

Optimization of microwave-assisted extraction of hydrophilic and lipophilic antioxidants from a surplus tomato crop by response surface methodology

5 José Pinela^{a,b}, M.A. Prieto^{a,c,*}, Maria Filomena Barreiro^d, Ana Maria Carvalho^a, M.
6 Beatriz P.P. Oliveira^b, J.A. Vázquez^e, Isabel C.F.R. Ferreira^{a,*}

⁹ ^a Mountain Research Centre (CIMO), ESA, Polytechnic Institute of Bragan , Campus
10 de Santa Apol nia, 1172, 5301-855 Bragan , Portugal

^b REQUIMTE/LAQV, Faculty of Pharmacy, University of Porto, Rua Jorge Viterbo Ferreira, nº 228, 4050-313 Porto, Portugal

¹³ ^c Nutrition and Bromatology Group, Faculty of Food Science and Technology,
¹⁴ University of Vigo, Ourense Campus, E32004 Ourense, Spain.

^d Laboratory of Separation and Reaction Engineering (LSRE), Associate Laboratory LSRE/LCM, Polytechnic Institute of Bragan , Campus de Santa Apol nia, 1134, 5301-857 Bragan , Portugal

¹⁸ ^e Grupo de Reciclado y Valorización de Materiales Residuales (REVAL), Instituto de
¹⁹ Investigacións Mariñas (IIM-CSIC), r/Eduardo Cabello, 6., Vigo 36208, Galicia, Spain.

* Authors to whom correspondence should be addressed: Isabel C.F.R. Ferreira (e-mail: iferreira@ipb.pt; telephone +351-273-303219; fax +351-273-325405) and M.A. Prieto (e-mail: michaelumangelum@gmail.com; telephone +34 654-694-616)

24 **Abstract**

25 Tomato is the second most important vegetable crop worldwide and a rich source of
26 industrially interesting antioxidants. Hence, the microwave-assisted extraction of
27 hydrophilic (*H*) and lipophilic (*L*) antioxidants from a surplus tomato crop was
28 optimized using response surface methodology. The relevant independent variables
29 were temperature (*T*), extraction time (*t*), ethanol concentration (*Et*) and solid/liquid
30 ratio (*S/L*). The concentration-time response methods of crocin and β -carotene
31 bleaching were applied, since they are suitable *in vitro* assays to evaluate the antioxidant
32 activity of *H* and *L* matrices, respectively. The optimum operating conditions that
33 maximized the extraction were as follows: *t*, 2.25 min; *T*, 149.2 °C; *Et*, 99.1 %; and *S/L*,
34 45.0 g/L for *H* antioxidants; and *t*, 15.4 min; *T*, 60.0 °C; *Et*, 33.0 %; and *S/L*, 15.0 g/L
35 for *L* antioxidants. This industrial approach indicated that surplus tomatoes possess a
36 high content of antioxidants, offering an alternative source for obtaining natural value-
37 added compounds. Additionally, by testing the relationship between the polarity of the
38 extraction solvent and the antioxidant activity of the extracts in *H* and *L* media
39 (polarity-activity relationship), useful information for the study of complex natural
40 extracts containing components with variable degrees of polarity was obtained.

41

42 **Keywords:** *Lycopersicon esculentum*; microwave-assisted extraction; β -carotene/crocin
43 bleaching assay; concentration-time response modelling; response surface methodology

44 **1. Introduction**

45 Tomato (*Lycopersicon esculentum* Mill.) is the second most important vegetable crop
46 worldwide after potato and is consumed either fresh or in the form of processed
47 products. In 2013, about 164 million of tones were produced in the world, having been
48 registered an increase of 2.6 million of tones over 2012 (FAOSTAT, 2015). Apart from
49 the large amounts of solid wastes produced by the processing industry, sometimes there
50 is also a surplus production that leads to glut in the market, distress sale and low profit
51 to the growers (Oliveira, 2006; Sashimatsung et al., 2011). One solution for the problem
52 of this glut may be its sustainable use for the recovery of value-added antioxidant
53 compounds with applications in food, pharmaceutical and cosmeceutical industries. In
54 fact, tomato is a rich source of hydrophilic and lipophilic antioxidants (Barros et al.
55 2012; Pinela et al., 2012). The hydrophilic fraction is constituted mainly by ascorbic
56 acid and soluble phenolic compounds, while the lipophilic fraction contains carotenoids
57 (mostly lycopene), tocopherols, sterols and lipophilic phenolics. Each of these
58 compounds has their own function in the human organism, acting at different locations,
59 but also working conjunctly, having the ability to offer protection against oxidative
60 stress and various degenerative diseases (Carocho and Ferreira, 2013a, 2013b;
61 Friedman, 2013). Besides, according to some reports, antioxidants belonging to the
62 hydrophilic fraction have a far more significant impact on total antioxidant activity than
63 does antioxidants of the lipophilic fraction (García-Valverde et al., 2013; Kotíková et
64 al., 2011).

65 The antioxidant activity can be monitored using a large variety of assays, each one
66 based on a specific mechanism of action, including hydrogen atom transfer, single
67 electron transfer, reducing power, and metal chelation, among others (Carocho and
68 Ferreira, 2013a; Shahidi and Zhong, 2015). For this reason, it is important to understand

69 the mechanisms behind the selected assay for a suitable evaluation of the antioxidant
70 potential. Crocin and β -carotene bleaching reactions are two *in vitro* assays appropriate
71 for the antioxidant activity evaluation of hydrophilic (*H*) and lipophilic (*L*) matrices,
72 respectively, and can provide useful information in the study of complex natural
73 extracts containing components with variable degrees of polarity (Prieto et al., 2013;
74 Prieto and Vázquez, 2014). Both assays are reproducible, especially accurate, and yields
75 a low experimental error (Prieto et al., 2014).

76 To recover antioxidants from plant-based products is necessary to follow suitable
77 extraction methods that ensure and preserve its integrity and bioactivity. That's why the
78 industry is looking for more efficient processes based on enhanced innovation capacity.
79 Among them, microwave-assisted extraction (MAE) has gained significance due to its
80 shortened extraction time, higher extraction rate, reduced solvent consumption and
81 superior product's quality at lower cost (Dahmoune et al., 2015; Gallo et al., 2010),
82 being one of the dominant trends of the "green chemistry" movement (Michel et al.,
83 2011). However, the extraction process efficiency depends on some variables and
84 operating conditions (Bhuyan et al., 2015; Dahmoune et al., 2015), which may not be
85 generalized for all plant materials due to the diverse nature of existing bioactive
86 phytochemicals. Therefore, selection and optimization of variables and operating
87 conditions for the MAE of antioxidants from tomato is necessary.

