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*Original*

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(Article begins on next page)

1 **CONVENTIONAL AND ULTRASOUND-ASSISTED EXTRACTION OF RICE**  
2 **BRAN OIL WITH ISOPROPANOL AS SOLVENT**

3

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15

16

17 **Abstract**

18 After cereal harvesting, rice is subjected to several milling processes to remove hull,  
19 germ, and bran and produce the final white rice. The bran represents around 10% of  
20 total grain weight and is usually considered as waste material. One of the most common  
21 rice bran applications is the extraction of rice bran oil, rich in  $\gamma$ -oryzanol, which has  
22 shown many health benefits including antioxidant, anti-inflammatory, and anti-  
23 hypercholesterolemic properties. Rice bran oil is usually extracted by organic solvents,  
24 which are toxic for health and the environment. In this work, rice bran oil was extracted

25 through isopropanol extraction, and the best-operating temperature and bran to solvent  
26 ratio have been identified. After that, an ultrasound-assisted extraction was conducted  
27 at room temperature and with the same rice bran to solvent ratio of the isopropanol  
28 extraction.

29 The kinetics evaluation through Peleg's model showed that the solvent extraction  
30 reaches the steady-state after 15 minutes while the ultrasound-assisted extraction  
31 reaches the steady-state after only 1 minute producing very similar yields in rice bran oil  
32 and  $\gamma$ -oryzanol. Comparing these two green extraction techniques through a life cycle  
33 assessment, it has emerged that with the same amount of rice bran oil produced, the  
34 ultrasound-assisted extraction is the less environmentally impacting process. The room  
35 temperature ultrasound-assisted extraction allows minimizing-the energy and time  
36 consumption demonstrating to be a sustainable process in line with the principles of  
37 green chemistry.

38

39 **Keywords:** Rice bran oil, green extraction, ultrasound-assisted extraction, life cycle  
40 assessment.

41

42

### 43 **1. Introduction**

44 Rice is one of the most important food crops in the world, representing a huge  
45 contribution to the dietary need of the global population [1]. According to FAO, world  
46 rice production exceeded 755 million tons in 2019 [2]. After harvesting, the rice grain  
47 undergoes a milling process to remove all the external layers making the edible white  
48 rice kernel [3]. During these milling operations, around 40% of the total grain is lost  
49 due to discarding the byproducts, including the husk, the bran, the germ, and the broken

50 rice [4]. Usually, these are burned or used as animal feed, but they may represent an  
51 excellent source of bioactive compounds, making them suitable for nutraceutical,  
52 cosmetic and pharmaceutical applications [2]. In particular, rice bran, which represents  
53 around 10% of the grain weight, contains proteins, fibers, and oil, this latter rich in  
54 bioactive and antioxidant compounds [5]. This rice bran oil (RBO) presents a balanced  
55 fatty acid composition and high levels of functional molecules, including phytosterols,  
56 tocopherols, tocotrienols, and other nutrients. Thanks to its exceptional properties,  
57 RBO is commonly used in Asian countries where it is considered as a "healthy oil".  
58 Indeed, it has been demonstrated that RBO has antihypertensive, antidiabetic, anti-  
59 obesity, and anticarcinogenic properties due to its significant antioxidant and anti-  
60 inflammatory activity [6–8]. Several studies confirmed that one of the main responsible  
61 for these beneficial effects is  $\gamma$ -oryzanol, an antioxidant mixture of ferulic acid esters of  
62 phytosterols, present at high levels in RBO [9]. The conventional method used for the  
63 commercial extraction of RBO is solvent extraction (SE) using hexane, a petroleum-  
64 derived, flammable, and toxic solvent, dangerous for human health and the  
65 environment. The disadvantages of this type of extraction led most researchers to look  
66 for alternative approaches, focusing on non-conventional and non-thermal extraction  
67 techniques, for RBO extractions [10]. These innovative techniques employ less  
68 dangerous solvents, often combined with one or more process intensification steps to  
69 reduce time and energy waste, obtaining high-quality extracts devoid of toxic residues  
70 [2, 11]. Some studies demonstrated that short-chain alcohols might represent an  
71 excellent alternative to hexane as a solvent in RBO extraction. In particular, the use of  
72 isopropanol allowed to obtain a high yield in oil and  $\gamma$ -oryzanol [12].

73 Moreover, in recent years, ultrasound-assisted extraction (UAE), thanks to the  
74 phenomenon of "acoustic cavitation," has become an effective green method for oil  
75 extraction. In their review, Mushtaq et al. [13] concluded that the UAE represents a  
76 good alternative for extracting edible oil. This technique allows operating at low  
77 temperatures and times, reducing solvent consumption, avoiding thermal damage and  
78 preserving their structural and bioactive properties. Indeed, UAE was successfully  
79 applied to extract bioactive compounds from several natural matrices, but only a few  
80 studies on RBO extraction focus on this green technique [14].

81 The present work aims to show the suitability of isopropanol as non-conventional  
82 solvent for RBO extraction and then demonstrate that the substitution of high  
83 temperature with room-temperature ultrasound in the extraction with isopropanol as the  
84 solvent allows minimizing the extraction time and energy consumption, obtaining  
85 comparable yields.

86 Isopropanol SE and room temperature UAE were compared in terms of oil yield and  $\gamma$ -  
87 oryzanol content. The extraction kinetics were determined using Peleg's model to  
88 identify the best extraction time. Moreover, a life cycle assessment (LCA) study was  
89 performed to compare the environmental sustainability of the two processes and to  
90 choose the most environmental-friendly extraction process.

91

## 92 **2. Materials and methods**

### 93 **2.1 Material and chemicals**

94 Cryo-milled rice bran sample, with a particle diameter of 500  $\mu\text{m}$ , were supplied by  
95 Agrindustria Tecco S.R.L. and stored at  $-20.0\text{ }^{\circ}\text{C}$  until extraction. HPLC grade hexane,  
96 methanol, acetonitrile, acid acetic and isopropanol used for extraction and high-

97 performance liquid chromatography (HPLC) analysis and  $\gamma$ -oryzanol standard for  
98 quantification were purchased from Merck (Darmstadt, Germany).

