylpropionate, 35) 3.7-506 dimethyloct-1-en-3,7-diol, 36) diethylhydroxybutanedioate, 37) caprylic acid, 38) glycerol. NI: non 507 identified compound.

508

509





Figure 3. Chromatogram corresponding to the extract recovered from red wine in S1
separator (Ext. 4 in Table 2).

516 1) ethyl acetate, 2) 2-methyl-1-propanol, 3) isoamyl acetate, 4) n-butanol, 5) 3-methyl,1-butanol, 6) ethyl 517 hexanoate, 7) 2-butanone, 3-hydroxy-, 8) propanoic acid, 2-hydroxy-, methyl ester, 9) 1-pentanol, 3-518 methyl-, 10) 2-hydroxy-isobutyric acid, methyl ester, 11) ethyl lactate, 12) 1-hexanol, 13) 3 ethoxy-1-519 propanol, 14) 3-hexen-1-ol, 15) ethyl octanoate, 16) tert-butoxymethoxy, methane, 17) 2-520 furancarboxaldehyde, 18) acetic acid, 19) butanoic acid, 3-hydroxy-ethyl ester, 20) 2,3-butanediol, 21) 521 butane,1-methoxy-3-methyl-, 22) etanol, 2-methoxyethanol, 23) propanoic acid, 2-methyl, 24) 2(3H)-522 furanone, dihydro-, 25)NI-I, 26) butanedioic acid, dietil ester, 27) isovaleric acid, 28) 3-methyl thiol 523 propanol, 29) NI-II, 30) N-(3-methylbutyl)acetamide, 31) NI-III, 32) phenylethyl alcohol, 33) ethyl-2-524 hydroxy-3-phenylpropionate, 34) diethylhydroxybutanedioate, 35) caprylic acid, 36) dietil-2-hydroxy-525 pentanedioate. NI: non identified compound. 526





532 Figura 4. Chromatogram corresponding to the extract recovered from rose wine (S1533 separator).

1: carbon dioxide, 2: acetaldehyde, 3: ethyl acetate, 4: 2-methyl-1-propanol, 5: isoamyl acetate, 6: nbutanol, 7: 3-methyl-1-butanol, 8: ethyl-hexanoate, 9: 3-hydroxy-2-butanoate, 10: ethyl lactate, 11: 1hexanol, 12: ethyl-octanoate, 13: acetic acid, 14: cis-5-hydroxy-2-methyl-1,3-dioxane, 15: 2,3-butanediol,
16: 2-methyl-propanoic acid, 17: 2(3H)-dihydro-furanone, 18: butyric acid, 19: dietil succinate, 20: 3methyl-mercapto-1-propanol, 21: metil-2-acetylhydroxy-palmitate, 22: butanedioic acid, dietil ester, 23:
hexanoic acid, 24: phenyl ethyl alcohol, 25: diethyl hydroxybutanedioate, 26: caprylic acid.