88 One-factor-at-a-time approaches are commonly used to optimize extraction processes;
89 but it is well-known that optimal operating conditions or interactions between variables
90 cannot be predicted with this methodology. Both problems may be overcome by
91 employing the response surface methodology (RSM), a powerful statistical tool used to
92 predict the optimum experimental conditions to maximize or minimize various
93 independent variables. Indeed, RSM describes the relationship between independent

94 variables and one or more responses, enabling process optimization such as the
95 extraction of bioactive molecules from natural sources with a reduced number of
96 experimental trials.

97 This study aimed at determining the optimal extraction conditions for *H* and *L*
98 antioxidants from a tomato surplus. Four independent variables (temperature, extraction
99 time, ethanol concentration and solid/liquid ratio) were studied and the extraction
100 process was optimized by RSM. The concentration-time response methods of β -
101 carotene and crocin bleaching were applied, which are appropriate for the evaluation of
102 antioxidant properties of *L* and *H* fractions, respectively.

103

104 **2. Material and methods**

105 **2.1. Equipment and reagents**

106 *Equipments*: Biotage Initiator Microwave (Biotage® Initiator⁺, Uppsala, Sweden) using
107 closed high precision glass vials. Multiskan Spectrum Microplate Photometer using 96-
108 well polypropylene microplates.

109 *Reagents*: Linoleic acid (CID 5280450); β -Carotene (CID 5280489); Crocin (CID
110 5281233); 2,2'-Azobis(2-amidinopropane) (AAPH or ABAP, CID 1969). All other
111 chemicals and solvents were of analytical grade and purchased from common sources.
112 Water was treated in a Milli-Q water purification system (Millipore, model A10,
113 Billerica, MA, USA).

114

115 **2.2. Plant material**

116 A common tomato farmers' variety known as "tomate redondo or batateiro" (round
117 tomato), and widely cultivated in rural communities from Miranda do Douro, North-
118 eastern Portugal, was chosen for this study. Surplus tomatoes at the ripe stage were

119 hand-harvested randomly from the middle of six plants, in selected homegardens of two
120 villages in the studied area. The ripening stage was established according to local
121 consumers' criteria based in morphological descriptors such as size, texture, and
122 pericarp colour. Six tomatoes (pericarps without jointed pedicels and seeds) were frozen
123 and lyophilized (Free Zone 4.5, Labconco, Kansas City, MO, USA), reduced to a fine
124 dried powder (20 mesh) using a grinding machine and kept at -20 °C until analysis.

125

126 **2.3. Microwave-assisted extraction of H and L antioxidants**

127 The MAE process was performed using a Biotage Initiator Microwave apparatus in
128 closed vials. The dried powdered samples were extracted at different time (*t*),
129 temperature (*T*), ethanol concentration (*Et*) and solid/liquid ratio (*S/L*) ranging as
130 defined by the RSM design (Fig. 1). The solvent volume was fixed at 20 mL. During
131 extraction, samples were stirred at 600 rpm using a magnetic stirring bar and irradiated
132 at 200 W. After that, the reaction mixture in the closed vial was quickly cooled in the
133 processing chamber and then centrifuged at 6000 rpm for 10 min. The pellet was
134 discarded and the supernatant was carefully collected, evaporated under reduced
135 pressure to remove the solvent and finally re-suspended in distilled water for further
136 analysis. The dry weight (dw) of the suspended solids in the supernatant of each
137 solution was determined to compute the extraction yield (g extract/g samples).

138

139 **2.4. Determination of the concentration-time dependency of L and H antioxidants**

140 β-Carotene (Marco, 1968) and crocin (Bors et al., 1984) methods (BCM and CM,
141 respectively) are widely used to evaluate the antioxidant activity of different matrices.
142 Both *in vitro* assays share some analytical similarities as depicted in the next points.

143

144 2.4.1. Reaction conditions

145 β CM conditions (Prieto et al., 2012): 2 mg of β -carotene (β C, 1 μ M in the final
146 reaction), 0.25 mL of linoleic acid and 2 g of Tween-40 were dissolved in 20 mL of
147 chloroform, vigorously mixed, followed by chloroform evaporation (45 °C/~15 min).
148 To the resulting oily residue were added 300 mL of buffered Mili-Q water (100 mM
149 Briton, pH=6.5) at 45 °C. The absorbance at 470 nm of the prepared reagent was ~1.40.
150 CM conditions (Prieto et al., 2015): 4 mg of crocin (Cr, 100 μ M in the final reaction)
151 and 75 mg of AAPH (7.68 mM in the final reaction) were dissolved in 30 mL of a 100
152 mM Briton buffer, pH=5.5, in Mili-Q water. The absorbance at 450 nm of the prepared
153 reagent was ~1.40.

154

155 2.4.2. Procedure

156 The procedure was performed by adding 50 μ L of sample extract and 250 μ L of reagent
157 into the wells (350 μ L) of the microplate. The microplate-reader was programmed at
158 intervals of 3, 5 and 10 min (initiation, propagation and asymptotic phases), during a
159 period of 200 min (total of 30 measures). The sample extracts were analyzed kinetically
160 for eight different concentrations obtained by serial dilution (1/1, 1/2, 1/4, 1/8, 1/16,
161 1/32, 1/64 and the control) in distilled water.

162

163 2.4.3. Quantification

164 The area under the curve (AUC) (Eq. 1), computed by any numerical integration method
165 such as the trapezoidal rule, proved to be a highly robust criterion, able to summarize in
166 a single and direct value the global feature of any kinetic profile.

$$AUC = \frac{R_1 \Delta t_1}{2} + \sum_{i=2}^{i=n-1} R_i \Delta t_i + \frac{R_n \Delta t_n}{2} \quad (1)$$

167 where i is the number of data measured over time t , R_i are the responses along an
168 arbitrary time series, and Δt is the interval of each measurement.

169 The AUC of a concentration-response of an antioxidant was normalized against to the
170 AUC obtained with the control, leading to the formulation of the relative area units or
171 protected substrate (\bar{P}) in percentage (Eq. 2), as defined similarly by other authors
172 (Dávalos, 2004; Huang et al., 2002; Naguib, 2000) for antioxidant responses.

$$\bar{P}(A) = S_0 \left(\frac{AUC_C - AUC_A}{AUC_C} \right) \frac{100}{S_0} \quad (2)$$

173 where AUC_C and AUC_A are the area units corresponding to the kinetic profiles found in
174 the absence (control, C) and presence of an antioxidant concentration A , respectively,
175 and S_0 is the initial substrate in the reaction (for the CM, the substrate is equivalent to
176 100 μM of Cr, and for the β CM it is equivalent to 1 μM of β C).

177 The relationship in Eq. (2) establishes that AUC_C (control) is also the maximum
178 response achievable; consequently the values obtained were also standardized in
179 percentage. In addition, by normalizing the response, the results obtained are less
180 dependent on the experimental conditions, which, in practice, is one of the common
181 problems when analyzing the efficacy of response factors.