## 99 **2.2 Isopropanol extraction**

100 Rice bran was mixed with isopropanol in a Pyrex reaction flask connected with a Liebig  
101 reflux condenser. The flask was put into a water bath with a magnetic stirrer. To choose  
102 the best condition, 1:3, 1:6, and 1:9 solid-to-solvent ratio (w/v) was used at different  
103 temperatures (30, 45, 60 °C). The extraction time was fixed for 1 h. After these  
104 preliminary studies, the best solid-to-solvent ratio was chosen. The temperature range  
105 was extended to determine the effect of temperature on the yield of extracted  
106 components. The temperature investigated were 30, 60, 90, and 120 °C. At the best  
107 temperature, a kinetic study was performed examining the yield in oil and  $\gamma$ -oryzanol  
108 against time from 1 minute to 120 min.

## 109 **2.3 Ultrasound-assisted extraction**

110 For the UAE experiments, a VCX750 Ultrasonic Processors (Sonic and Materials Inc.),  
111 with a frequency of 20 kHz and 40 % of amplitude, equipped with a 13 mm probe was  
112 used. Rice bran (5 g) was mixed with 45 mL of isopropanol (1:9 w/v) in a Pyrex  
113 reaction flask put into a water bath with a magnetic stirrer. The temperature was  
114 maintained at room temperature (25 °C) and controlled with a thermocouple. A kinetic  
115 study was performed investigating the yield in oil and  $\gamma$ -oryzanol against time from 10 s  
116 to 30 min.

## 117 **2.4 Hexane extraction**

118 Isopropanol and UAE were compared with a conventional hexane extraction following  
119 the method of Pengkumsri et al. [15]. Rice bran was mixed with hexane in a 1:10 (w/v)

120 ratio in a Pyrex reaction flask connected with a Liebig reflux condenser. The flask was  
121 put into a water bath with a magnetic stirrer at 40 °C for 30 min.

## 122 **2.5 Determination of RBO Yield**

123 After the extractions, all the samples were filtered two times using a paper filter  
124 Whatman grade 1 to separate the liquid phase from the exhausted rice bran. Then the  
125 solvent was evaporated using a Heidolph Rotary Evaporator, Laborota 4000. The RBO  
126 yield was calculated on the base of Eq. 1:

127

$$RBO\ Yield\ (\%) = \frac{RBO\ (g)}{Rice\ bran\ (g)} \times 100 \quad (1)$$

## 128 **2.6 Determination of $\gamma$ -oryzanol content**

129 After the evaporation of the solvent and the determination of the oil yield, the RBO  
130 samples were resuspended in 15 mL of isopropanol, and  $\gamma$ -oryzanol content was  
131 determined by reversed-phase HPLC. The HPLC system (Shimadzu 20A Prominence)  
132 was equipped with a Kinetex C18 column (5  $\mu$ m, 150 x 4.6 mm) by Phenomenex and  
133 photodiode array (PDA) detector using an isocratic elution. The mobile phase was  
134 composed by methanol, acetonitrile and 0.03 % acid acetic at a ratio 52:45:3 (v/v/v) [16,  
135 17]. The flow rate was maintained at 0.8 mL/min, and the column oven was  
136 thermostated at 30 °C.  $\gamma$ -oryzanol content was determined through a calibration curve  
137 prepared using 8 different concentrations of  $\gamma$ -oryzanol standard (0.01-0.8 mg/mL) in  
138 isopropanol. The limit of detection (LOD) and the limit of quantification (LOQ) were  
139 calculated with the following equation, where  $\sigma$  is the standard deviation of the response

140 and S is the slope of the calibration curve [18]. LOD and LOQ were respectively 0.01  
141 and 0.04 mg/mL.

142

143

$$LOD = \frac{3.3 \times \sigma}{S} \quad (2)$$

$$LOQ = \frac{10 \times \sigma}{S} \quad (3)$$

144 The  $\gamma$ -oryzanol yield was calculated using the formula:

$$\gamma - \text{oryzanol yield} = \frac{\gamma - \text{oryzanol (mg)}}{\text{Rice bran (g)}} \quad (4)$$

145

## 146 **2.7 Statistical analysis**

147 All the extraction experiments were performed in triplicate and analyzed by one-way  
148 ANOVA (analysis of variance) with a Tukey's posthoc test ( $P \leq 0.05$ ), after the  
149 assessment of the fundamental assumptions of ANOVA: the normality of distributions  
150 (Shapiro-Wilk test, p-value  $N 0.05$ ) and the homogeneity of the variances of the  
151 residuals (Levene's test with  $P(NF) N 0.05$ ). The statistical software R (version 4.0.4 -  
152 Feather Spray - 2021) was used for all.

153

154

155 **2.8 Extraction kinetics**

156 To describe the SE and UAE kinetics of RBO and  $\gamma$ -oryzanol from rice bran was used  
157 the model proposed by Peleg [19], a two-parameters, non-exponential empirical  
158 equation, originally proposed to describe sorption curves and adapted for the extraction  
159 process in the form:

$$C(t) = C_0 + \frac{t}{K_1 + K_2 \times t} \quad (5)$$

160

161 where  $C(t)$  is the RBO or  $\gamma$ -oryzanol yield [(g RBO / g bran)·100] or [mg  $\gamma$ -oryzanol / g  
162 bran], respectively, at time  $t$ ;  $t$  is the extraction time [min],  $C_0$  is the yield at time  $t = 0$ ,  
163  $K_1$  is Peleg's rate constant [min· (g bran / g RBO) ·100] or [min· (g bran / mg  $\gamma$ -  
164 oryzanol)], and  $K_2$  is Peleg's capacity constant [g bran · 100 / g RBO] or [g bran / mg  $\gamma$ -  
165 oryzanol]. Since,  $C_0$  is considered zero, and this term can be omitted from Peleg's  
166 equation, the final form of the equation used is:

$$C(t) = \frac{t}{K_1 + K_2 \times t} \quad (6)$$

167

168 It should be noted that a lower  $K_1$  value means a faster rate of extraction, and a lower  $K_2$   
169 value suggests maximum yield [20]. The Peleg's rate constant  $K_1$  relates to the  
170 extraction rate ( $B_0$ ) at the start ( $t = t_0$ ).