182 The asymptotic variation of \bar{P} as function of an antioxidant compound suggests that
183 some radical-generating property of the system can be saturated (Gieseg and Esterbauer,
184 1994). This type of concentration-response patterns should behave in a similarly
185 accumulative way with a number of different antioxidant compounds found in the
186 extract material. Therefore, in general, this patterns can be adjusted by a group of
187 mathematical expressions (mechanistic or not) that translates the pattern of the response
188 into parameters that allow to deduce the meaning and/or quantify the effect of the
189 dependent variable in a simple and global mode. Previous researchers discussed the

190 applicability of different mathematical expressions (Prieto et al., 2014); therefore,
 191 following their views, the Weibull cumulative distribution function was selected
 192 (Weibull and Sweden, 1951). Thus, the variation of \bar{P} as function of increasing
 193 concentrations of an antioxidant (A) can be described satisfactorily using the Weibull
 194 model rearranged for our own purposes as follows in Eq. (3).

$$\bar{P}(t, A) = P_m \left\{ 1 - \exp \left[-\ln(2)^{1-a} \left(\frac{2V_m}{P_m a} A \right)^a \right] \right\} \quad (3)$$

195 The parameter P_m is the averaged maximum protected substrate, asymptotic value of the
 196 response (% μ M of β C or Cr), which is specific of each A agent. The parameter V_m
 197 corresponds to the average amount of protected molecules per gram of extracted
 198 material (% μ M of protected substrate/g extract). The parameter a is a shape parameter
 199 related to the slope that can produce potential profiles ($a < 1$), first order kinetic ones
 200 ($a = 1$) and a variety of sigmoidal profiles ($a > 1$).

201 In addition, the concentration needed to reach 50% of the maximum protective effect
 202 (the so called IC_{50}) can be determined according to Eq. (4).

$$IC_{50} = \frac{Ka \ln 2}{2V_m} ; \text{ therefore } \bar{P}(t, A) = P_m \left\{ 1 - \exp \left[-\ln(2)(t/IC_{50})^a \right] \right\} \quad (4)$$

203 where IC_{50} is the concentration producing the half-maximal response and all other
 204 notations remain with the same meaning as above.

205

206 2.5. Response surface methodology

207 The RSM family designs are used for modelling and analyzing problems in which a
 208 response of interest is influenced by a set of variables. RSM was applied to optimize the
 209 MAE process with the purpose of finding the favourable processing conditions that
 210 would result in a higher extraction rate of H or L antioxidants.

211

212 *2.5.1. Response criteria for evaluating the antioxidant capacity*

213 The responses used in the RSM analysis were based in the numerical values of the
214 parametric coefficients P_m , V_m and IC_{50} of Eqs. (3) and (4). The information provided
215 by the combination of the values of the three response criteria represents a robust tool to
216 compare the activity of different antioxidant agents based on the parametric
217 concentration-time estimations.

218

219 *2.5.2. Preliminary experiments*

220 Preliminary single-factor experiments were conducted in order to select the significant
221 variables and/or collateral factors in extraction process and to determine the preliminary
222 range of the optimum level of each factor for an appropriate experimental RSM design.

223 In this primary screening trial, the following variables and factors were considered:

- 224 - Internal independent variables of the microwave equipment: Pressure (1-30 bar),
225 stirring rate (0-1000 rpm), microwave power (0-400 W), temperature (40-300 °C)
226 and extraction time (no limits).
- 227 - Internal factors of the instrument software: Absorption level (*very low, low, normal,*
228 *high, or very high*), fixed hold time (if *on* the time countdown starts when the target
229 temperature or pressure is reached, *i.e.*, the initial time taken to reach the set
230 conditions is not included in the heating time; if *off* the time countdown starts when
231 the heating starts), cooling (*on* or *off*), pre-stirring (during the fixed hold time, if
232 selected; *on* or *off*) and vial type (2-5 mL or 10-20 mL).
- 233 - External independent variables and factors: Solid/liquid ratio and ethanol
234 concentration. The type of solvent used in the extraction.

235

236 2.5.3. Experimental design

237 From the preliminary study, the independent variables X_1 (extraction time, min), X_2
238 (temperature, °C), X_3 (ethanol concentration, %) and X_4 (solid/liquid ratio, g/L) were
239 selected. Then, the combined effects of these variables on the extraction yield of H and
240 L antioxidant were studied using *central composite design* as proposed by Box et al.
241 (1957). In this design, the points of experiments are generated on a sphere around the
242 centre point. The centre point is supposed to be an optimum position for the response
243 and is repeated in order to maximize the prediction precision (Box and Hunter, 2005).
244 This design also requires five levels for each factor. The number of repetitions n_0 of the
245 centre point is calculated by the formulas present in Eq. (5) for k factors based on the
246 uniform precision.

$$\gamma = \frac{(k+3) + \sqrt{9k^2 + 14k - 7}}{4(k+2)}; \text{ where: } n_0 = \text{floor} \left(\gamma \left(\sqrt{2^k} + 2 \right)^2 - 2^k - 2k \right) \quad (5)$$

247 where *floor* designates the highest integer value smaller than the argument. The number
248 of experiments n for k factors is given as:

$$n = 2^k + 2k + 1 \quad (6)$$

249 Experimental runs were randomized, to minimize the effects of unexpected variability
250 in the observed responses. Independent variables coded values and natural ones of the
251 factorial design are coded and decoded by the expressions in Eq. (7).

$$v_c = (v_n - v_0) / \Delta v_n \quad \text{and} \quad v_n = v_0 + \Delta v_n \times v_c \quad (7)$$

252 where v_n and v_c is the natural (n) and coded (c) value in the centre of the experimental
253 domain, v_0 is the initial value and Δv_n is the increment of v_n for unit of v_c .
254

255 2.5.4. Box-Behnken mathematical model

256 Response surface models were fitted by means of least-squares calculation using the
257 following Box-Behnken equation:

$$Y = b_0 + \sum_{i=1}^n b_i X_i + \sum_{\substack{i=1 \\ j>i}}^{n-1} \sum_{j=2}^n b_{ij} X_i X_j + \sum_{i=1}^n b_{ii} X_i^2 \quad (8)$$

258 where Y is the dependent variable (response variable) to be modelled, X_i and X_j define
259 the independent variables, b_0 is the constant coefficient, b_i is the coefficient of linear
260 effect, b_{ij} is the coefficient of interaction effect, b_{ii} the coefficients of quadratic effect
261 and n is the number of variables. As pointed out, three different response formats based
262 in the parametric estimations (P_m , V_m and IC_{50}) of Eqs. (3) and (4) were used as the
263 dependent variable for each H and L antioxidant analytical reaction (Y_{Pm}^H ; $Y_{V\tau}^H$; $Y_{IC_{50}}^H$;
264 Y_{Pm}^L ; $Y_{V\tau}^L$; and $Y_{IC_{50}}^L$).

265

266 **2.6. Numerical methods and statistical analysis**

267 All fitting procedures, coefficient estimates and statistical calculations were performed
268 on a Microsoft Excel spreadsheet. Fitting and statistical analysis of the experimental
269 results to the proposed equations were carried out in four phases:

- 270 1) Coefficients determination: Parametric estimates were obtained by minimization of
271 the sum of the quadratic differences between observed and model-predicted values,
272 using the nonlinear least-squares (quasi-Newton) method provided by the macro
273 *Solver* in *Microsoft Excel* 2003 (Kemmer and Keller, 2010), which allows a quick
274 testing of a hypotheses and its consequences (Murado and Prieto, 2013).
- 275 2) Coefficients significance: The determination of the parametric confidence intervals
276 were calculated using the ‘*SolverAid*’ (Prikler, 2009). The model was simplified by
277 dropping terms, which were not statistically significant p -value > 0.05 .

278 3) Model consistency: The Fisher *F*-test ($\alpha=0.05$) was used to determine whether the
279 constructed models were adequate to describe the observed data (Shi and Tsai,
280 2002).

281 4) Other statistical assessment criteria: To re-check the uniformity of the model the
282 following criteria were applied: a) The ‘SolverStat’ macro was used for the
283 assessment of the parameter and model prediction uncertainties (Comuzzi et al.,
284 2003); b) The R^2 was interpreted as the proportion of variability of the dependent
285 variable explained by the model; c) The adjusted coefficient of determination (R^2_{adj})
286 was a correction to R^2 taking into account the number of variables used in the
287 model; d) Bias and accuracy factors of all equations were calculated to evaluate the
288 fittings to experimental data, such as the Mean Squared Error (MSE), the Root
289 Mean Square of the Errors (RMSE) and the Mean Absolute Percentage Error
290 (MAPE); e) The Durbin-Watson coefficient (DW) was used to check if the residuals
291 of the model are not autocorrelated; and f) The analysis of variance table (ANOVA)
292 was used to evaluate the explanatory power of the variables.

293

294 **3. Results and discussion**

295 **3.1. Preliminary study**

296 As listed in the material and methods section, the MAE efficiency may be affected by
297 five internal (pressure, stirring rate, microwave power, extraction time, and temperature)
298 and two external (solid/liquid ratio and ethanol concentration) independent variables
299 and five instrument/software factors (absorption level, fixed hold time, vial type,
300 cooling option, and solvent type). Although there are previous research examples
301 (Bhuyan et al., 2015; Dahmoune et al., 2015), the results are not generalizable for all
302 plant materials due to the diverse nature of existing bioactive phytochemicals.

303 Therefore, this preliminary study allowed screening of appropriate independent
304 variables and determining their optimum experimental domain for an appropriate
305 experimental RSM design. Variables/factors were investigated by testing a broad range,
306 keeping other ones constant and analyzing their antioxidant responses.

307 - Type of extracting solvent is the key for separating *H* and *L* compounds. Water is
308 the polar solvent with greater interest in biological processes than any other solvent.
309 Ethanol has a polar hydroxyl group and dissolves many ionic compounds, but also
310 has a non-polar end, which will contribute to dissolve non-polar substances. In this
311 study, binary interactions of ethanol-water mixtures were selected due to their
312 straight *H* affinity, but also because ethanol increases the *L* character of the aqueous-
313 ethanolic mixture. All the tested ethanol concentrations give rise to significant
314 differences; thus, the range from 0 to 100 % was selected.

315 - Pressure and temperature were correlated. The selected irradiation power was
316 applied in the early stage of the extraction process to reach the selected
317 temperature/pressure in a short period of time. After that, it was automatically
318 applied in less intensity (estimated by the microwave system) to keep constant the
319 solution temperature/pressure. In consequence, the microwaves power was set at
320 200 W and the temperature was selected as the main controlled variable, since
321 relevant differences were found within the range 60 to 180 °C.

322 - Lower solid/liquid ratios can lead to a more efficient dissolution of constituents, but
323 also to a waste of solvent. At an industrial scale, higher ratios are desirable since it
324 is important to maximize the extraction yield (thus productivity) with a minimal
325 solvent consumption (more sustainable process). Significant differences were found
326 for all tested ratios in this preliminary study, being the range from 5 to 45 g/L
327 selected for RSM analysis.

328 - The temperature caused strong decomposition phases of antioxidants for extraction
329 times higher than 20 min, but this effect depended on the other variables that
330 remained constant. Therefore, extraction times ranging from 0 to 20 min were
331 selected.

332 - The cooling option showed relevant effects. It is used at the end of the MAE process
333 to cool the sample. When *off*, the cooling time of the solution after processing was
334 longer, affecting the extraction of antioxidants. Therefore, it was used *on* to quickly
335 cool the solution and stop faster the extraction process, making the process more
336 accurate.

337 - When testing the effects of other factors, such as the absorption level, fixed hold
338 time and vial type, no significant changes were found. Therefore, a *normal*
339 absorption level and vials of 10-20 mL were selected for further analysis. The fixed
340 hold time was turned *off*, since the initial time taken to reach the set temperature or
341 pressure was negligible.

342 Therefore, the RSM experiment was designed based on these preliminary results, using
343 five variation levels of extraction time (0-20 min), temperature (60-180 °C), ethanol
344 concentration (0-100%) and solid/liquid ratio (5-45 g/L) as independent variables to
345 optimize efficiently the MAE process, regarding the *H* and *L* antioxidant properties of
346 the extracts. The coded values and their natural values are presented in Fig. 2. Note that,
347 for simplification reasons, the RSM design reduces the number of experimental trials.
348 When studying 5 levels of 4 independent variables, the response would imply 625
349 possible combinations, but using RSM the experiment could be solved in 25
350 independent combinations and 7 replicates at the centre of the experimental domain.

351

352 **3.2. Concentration-time antioxidant responses for RSM**

353 Several reviews have discussed the numerous *in vitro* methods developed to evaluate
354 the antioxidant activity of plant extracts and their controversial aspects regarding
355 differences in the generated radicals, variables (mainly pH and temperature), reagents,
356 and quantification procedures (Carocho and Ferreira, 2013a; Frankel and Meyer, 2000;
357 Jiménez-Escrig et al., 2000). However, when determining the bioactivity of a sample,
358 the final activity response also depends on the degree of polarity of the intermediate
359 reaction components of the applied method. For example, when evaluating the
360 bioactivity of extracts obtained with *H* and *L* extraction solvents using the oxidative
361 hemolysis inhibition assay (OxHLIA) (method in which the thermal decomposition of
362 AAPH generates *H* radicals and the lipid peroxidation of the erythrocytes membranes
363 generates *L* radicals (Niki et al., 1988)), no clear conclusions can be made because both
364 *H* or *L* antioxidants delay the hemolysis and, therefore, produce an antioxidant response.
365 To the best of our knowledge only few articles have addressed the *H* and *L* intermediate
366 components activity (Arnao et al., 2001; Prieto et al., 2013; Prior et al., 2003).
367 Therefore, the lack of selective methods to differentiate the activity of *H* and *L*
368 antioxidants is a current issue in the evaluation of antioxidant responses that need to be
369 outlined in the following years.