$$B_0 = \frac{1}{K_1} \quad (7)$$

171

172 The Peleg capacity constant  $K_2$  relates to a maximum of extraction yield,  $C_e$  at  
173 equilibrium when  $t = \infty$ .

$$C_{t \rightarrow \infty} = C_e = \frac{1}{K_2} \quad (8)$$

174

175 The accordance of experimental ( $\hat{y}_i$ ) data and model-predicted results ( $y_i$ ) were  
176 established by correlation coefficient ( $R^2$ ), and root mean square error (RMSE) as  
177 follows, where  $n$  represents the number of experiments.

$$RMSE = \sqrt{\frac{\sum_{i=1}^n (\hat{y} - y)^2}{n}} \quad (9)$$

178

## 179 **2.9 Life cycle assessment**

180 LCA was performed with SimaPro 9.0.48 software, database Ecoinvent 3.0

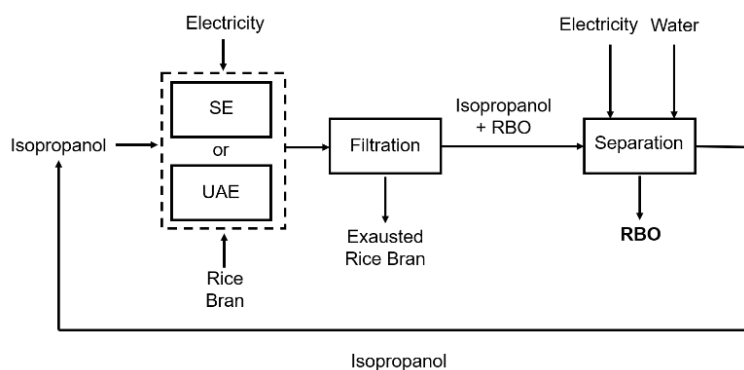
### 181 **2.9.1 Goal and scope**

182 The goal of LCA was to compare SE vs. UAE of RBO, to choose the best extraction  
183 process in terms of environmental sustainability.

184 The functional unit (FU) was 1 g of RBO produced.

185 The boundary conditions are shown in Fig. 1. Briefly, the entire process is divided in  
186 three steps: extraction (SE or UAE), filtration and separation. In the extraction step,  
187 the entered flows of matter and energy are isopropanol, rice bran and electricity. The  
188 rice bran and the isopropanol come out of the extraction together and enter the second  
189 step, where filtration occurs. Here the exhausted rice bran, which is discarded, and the

190 isopropanol containing the RBO are separated. Isopropanol and RBO enter the last step  
 191 where they are separated by a rotavapor which needs cooling water and electricity. In  
 192 this step, the isopropanol is removed from the RBO and recirculated for a new  
 193 extraction.



194

195 **Fig. 1** LCA boundary conditions.

196

197 **2.9.2 Life cycle inventory**

198 The life cycle inventory (LCI) defined all inputs and outputs involved in the processes.

199 The primary data came from the present study, the produced emissions, the consumed  
 200 material, and the required energy were referred to this FU.

201 Expansion system methodology was applied to the recovery of isopropanol in the  
 202 separation step. The secondary data were taken from Ecoinvent 3.01 and reported in  
 203 Table 1.

204

205

206

207

208

209

210 **Table 1** Secondary data from Ecoinvent 3.01

Electricity	Electricity, medium voltage {Europe without Switzerland market}  Alloc Def Unit
Isopropanol	Isopropanol {GLO} market for  Alloc Def Unit
Rice bran	Rice bran from dry milling at plant   CN mass
Water	Tap Water from natural resource

211

212

### 213 **2.9.3 Life cycle impact assessment**

214 Life cycle impact assessment (LCIA) was performed with the ReCIPE Midpoint (H)

215 method. In the present study, the analyzed impact category were: Climate change (kg

216 CO<sub>2</sub> eq), Ozone depletion (kg CFC-11 eq), Human toxicity (kg 1,4-DB eq) and

217 Freshwater eutrophication (kg P eq).

218

## 219 **3. Results and discussion**

### 220 **3.1 Solvent extraction**

221 The solvent chosen for all the extractions experiments was isopropanol. Traditionally,

222 RBO was extracted using hexane as the solvent because it presents a low corrosiveness,

223 high stability, and a good capacity for dissolving oil and relevant compounds such as  $\gamma$ -

224 oryzanol [12, 21–23]. Despite these advantages, hexane presents health and

225 environmental risks, therefore researchers and oil industries are focusing on reliable  
226 alternative solvents. Moreover, hexane derives from a non-renewable source and  
227 presents high flammability and toxicity for the environment and human health [22]. A  
228 few types of solvents have been tested to substitute hexane as an extractant of vegetable  
229 oil. In particular, short-chain alcohols, including isopropanol, are particularly promising  
230 due to their low toxicity [23]. Isopropanol is labeled as "recommended" or "preferred"  
231 in several green-solvent selection guidelines, such as the GlaxoSmithKline (GSK),  
232 Pfizer, Sanofi, AstraZeneca, and Green Chemistry Institute-Pharmaceutical Roundtable  
233 (GCI-PR) [2]. Furthermore, all the components of  $\gamma$ -oryzanol present an alcohol group  
234 deriving from the ferulic acid; this increases the polarity, making them more soluble in  
235 polar solvents such as short-chain alcohols, including isopropanol [21].