370 In order to reduce the variability of experimental conditions, allowing meaningful
371 comparisons, and to quantify the power of the antioxidants in function of the degree of
372 polarity, the response models of CM and β CM were selected because they provide a
373 microsystem for *H* and *L* oxidation processes, respectively (Prieto et al., 2013). The β C
374 is an *L* oxidizable substrate that can join the system of lipid micelles in which the
375 oxidation reaction is accomplished. The method is especially sensitive to antioxidants in
376 a lipidic environment, producing a very low response to *H* antioxidants, even to

377 powerful ones. In turn, Cr is an *H* oxidizable substrate and *L* antioxidants produce very
378 low responses in the reaction system.

379 Fig. 3 and Fig. 4 show an illustration of the antioxidant responses obtained for the
380 tomato extracts produced under the experimental RSM design presented in Fig. 2 for
381 each *H* and *L* antioxidant reaction (CM and β CM), respectively. In both figures, two
382 well differentiated sections are presented at the left- and right-hand sides, showing the
383 visual variable distribution of the 25 genuine combinations. The left-hand side shows
384 the combinations of the concentration-time responses of seven serial dilutions (\bigcirc :1/1,
385 \blacktriangle : 1/2, \triangle : 1/4, \blacksquare : 1/8, \square : 1/16, \blacklozenge : 1/32, \lozenge : 1/64) and the control (\bullet) for the
386 remaining substrates (% μ M Cr and β C). Meanwhile, the right-hand side shows the
387 concentration-time transformation into the concentration-response values of the *AUC*
388 computed by the numerical integration method in Eq. (1) and standardized in percentage
389 of protected substrate (Cr or β C) by Eq. (2). The dots (\bullet) are the raw values and lines
390 (—) the fitted responses to the mathematical model of Eq. (3) or (4). The parametric
391 fitting values of Eqs. (3) and (4) are presented in Table 1. The estimated numerical
392 values of P_m , V_m and IC_{50} were the three meaningful ways considered to evaluate the
393 effectiveness of the antioxidant response by RSM.

394

395 **3.3. Development of the theoretical response surface models and statistical
396 verification**

397 Table 1 shows the results of the parametric fitting coefficients (data presented in Fig. 3
398 and Fig. 4) for each *H* and *L* reaction obtained after running 32 trials (25 genuine
399 combinations and 7 replicates) following the experimental RSM design. Estimated
400 coefficient values of Eq. (8), parametric intervals and numerical statistical criteria are
401 shown in Table 2, for each coefficient used as response criteria and for *H* and *L*

402 reactions. The coefficients that showed effects with *p-values* higher than 0.05 are not
 403 significant (*ns*) at the 95% confidence level and consequently were discarded for model
 404 development.

405 Mathematical models were built through nonlinear least-squares estimations based on
 406 the coded experimental plan and the response results, obtaining the following second-
 407 order polynomial Eq. (8):

408 when the hydrophilic *CM* was considered:

$$Y_{Pm}^H = 30.5 + 8.7x_3 + 6.9x_4 + 7.3x_1^2 + 13.1x_4^2 + 13.1x_1x_2 - 4.5x_2x_3 \quad (9)$$

$$Y_{V\tau}^H = 26.5 - 8.1x_1 - 17.2x_3 - 6.9x_2^2 + 9.1x_3^2 + 7.0x_4^2 \quad (10)$$

$$Y_{IC_{50}}^H = 0.40 + 0.04x_2 + 0.27x_3 + 0.05x_2^2 + 0.05x_3^2 + 0.16x_1x_2 - 0.05x_2x_3 \quad (11)$$

409 when the lipophilic β *CM* was considered:

$$Y_{Pm}^L = 96.1 + 8.1x_1 - 7.1x_2 + 5.1x_3 - 5.6x_4 - 10.7x_1^2 - 9.8x_2^2 - 8.6x_3^2 - 11.8x_4^2 - 7.7x_1x_2 + 13.1x_2x_3 \quad (12)$$

$$Y_{V\tau}^L = 13.1 - 2.9x_1 - 3.2x_3 - 2.0x_4 + 5.9x_2^2 + 2.3x_1x_3 + 1.6x_1x_4 - 1.7x_2x_3 + 6.2x_2x_4 \quad (13)$$

$$Y_{IC_{50}}^L = 2.97 + 0.76x_3 + 0.31x_4 - 0.24x_1^2 - 0.07x_3^2 - 0.22x_4^2 - 0.24x_1x_4 + 0.43x_2x_3 + 0.27x_3x_4 \quad (14)$$

410 where X_1 (extraction time), X_2 (temperature), X_3 (ethanol concentration), X_4 (solid/liquid
 411 ratio), Y is the response, sub-indices indicates the coefficient criteria (P_m , V_m and IC_{50})
 412 used as responses for RSM and super-indices H and L accounts for the H (CM) and L
 413 (β CM) reactions.

414 The multivariable characterization of the Box-Behnken second-order polynomial model
 415 is especially robust, minimizing the experimental errors, allowing explain the utmost of
 416 the results. In addition, once a model is designed, if the experimental data obtained do
 417 not span the full range and some of them fail to provide information about one or more
 418 of the parameters of the equation, the multivariable application describes simply and

419 accurately all the areas. As explained, not all the parameters of Eq. (8) were used for
420 building the model, since some terms were non-significant (Table 2). Model coefficients
421 obtained are empirical and cannot be associated with physical or chemical significance.
422 However, they are useful to predict the results of untested operation conditions (Ranic
423 et al., 2014). The sign of the effect marks the performance of the response. In this way,
424 when a factor has a positive effect, the response is higher at the high level and when a
425 factor has a negative effect, the response is lower at high level. The higher the absolute
426 value of a coefficient, the more important the weight of the corresponding variable.
427 Based in the mathematical expressions, it was found that the responses in the *L*
428 environment were much more complex than those found in the *H* one.
429 The statistic lack of fit, used to test the adequacy of the obtained models, demonstrated
430 that no considerable improvement was achieved by the inclusion of the statistically non-
431 significant effects (Table 2). This was also verified by the high R^2 and R^2_{adj} values
432 indicating the percentage of variability of each response that is explained by the model
433 (Table 2). The distribution of residuals always randomly scattered around zero and
434 grouped data and autocorrelations were not observed. This means that these models are
435 workable and can be applied in the subsequent prediction and optimization stages.
436 Finally, the analysis of variance (ANOVA) was computed for the regression equations.
437 The lack of fit was used to verify the adequacy of the model and was not significant (p
438 > 0.05), indicating that Eqs. (12) to (14) adequately fit the experimental data.

439

440 **3.4. Effect of *Et* and *T* variables as representative case of the typical *H* and *L*
441 trends**

442 The three response criteria (P_m , V_m and IC_{50}) characterize singular features of the
443 response. Previous to the complete analysis of the *H* and *L* antioxidant extraction trends,

444 the information provided by each parametric response criteria, which were used in the
445 RSM design, was individually analyzed. As an illustrative case study, it was selected
446 the effect of the variables Et and T , meanwhile the variables t and S/L were positioned at
447 the centre of their experimental domain ($t=10$ min and $S/L=25$ g/L). Graphical 3D
448 representations are displayed in Fig. 5 and the parametric fitting values are present in
449 Table 2. In general, it can be observed that the H and L antioxidant activity of the
450 tomato extracts have opposite trends for Et and T . In more specific terms, for each
451 criterion it can be concluded that:

- 452 a) The parameter P_m of Eq. (3) shows the maximum specific capability of the
453 antioxidant agent to protect the substrate (% μM of Cr or βC) and, the higher the P_m
454 value, the more powerful the protective capability of the antioxidant. In general, we
455 can speculate that the more complex the content in antioxidant molecules in the
456 extract (which act at different H or L oxidation levels), higher the parameter P_m .
457 These types of extracts are usually obtained with longer extraction times. The
458 conditions that favour the H activity of the P_m value were at high ranges (\uparrow) of Et
459 and low ranges (\downarrow) of T or, in a much less active manner, at $\downarrow Et$ and $\uparrow T$. In contrast,
460 the L activity was found at intermediate ranges (\leftrightarrow) of Et and T , leading to a clear
461 optimum at 50 % Et and 120 °C. The inversion effect of the polarity-activity
462 relationship proposed by the polar paradox theory is visible in these results (Porter,
463 1993). Actually, the speculated effect of the non-polar end of ethanol on the activity
464 of the tomato extracts was not observed; they even showed an improvement of the H
465 antioxidant activity at $\uparrow Et$, while the L antioxidant activity decreased sharply at
466 both ends of Et .
- 467 b) The parameter V_m of Eq. (3) corresponds to the average amount of protected
468 molecules of Cr or βC per gram of extracted material (% μM of protected

469 substrate/g extract), which is a value of maximal predictability. The higher the V_m
470 value, the more powerful the antioxidant. There are a diverse number of compounds
471 that would present a high specific protection, but only few would be present in an
472 enough amount to show its activity. Therefore, the highest values should appear
473 when an extraction peak of an antioxidant with a high specific protection would be
474 found. The conditions that maximize the V_m response of the H activity were at $\downarrow Et$
475 and $\leftrightarrow T$; while for the L activity were found at $\downarrow Et$ and $\uparrow T$. The effect of the
476 inversion of the polar activity on the optimal response was not as evident as for the
477 parameter P_m , but the opposite trends remain present as can be seen in each 3D
478 surface.

479 c) The parameter IC_{50} of Eq. (4) provides directly the classical IC_{50} (g of extract),
480 which will effectively summarize all effects of the other two responses. It provides
481 the amount of extract needed to achieve a very specific response (50%). The lower
482 the IC_{50} value, the more powerful the antioxidant. The lowest values should be
483 found in an intermediate position between those speculated in the previous criteria.
484 For the IC_{50} criteria, the conditions that maximize the response for the H antioxidant
485 activity were at $\downarrow Et$ and $\downarrow T$, while for the L activity were at $\downarrow Et$ and $\uparrow T$.

486 The data published in literature often focus on only one response parameter, but each of
487 them describes different intrinsic characteristics of the response. The information
488 provided by the combination of the three values represents a robust tool to compare the
489 activities of different antioxidant agents based on the parametric concentration-time
490 estimations. By analyzing all parametric nonlinear values for the experimental RSM
491 design, a more rigorous evaluation of the extraction efficiency of H and L antioxidants
492 is accomplished.

493

494 **3.5. Nonlinear relationship between extraction solvent polarity and antioxidant**
495 **activity**

496 Matrix combination of the 3D responses for the *H* and *L* environmental reactions
497 obtained for the P_m , V_m and IC_{50} are presented in Fig. 6, Fig. 7 and Fig. 8, respectively.
498 In addition, a simplified way to present the results in a 2D format for all responses is
499 presented in Fig. S1 of the supplementary material. Eqs. (12) to (14) were used to
500 simulate the surfaces. In each graphical illustration, the top diagonal part presents the
501 response surfaces for *L* reactions and the bottom diagonal part presents the response
502 surfaces for *H* reactions. The variables excluded in each 3D graph were positioned at
503 the centre of their experimental domain ($t=10$ min; $T=120$ °C; $Et=50$ %; and $S/L=25$
504 g/L).

505 In general, the inversion effect of the polarity-activity relationship was observed in
506 almost all responses. The effects accounted between the *t*, *T* and *S/L* variables would
507 describe the conditions that optimize the *H* and *L* antioxidant responses of the tomato
508 extracts. However, the variable *Et* did not perform as theoretically expect. Such
509 fuzziness between the polarity of extraction solvent and the antioxidant activity of the
510 extracts in *H* and *L* environments (the so-called polarity-activity relationship) was found
511 interesting enough to be considered. Generally, the extraction ability of solvents can be
512 grouped in three main types: non-polar, polar aprotic and polar protic solvents
513 (Huffman et al., 2012; Kislik, 2012). The choice of the extracting solvent is the first
514 crucial step towards the optimization of any extraction method (Sultana et al., 2009),
515 which has a strong impact on the type of molecules that would be separated. In turn,
516 antioxidants are classified into two broad divisions (Arnao et al., 2001), depending on
517 whether they are soluble in water (*H*, such as ascorbic acid) or in lipids (*L*, such as α -
518 tocopherol). When performing an extraction, it is well known that *L* antioxidant

519 molecules are mostly extracted in non-polar solvents (*i.e.*, *n*-hexane) and *H* antioxidant
520 molecules in polar ones (*i.e.*, water), according to the “like dissolves like” principle, as
521 confirmed by several authors that separated effectively the molecular *H* and *L* character
522 of molecules by applying different solvent combinations in conjunction with different
523 extraction procedures (Watanabe et al., 2014). However, based on the achieved results,
524 the separation of extracts according to their molecular polarity character does not
525 guarantee that their polar or non-polar target activity can be separated as well. As stated
526 before (Prieto et al., 2013), when testing the activity of *H* and *L* antioxidant extracts
527 (hexane and methanol solvents, respectively), it was confirmed that *H* and *L* antioxidant
528 extracts, normally in a much lesser extent, remain active in the opposite environment. In
529 addition to that complex scenery, there are amphiphilic molecules presenting an affinity
530 with solvents of various polarities (Taresco et al., 2015).