236 At the beginning of this work, a preliminary study on the effects of different solid-to-  
237 solvent ratios at various temperatures was performed. After that, using the best solvent-  
238 to-solid ratio condition, the temperature range was extended to determine the effect of  
239 temperature on the yield of extracted components. At this step, all the extractions were  
240 conducted for 1 hour. Figure 2 a) shows the oil yield using 1:3, 1:6, and 1:9 (w/v) solid-  
241 to-solvent ratios at three different temperatures (30, 45, and 60 °C). Remarkably, the  
242 yield of RBO grows with the increase of the solvent volume at any temperatures tested.  
243 This result is in line with Ruen-Ngam et al. [24], which investigated the use of different  
244 solvents with an increasing solid-to-solvent ratio and stated that a low volume of solvent  
245 does not allow its complete penetration into the rice bran material. The same trend was  
246 noted by Hu et al. [12], which obtained a higher yield of RBO and  $\gamma$ -oryzanol,  
247 increasing the solvent-to-bran ratio, using both hexane and isopropanol. Indeed the  
248 solvent extraction of RBO is mainly considered as a mass transfer process between the

249 solid and liquid phases, where the oil passes from the bran powder to the solvent  
250 through a diffusion mechanism [25]. The effect of the solid-to-solvent ratio on the  
251 extraction yield is coherent with the mass transfer process. The concentration gradient  
252 between the solid and the liquid represents the driving force of this phenomena [26],  
253 and it is more significant when the solid is in contact with a large volume of solvent.  
254 The high quantity of isopropanol reduces the saturation level of the solvent, increasing  
255 oil extraction yields [26]. The same results have been found for the alcoholic SE of  
256 antioxidants compounds from different natural sources such as grape pomace, olive  
257 leaves or stonebreaker [27–29].

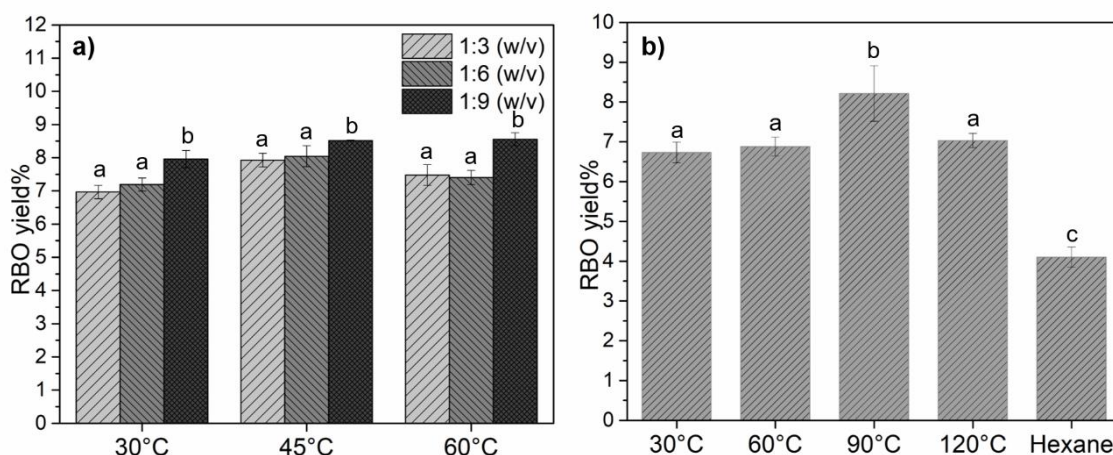
258 Because no striking differences were noted between yields at the previously mentioned  
259 temperatures, it was chosen to test the 1:9 solid-to-solvent ratio at 30, 60, 90, and 120  
260 °C to understand if the high temperatures can affect the extraction. As reported in Fig.  
261 2b, a temperature of 90 °C ( $p < 0.05$ ) gave the best results in terms of RBO extraction  
262 yield.

263 Indeed, the temperature is one of the most significant factors influencing the mass  
264 transfer and the RBO yield [25]. Capellini et al. [23] performed RBO extraction using  
265 isopropanol and ethanol at different temperatures (50, 60, 70, and 80 °C). They found  
266 that temperature increase resulted in a growth in oil yield, regardless of the solvent. The  
267 same results had been found by, Imsanguan et al.[28], Xu and Godber [19], and Oliveira  
268 et al.[20]. All these authors [19, 20, 28] agree that the diffusivity and the solubility of  
269 the compounds to extract increase with the increase of temperature, and the extraction  
270 output is enhanced [30]. At high temperatures, degradation of the sample matrix  
271 structure occurs, making it more permeable to the solvent. Moreover, a temperature rise  
272 produces a decrease in solvent viscosity, increasing its diffusivity. However, the high

273 temperature may cause the degradation of thermolabile compounds and the evaporation  
274 of the solvent. This has been reported for different plant matrices such as tomatoes,  
275 grape waste material and muiltle [20, 31–33]. Furthermore, all these extractions have  
276 been compared to hexane extraction. As can be noted from Fig. 2, the conventional  
277 hexane extraction, proposed by Pengkumsri et al. [15], and used as the reference  
278 method, produced only  $4.1 \pm 0.25$  g of RBO/100 g of rice bran. This value is lower than  
279 the yields obtained with isopropanol in any operating condition, which reaches the  
280 maximum of  $8.21 \pm 0.69$  g of RBO/100 g of rice bran at  $90^{\circ}\text{C}$  and bran to solvent ratio  
281 of 1:9, demonstrating that isopropanol is a solvent suitable to replace hexane in RBO  
282 extractions. These results are confirmed by Ruen-Ngam et al., Oliveira et al. and  
283 Capellini et al. [22, 24, 34] which affirm that alcoholic solvents like isopropanol, can  
284 extract higher amounts of oil than hexane due to their high polarity. In particular,  
285 isopropanol can extract higher quantities of phospholipids and unsaponifiable material  
286 from solid matrices and obtain RBO richer in  $\gamma$ -oryzanol. Indeed, the structure of  $\gamma$ -  
287 oryzanol presents both sides of polar and non-polar (ferulic acid and sterol) making this  
288 group of molecules particularly soluble in short-chain alcohols which show dielectric  
289 constant higher than hexane and lower than water [24].

290

291



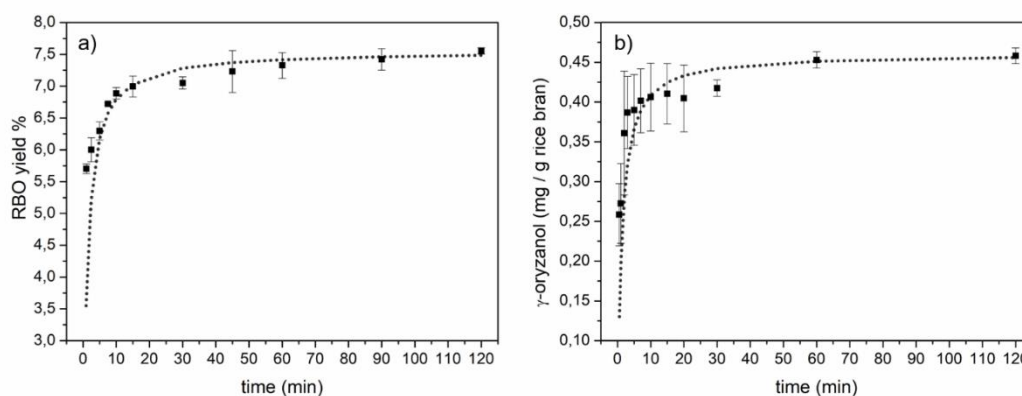
292

293 **Fig. 2** a) Effect of bran to solvent ratio at different temperatures on RBO yield (%). b)  
 294 effect of higher temperature on the RBO yield (%) with 1:9 (w/v).

### 295 3.2 SE kinetic

296 The kinetic study was performed measuring the yields in RBO and  $\gamma$ -oryzanol against  
 297 the time, starting from 1 min up to 2 h. RBO yield % and  $\gamma$ -oryzanol were plotted in  
 298 function of time (Fig. 3). The shapes of the graphs indicate that the extraction yields are  
 299 significantly time-dependent. The yields in RBO and  $\gamma$ -oryzanol rise rapidly with time  
 300 initially, and then, after the knee at about 15 min, start to keep constant producing  $6.89$   
 301  $\pm 0.089$  g of RBO/ 100 g of rice bran and  $0.41 \pm 0.038$  mg of  $\gamma$ -oryzanol/g of rice bran.  
 302 These profiles fit with the typical SE curve composed of a fast extraction step called  
 303 "washing phase", followed by a slower extraction step called "diffusion step" [35].  
 304 Indeed, the mass transfer rate into the solvent is exceptionally high at the beginning of  
 305 extraction, when the solvents penetrate the rice bran, thanks to the elevated  
 306 concentration gradients. Progressing with the extraction, the solutes diffuse from the  
 307 interior of the bran to the solvent. The mass transfer of solutes becomes more difficult

308 because the concentration gradient between the solid and the liquid phase decreases and  
309 the extraction rate becomes slower [20, 35]. The obtained constants of the model (rate  
310 constant  $K_1$ , constant capacity  $K_2$ ) and the calculated parameters (initial extraction rate  
311 ( $B_0$ ) and the maximum yield extraction ( $C_e$ ), regression coefficient ( $R^2$ ), and the root  
312 mean square error (RMSE) are reported in Table 2. The model fitted well with the  
313 experimental data, with reasonable accuracy, as evidenced by high  $R^2$  and low RMSE.  
314 In particular, experimental data fit slightly worse in the case of  $\gamma$ -oryzanol yield, but this  
315 could be due to the multiple treatments of the sample before the HPLC analysis, causing  
316 uncertainty in measurement and high standard deviation. In both cases, there was  
317 concordance between experimental and predicted yield values. Yields are lower than  
318 those reported in the literature [5, 36, 37], but this is certainly due to different  
319 experimental conditions and, above all, to the type of rice bran used for the extraction.  
320 Indeed the composition of rice bran changes based on the rice variety, the growing  
321 conditions, the milling system employed and the stabilization process [2]. Peleg's  
322 equation is one of the most suitable models to describe the SE from plant matrix as  
323 demonstrated by several studies. Karacabey et al. [38] compared the first-order kinetic  
324 model, Peleg's model, two-site kinetic model and modified Gompertz equation to  
325 describe solid–liquid extraction kinetics of trans-resveratrol from grape cane. Jurinjak  
326 Tusek et al. [39] compared Peleg's, Page's, and Logarithmic model for total  
327 polyphenols, antioxidants extraction yield from Asteraceae plants. Poojary and  
328 Passamonti [20] the first-order kinetic model, the mass transfer model, and Peleg's  
329 model for understanding the behavior of lycopene extraction from tomato processing  
330 waste. All these authors agree that Peleg's model showed a better fit to the experimental  
331 data than other models investigated in their studies.