531 Thus, if their polarity-activity is not totally related with their distribution in the
532 extracting solvents as defined by the polarity index (*i.e.*, dielectric constant), we may be
533 using extracts for *L* environments (*i.e.* oils) with a high content in molecules with a *H*
534 antioxidant activity and vice versa. Nonetheless, it is recognized that antioxidants with a
535 clear *H* and *L* character can cause the opposite effect when applied in the opposite
536 environment (*i.e.*, ascorbic acid can initiate lipid oxidation in conjunction with metal
537 cations) (Zhang and Omaye, 2001). Actually, according to the polar paradox theory
538 (Porter, 1993), polar antioxidants are more effective in less polar media, while non-
539 polar or amphiphilic antioxidants tend to be more effective in a media of relatively
540 higher polarity. The higher efficiency of *L* antioxidants in oil-in-water emulsions would
541 be due to their tendency to concentrate at the interfacial membrane where the oxidation
542 is supposed to occur, while more *H* antioxidants would tend to segregate into the

543 aqueous phase where they would be much less effective (Frankel et al., 1994). Our
544 results support this phenomenon.

545 A possible hypothetical foundation behind the mechanisms that caused this effect could
546 be the microwave absorbing properties of the solvent (Dahmoune et al., 2015). Polar
547 molecules strongly absorb microwave energy because of the permanent dipole moment,
548 and the degree of absorption increases with the dielectric constant. A simple comparison
549 between water and ethanol shows that ethanol has a lesser ability to obstruct the
550 microwaves as they pass through, but has a greater ability to dissipate the microwave
551 energy into heat. This strong absorption provides an increase of the temperature inside
552 the sample, leading to the rupture of cells by the *in situ* water. In some cases it can
553 promote the degradation of the target antioxidants and, in other cases, can increase the
554 diffusivity of the target antioxidants in the matrix.

555 Knowing all that, when describing the antioxidant activity of components of a complex
556 natural extract as a function of the degree of polarity, scientific studies typically involve
557 a first extraction step with solvents with different polarity index, followed by testing
558 their activity by different analytical procedures. However, such a link between polarity-
559 activity cannot be straightforward performed and further analyses are need. In the
560 literature there are few reports addressing the previously mentioned associated issues
561 (Jayasinghe et al., 2013; Li et al., 2015). However, it would be interesting to perform
562 studies considering the following issues: a) a well-defined group of *in vitro* methods
563 that could separate the polar activity of compounds in *H* and *L* antioxidants; b) a
564 representative set of natural materials sources extracted with a set of solvents
565 demonstratives of the different polarity index; c) a complex optimization of variable
566 conditions that affect the extraction of *H* and *L* antioxidants to ensure and preserve its
567 integrity and bioactivity; and d) clear target applications with a marked *H* and *L*

568 character to prove in an *in vivo* form, whether or not the relation between the aspects
569 stated in a), b) and c) are validated. The combination of all these requirements seems to
570 be a labour-intensive approach, being out the context of this work.

571

572 **3.6. Optimal extraction conditions for *H* and *L* antioxidants**

573 The fitting results (Table 2) obtained by applying Eq. (8) to all the response criteria (P_m ,
574 V_m and IC_{50}) are presented in Eqs. (9) to (11) for the *H* reaction and in Eqs. (12) to (14)
575 for the *L* reaction. By finding the partial derivatives of these regression equations,
576 equating them to zero (Table S1 of the supplementary material) and solving the
577 equations system, the coded values that optimize the response criteria were obtained.
578 Then, the coded variable values were introduced in the original Eqs. (9)-(14) and the
579 optimal response values were found. Finally, by decoding the coded values, the
580 conditions that maximize the response were transformed into natural values.

581 The operating conditions that maximize the extraction of the tomato antioxidants and
582 the optimal response values are presented in Table 3, for each parametric estimation
583 criteria (P_m , V_m and IC_{50}) and analytical reaction (*H* and *L*). For *H* antioxidants, the
584 optimal conditions for P_m were at 180.0 °C, with 56.8 % ethanol and 45.0 g/L of
585 sample, during 18.7 min; for V_m were at 120.0 °C, with 0.0 % ethanol and 5.0 g/L of
586 sample, during 2.5 min; and for IC_{50} were at 90.0 °C, with 44.0 % ethanol and 17.0 g/L
587 of sample, during 14.5 min; and for *L* antioxidants, the optimal conditions for P_m were
588 at 93.6 °C, with 44.0 % ethanol and 21.3 g/L of sample, during 13.4 min; for V_m were at
589 180.0 °C, with 100 % ethanol and 5.0 g/L of sample, during 2.2 min; and for IC_{50} were
590 at 169.1 °C, with 91.7 % ethanol and 10.9 g/L of sample, during 2.6 min. Optimal
591 extraction conditions based on all the response criteria (P_m , V_m and IC_{50}) were also
592 determined for *H* and *L* antioxidants. Based on these values, it was found that the

593 extraction of *L* antioxidants demands a longer *t* (15.4 min) but a lower *T* (60.0 °C), *Et*
594 (33.0 %) and *S/L* (15.0 g/L), comparing to the operating conditions outlined for *H*
595 antioxidants (*i.e.*, *t*, 2.25 min; *T*, 149.2 °C; *Et*, 99.1 %; and *S/L*, 45.0 g/L). These
596 intermediate extraction conditions, and others that were optimized for each response
597 criteria (P_m , V_m and IC_{50}) of both *H* and *L* antioxidants, were depicted using a simplex
598 method tool to solve linear problem. Restrictions were made to the variable coded
599 values that did not allowed the set of equations consider unnatural conditions (*i.e.*,
600 lower times than 0). Additionally, optimal extraction conditions for both *H* and *L*
601 antioxidants based on all the response criteria were determined (*i.e.*, *t*, 12.1 min; *T*,
602 122.3 °C; *Et*, 100 %; and *S/L*, 27.2 g/L), which allow to obtain the maximum extraction
603 yield of both antioxidants simultaneously.

604

605 **4. Conclusions**

606 Optimal MAE conditions for *H* and *L* antioxidants from a surplus tomato crop were
607 determined in this study. A five-level full factorial Box-Behnken design was
608 successfully implemented and RSM used for analysis. The independent variables of *t*, *T*,
609 *Et* and *S/L* had significant effects on MAE. To predict the optimal extraction conditions,
610 a second-order polynomial model assuming interactive effects was fitted to each
611 response and the regression coefficients were determined using the least-squares
612 method. Optimal MAE conditions for *H*, *L* and both antioxidants were determined
613 based on the parametric response criteria P_m , V_m and IC_{50} . Overall, MAE proved to be a
614 powerful and efficient innovative methodology to extract the tomato antioxidants. In
615 statistical terms, the high values of the adjusted coefficient of determination ($R^2_{adj} >$
616 0.90) and the non-significant difference between predicted and experimental values
617 demonstrated the validity of the proposed optimization model. The results also indicated

618 that the antioxidant capacity of the *H* fraction was much higher than the *L* one.
619 Additionally, a discussion on the relationship between the extraction capacity of the
620 solvent in function of its polarity and the antioxidant activity of the extracts in *H* and *L*
621 media (the so-called polarity-activity relationship) was initiated, providing useful
622 information in the study of complex natural extracts containing ingredients with
623 opposite degrees of polarity.