332

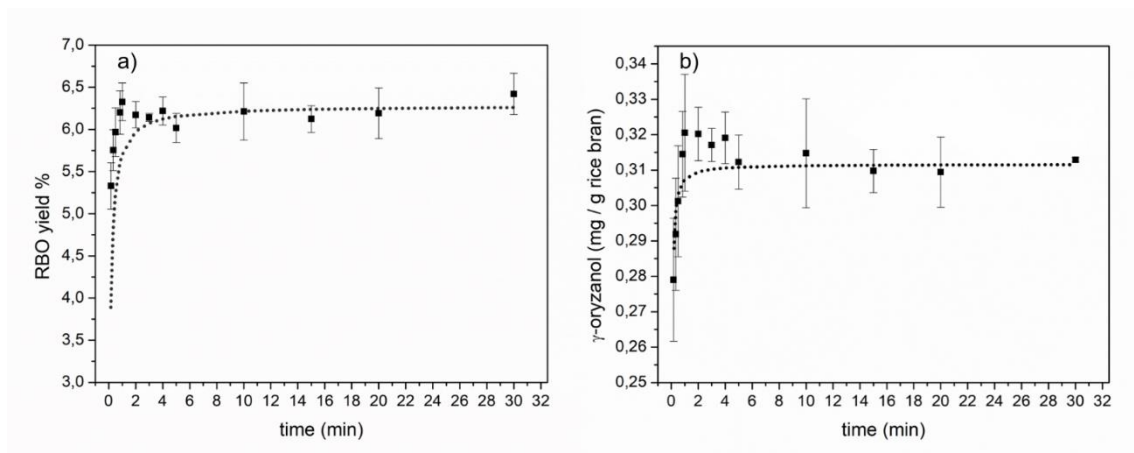
333 **Fig. 3** Isopropanol extraction (SE) kinetics of a) RBO and b)  $\gamma$ -oryzanol. Black squares  
 334 represent the experimental values. Each experiment was conducted in triplicate, and the  
 335 error bars correspond to standard error. Dotted lines represent the modeled values by  
 336 Peleg's equation.

337

### 338 3.3 UAE kinetic

339 After determining the best isopropanol extraction conditions in terms of temperature  
 340 and bran to solvent ratio, an intensification process step was performed to work at room  
 341 temperature and reduce the reaction time. The extraction was conducted with the help of  
 342 ultrasound as described in Paragraph 2.3, maintaining the bran to solvent ratio of 1:9, as  
 343 in the previous experiments. The extraction yields in RBO and  $\gamma$ -oryzanol in the  
 344 function of time were plotted to find the best extraction time and to understand the  
 345 process kinetic (Fig. 4). These two yields were initially measured every 10 s up to 1 min  
 346 and then less frequently up to 30 min. As shown in Fig.4 the UAE reaches the steady-  
 347 state after only 1 minute producing  $6.33 \pm 0.22$  g of RBO/100 g of rice bran and  $0.32 \pm$   
 348  $0.016$  mg of  $\gamma$ -oryzanol/g of rice bran. Peleg's model was adapted to experimental  
 349 conditions and used for data approximation (see paragraph 3.2) and the results are

350 reported in Tab. 2. The initial extraction rate ( $B_0$ ) is much higher than the  $B_0$  found for  
 351 the isopropanol extraction. Indeed, the knee of the curve occurs at about 1 min, where  
 352 yields reach the maximum values and then stabilize. The maximum yield extraction ( $C_e$ )  
 353 is slightly lower than SE and the  $R^2$ , but this may be due to the difficulty of manually  
 354 measuring the yield every 10 s. This difficulty caused a high uncertainty and an elevated  
 355 standard deviation producing the worst fit with the model. Until now, to the best of the  
 356 authors' knowledge, previous studies on the combination of isopropanol and UAE do  
 357 not exist. Cravotto et al. [40] and Khoei et al. [41] studied the RBO extraction using  
 358 UAE and water as the solvent, demonstrating that the ultrasound is suitable for aqueous  
 359 extracting rice bran oil. Other authors studied the effect of UAE of RBO combined with  
 360 short-chain alcohols like methanol and ethanol [42, 43], demonstrating the feasibility of  
 361 this kind of extraction. Still, no one managed to complete the extraction in such a short  
 362 time. Kumari et al. [41] and Galvan et al. [42] showed in their works that Peleg's model  
 363 efficiently describes the kinetic of UAE for other plant matrixes such as potato peels  
 364 and black chokeberry wastes.



365  
 366 **Fig. 4** UAE kinetics of a) RBO and b)  $\gamma$ -oryzanol. Black squares represent the  
 367 experimental values. Each experiment was conducted in triplicate, and the error bars

368 correspond to standard error. Dotted lines represent the modeled values by Peleg's  
369 equation.

### 370 **3.4 SE and UAE kinetics comparison**

371 The obtained constants of the model (rate constant  $K_1$ , constant capacity  $K_2$ ) and the  
372 calculated parameters, initial extraction rate ( $B_0$ ) and the maximum yield extraction  
373 ( $C_e$ ), regression coefficient ( $R^2$ ), and the root mean square error (RMSE), for the two  
374 kinds of extractions, are reported in Table 2. As can be noted, the initial extraction rate  
375 is higher for the UAE than SE, demonstrating that UAE can reach a steady state in a  
376 shorter time. Assuming infinite extraction time, the maximum yield obtained is 7.55 g  
377 RBO/100 g bran and 0.46 mg  $\gamma$ -oryzanol/g bran for the SE and 6.34 g RBO/100 g bran  
378 and 0.31 mg  $\gamma$ -oryzanol/g bran for the UAE. SE allows obtaining slightly higher  
379 quantities of RBO and  $\gamma$ -oryzanol at 90°C, but UAE allows to reach the maximum yield  
380 in just one minute and therefore in much shorter times, operating at room temperature.  
381 In their work, Mohammed Danlami et al. [43] and Zhang et al. [44] compared  
382 traditional SE with that of other extraction techniques to extract valuable components  
383 from plants. They affirmed that ultrasound facilitates the extraction of thermally  
384 sensitive compounds enhancing the extraction rate and reducing the extraction  
385 temperature. Khoei and Chekin [44], in their works, extracted RBO using aqueous  
386 extraction and compared the conventional SE with UAE. The two extraction techniques  
387 allowed to obtain very similar RBO yield and they demonstrated that the application of  
388 ultrasound permitted to work at room temperature in a shortened extraction time. The  
389 global production of RBO exceeds 1.7 million tons per year [11]. Although the use of  
390 ultrasound at room temperature leads to a slightly lower oil yield, on such a large annual  
391 production this decrease may not be relevant. Using UAE instead of SE would allow to

392 increase the global production of RBO as the extraction has a shorter duration and it is  
 393 therefore possible to increase the number of annual extractions. The results showed in  
 394 the present work are in accordance with the literature studies cited, the yields obtained  
 395 with the two extraction techniques are very similar, but the UAE seems to be the most  
 396 promising as it allows to reduce the time and energy costs derived from the use of high  
 397 temperatures. An LCA study will be described in the next paragraph to verify if the  
 398 UAE is the most sustainable extraction, even environmentally.

399 **Table 2** Peleg's parameters for SE and UAE. Rate constant  $K_1$ , constant capacity  $K_2$ ,  
 400 initial extraction rate ( $B_0$ ), maximum yield extraction ( $C_e$ ), regression coefficient ( $R^2$ ),  
 401 and the root mean square error (RMSE).

		$K_1$ [min · (g bran / g RBO) · 100 or min · (g bran / mg $\gamma$ -oryzanol)]	$K_2$ [g bran · 100 / g RBO or g bran / mg $\gamma$ -oryzanol]	$B_0$ [g RBO · 100 / min g bran or mg $\gamma$ -oryzanol / min g bran]	$C_e$ [(g RBO / g bran) · 100 or mg $\gamma$ -oryzanol / g bran]	$R^2$	RMSE
<b>SE</b>	RBO	0,1506	0,1324	6,6401	7,5529	0,9302	0,0534
	$\gamma$ -oryzanol	2,7668	2,1712	0,3614	0,4606	0,8919	0,7076
<b>UAE</b>	RBO	0,0175	0,1577	57,1429	6,3412	0,7743	0,6070
	$\gamma$ -oryzanol	0,0442	3,2091	22,6244	0,3116	0,7947	0,0070

402

### 403 3.5 Life cycle assessment

404 After evaluating the technical feasibility of oil extraction from rice bran, the  
 405 environmental sustainability of the different approaches was analyzed through LCA.

406 This analysis aimed to understand if to produce one gram of RBO, it is more  
407 environmentally friendly to heat at 90°C for 15 minutes or to generate ultrasounds  
408 (20kHz) for one minute at room temperature.

409 Fig. 5 shows the results of comparative LCA between SE and UAE in terms of four  
410 impact categories: Climate change (kg CO<sub>2</sub> eq), Ozone depletion (kg CFC-11 eq),  
411 Human toxicity (kg 1,4-DB eq), and Freshwater eutrophication (kg P eq). In each  
412 impact category, the single contribution of the filtration steps, electricity, and  
413 isopropanol to the emission and the total emission of all the two extraction procedures  
414 are shown.

415 Regarding the Climate change impact category, to produce 1 gram of RBO, the SE and  
416 UAE produce 0.206 and 0.156 kg of CO<sub>2</sub> eq, respectively; hence, the application of  
417 UAE allows a reduction of the total impact of 25 %. In this impact category, the most  
418 impacting step for both processes is filtration, which produces a considerable amount of  
419 exhausted rice bran as waste material.

420 The SE process emits to the atmosphere  $1.95 \cdot 10^{-8}$  kg CFC-11 eq while the UAE process  
421 emits only  $1.46 \cdot 10^{-8}$  kg CFC-11 eq reducing the contribution to the Ozone depletion of  
422 25 %. To evaluate if the difference among the two adopted techniques was statistically  
423 significant a test of student was carried out considering  $p < 0.05$ . In this case, the most  
424 significant contribution is due to electricity consumption, and the value is almost the  
425 same in the two treatments. SE and UAE differ in the emissions in the filtration step  
426 because the amount of RBO + isopropanol is higher in the second one, which produces  
427 a minor quantity of exhausted rice bran.

428 Concerning the human toxicity impact category, no significant difference between SE  
429 and UAE has been found, because the solvent employed was the same in both the  
430 extractions.

431 Regarding the freshwater eutrophication impact category, SE and UAE produce  $7.13$   
432  $\cdot 10^{-5}$  and  $1.31 \cdot 10^{-5}$  kg P eq, respectively. Therefore, there is a massive reduction of the  
433 impact of applying ultrasound instead of conventional extraction.

434 In all impact categories, the recycling of isopropanol originates avoided emissions  
435 represented in Fig. 5 by negative bars.

436 For the best of author's knowledge, in the scientific literature LCA studies on oil  
437 extraction from rice bran with SE and UAE techniques were not available.

438 To discuss the achieved results, the comparisons with other studies were performed  
439 considering studies with cradle to gate approach, the product extracted as a functional  
440 unit and midpoint as method to analyse the data coming from life cycle inventory.

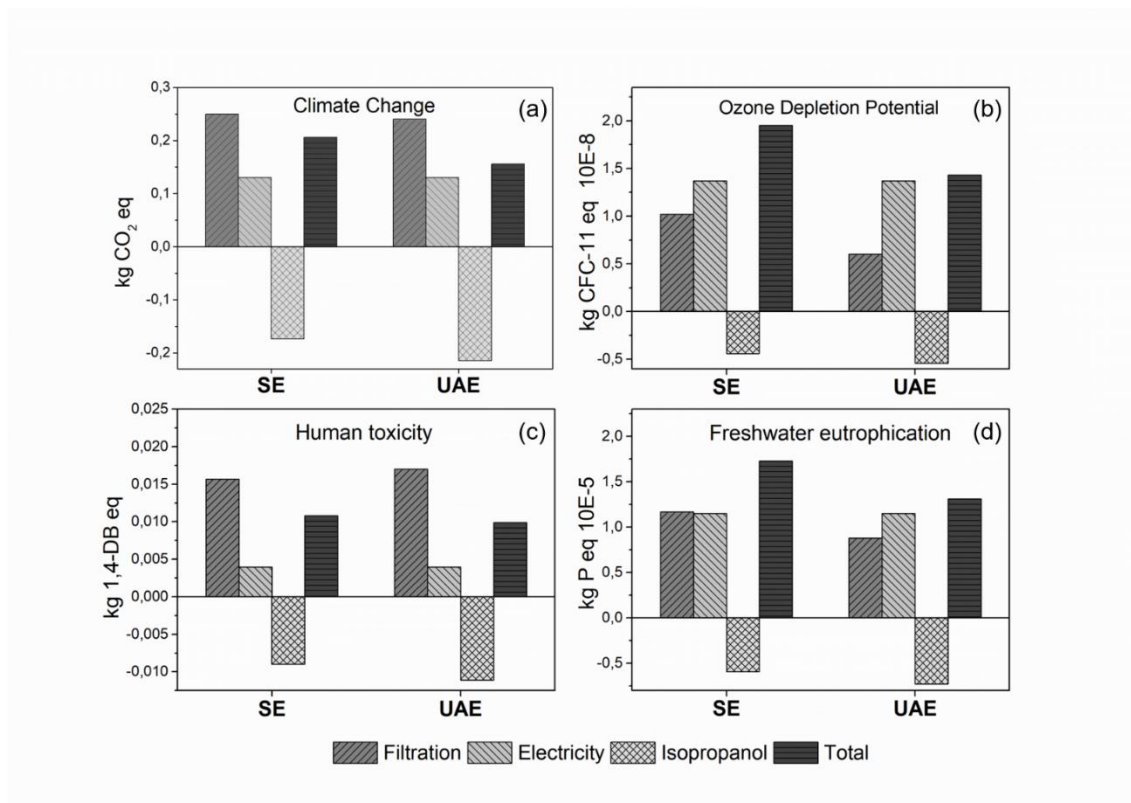
441 Papadaki et al. [45] carried out an LCA study comparing SE, micro-waves, and  
442 ultrasounds to recover the bioactive compounds from microalgae. The authors  
443 demonstrated that among the three extraction techniques, ultrasound was the most  
444 suitable one, since it reached the highest yielding, the lowest economic cost and  
445 medium environmental impacts. Such results agree with the outputs of the present work.

446 Castro-Puyana et al., [46] performed bio-compounds extraction from rosemary plant by  
447 means of green solvent and pressurized hot water extraction. The SE impacts obtained  
448 by the authors were in line with the impacts of SE technique achieved in the present  
449 work.

450 Amiri et al. [47] carried out an LCA about alkaloids extracted from the *Atropa*  
451 *belladonna* by methanol. The global warming potential (GWP) and ozone depletion

452 (OD) reported by the authors were equal to 0.899 kg CO<sub>2</sub> eq and to 0.00015 kg CFC-11  
453 eq, respectively, whereas in the present work the GWP was 0.206 kg CO<sub>2</sub> eq and the  
454 OD was  $1.43 \cdot 10^{-8}$  kg CFC-11 eq. Hence, the present study reached potential impacts  
455 lower than Amiri et al. [45] ones in a range between 77.00%-99.98 %.

456 Barjovenu et al. [48][48] performed an LCA study on polyphenols extraction from  
457 spruce bark, by means of SE using ethanol and UAE. The difference between SE using  
458 ethanol and UAE was 70.00%, whereas in the present study was 24%. The main  
459 difference with the present work is due to the different solvent used. However, the UAE  
460 technique resulted in an environmental impact lower than SE. Thus, the present work  
461 proved that UAE technique is both technically more efficient and environmentally more  
462 sustainable than SE technique. The present study proved the technical feasibility of the  
463 two proposed techniques SE and UAE and the feasible scale up, furthermore the LCA  
464 study, performed considering the data at laboratory scale, underscored the bottleneck of  
465 the processes, which are the filtration steps and the energy consume for both the  
466 techniques. Hence, the recommendation and future prospective are the minimisation of  
467 waste production at filtration step improving the technique and the optimisation of the  
468 energy consume by doing a proper design of the plant.



469

470 **Fig. 5** Comparative LCA results between SE and UAE. Four main impact categories are  
 471 illustrated: Climate change, Ozone depletion, Human toxicity and Freshwater  
 472 eutrophication. Each graph reports the result for SE (on the left) and for UAE (on the  
 473 right); for each extraction process, the contribution of filtration, electricity, isopropanol  
 474 and their sum (total) are reported.

475

#### 476 **4. Conclusions**

477 In the present work, the isopropanol SE of RBO was optimized in terms of temperature  
 478 and bran to solvent ratio. The best RBO yield was obtained at 90°C and 1:9 bran to  
 479 solvent ratio. The results were compared with the RBO yield obtained in a standard  
 480 hexane extraction, demonstrating that isopropanol is suitable to RBO extraction, making  
 481 the substitution of organic and toxic solvents possible. The kinetics of isopropanol SE at

482 the best-operating conditions was evaluated and compared with a room temperature  
483 UAE using a 1:9 bran to solvent ratio. The two extraction techniques produced similar  
484 yields in terms of RBO and  $\gamma$ -oryzanol, but UAE reduced remarkably the extraction  
485 time. A comparative LCA between the two extraction techniques showed that UAE  
486 allows lower the emission contribution to climate change, ozone depletion, and  
487 freshwater eutrophication compared to SE, to produce 1 gram of RBO generating high  
488 yield, operating at room temperature in a very short time, in line with the principles of  
489 green chemistry. To the best of the authors' knowledge, this paper shows for the first  
490 time a comparison between the extraction of RBO with isopropanol at 90°C and with  
491 isopropanol at room temperature assisted by ultrasounds both from a technical and  
492 environmental point of view. The evaluation carried out and the results obtained can be  
493 the basis for new experimental campaigns or to design a scale-up RBO production  
494 plant.

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