624

625 **Acknowledgments**

626 The authors are grateful to the Foundation for Science and Technology (FCT, Portugal)
627 for financial support to CIMO (PEst-OE/AGR/UI0690/2014), REQUIMTE
628 (UID/QUI/50006/2013) and J. Pinela (SFRH/BD/92994/2013); to FCT/MEC and
629 FEDER under Programme PT2020 for financial support to LSRE
630 (UID/EQU/50020/2013), and to QREN, ON2 and FEDER (Project NORTE-07-0162-
631 FEDER-000050); to the Xunta de Galicia for financial support for the post-doctoral
632 researcher of M.A. Prieto.

633 **References**

- 634 Arnao, M.B., Cano, A., Acosta, M., 2001. The hydrophilic and lipophilic contribution to
635 total antioxidant activity. *Food Chem.* 73, 239–244.
- 636 Barros, L., Dueñas, M., Pinela, J., Carvalho, A.M., Santos Buelga, C., Ferreira, I.C.F.R.,
637 2012. Characterization and quantification of phenolic compounds in four tomato
638 (*Lycopersicon esculentum* L.) farmers' varieties in northeastern Portugal
639 homegardens. *Plant Foods Hum. Nutr.* 67, 229–234.
- 640 Bhuyan, D.J., Van Vuong, Q., Chalmers, A.C., van Altena, I. A., Bowyer, M.C.,
641 Scarlett, C.J., 2015. Microwave-assisted extraction of *Eucalyptus robusta* leaf for
642 the optimal yield of total phenolic compounds. *Ind. Crops Prod.* 69, 290–299.
- 643 Bors, W., Michel, C., Saran, M., 1984. Inhibition of the bleaching of the carotenoid
644 crocin a rapid test for quantifying antioxidant activity. *Biochim. Biophys. Acta*
645 796, 312–319.
- 646 Box, G., Hunter, J., 1957. Multi-factor experimental designs for exploring response
647 surfaces. *Ann. Math. Stat.* 28, 195–241.
- 648 Box, G., Hunter, J., Hunter, W., 2005. *Statistics for experimenters: design, innovation,*
649 and discovery.
- 650 Wiley. ISBN: 978-0-471-71813-0
- 651 Carocho, M., Ferreira, I.C.F.R., 2013a. A review on antioxidants, prooxidants and
652 related controversy: Natural and synthetic compounds, screening and analysis
653 methodologies and future perspectives. *Food Chem. Toxicol.* 51, 15–25.
- 654 Carocho, M., Ferreira, I.C.F.R., 2013b. The role of phenolic compounds in the fight
655 against cancer--a review. *Anticancer. Agents Med. Chem.* 13, 1236–58.
- 656 Comuzzi, C., Polese, P., Melchior, A., Portanova, R., Tolazzi, M., 2003.
657 SOLVERSTAT: a new utility for multipurpose analysis. An application to the

- 657 investigation of dioxygenated Co (II) complex formation in dimethylsulfoxide
658 solution. *Talanta* 59, 67–80.
- 659 Dahmoune, F., Nayak, B., Moussi, K., Remini, H., Madani, K., 2015. Optimization of
660 microwave-assisted extraction of polyphenols from *Myrtus communis* L. leaves.
661 *Food Chem.* 166, 585–595.
- 662 Dávalos, A., 2004. Extending applicability of the oxygen radical absorbance capacity
663 (ORAC-fluorescein) assay. *J. Agric. Food Chem.* 52, 48–54.
- 664 FAOSTAT, 2015. Food and Agriculture Organization of the United Nations. Statistics
665 division. Available at: <http://faostat3.fao.org/browse/Q/QC/E>. Accessed 5 March
666 2015.
- 667 Frankel, E.N., Meyer, A.S., 2000. The problems of using one-dimensional methods to
668 evaluate multifunctional food and biological antioxidants. *J. Sci. Food Agric.* 80,
669 1925–1941.
- 670 Frankel, E.N., Huang, S.W., Kanner, J., German, J.B., 1994. Interfacial phenomena in
671 the evaluation of antioxidants: bulk oils vs. emulsions. *J. Agric. Food Chem.* 42,
672 1054-1059.
- 673 Friedman, M., 2013. Anticarcinogenic, cardioprotective, and other health benefits of
674 tomato compounds lycopene, α -tomatine, and tomatidine in pure form and in fresh
675 and processed. *J. Agric. Food Chem.* 61, 9534–9550.
- 676 Gallo, M., Ferracane, R., Graziani, G., Ritieni, A., Fogliano, V., 2010. Microwave
677 assisted extraction of phenolic compounds from four different spices. *Molecules*
678 15, 6365–74.
- 679 García-Valverde, V., Navarro-González, I., García-Alonso, J., Periago, M.J., 2013.
680 Antioxidant bioactive compounds in selected industrial processing and fresh
681 consumption tomato cultivars. *Food Bioprocess Technol.* 6, 391–402.

- 682 Gieseg, S.P., Esterbauer, H., 1994. Low density lipoprotein is saturable by pro-oxidant
683 copper. FEBS Lett. 343, 188–194.
- 684 Huang, D., Ou, B., Hampsch-Woodill, M., Flanagan, J., Prior, R., 2002. High-
685 throughput assay of oxygen radical absorbance capacity (ORAC) using a
686 multichannel liquid handling system coupled with a microplate fluorescence reader
687 in 96-well format. J. Agric. Food Chem. 50, 4437–4444.
- 688 Huffman, B.A., Poltash, M.L., Hughey, C.A., 2012. Effect of polar protic and polar
689 aprotic solvents on negative-ion electrospray ionization and chromatographic
690 separation of small acidic molecules. Anal. Chem. 84, 9942–9950.
- 691 Jayasinghe, C., Gotoh, N., Wada, S., 2013. Pro-oxidant/antioxidant behaviours of
692 ascorbic acid, tocopherol, and plant extracts in n-3 highly unsaturated fatty acid
693 rich oil-in-water emulsions. Food Chem. 141, 3077–3084.
- 694 Jiménez-Escríg, A., Jiménez-Jiménez, I., Sánchez-Moreno, C., Saura-Calixto, F., 2000.
695 Evaluation of free radical scavenging of dietary carotenoids by the stable radical
696 2,2-diphenyl-1-picrylhydrazyl. J. Sci. Food Agric. 80, 1686–1690.
- 697 Kemmer, G., Keller, S., 2010. Nonlinear least-squares data fitting in Excel spreadsheets.
698 Nat. Protocols 5, 267–281.
- 699 Kislik, V.S., 2012. Solvent extraction: classical and novel approaches. Elsevier. ISBN:
700 978-0-444-53778-2
- 701 Kotíková, Z., Lachman, J., Hejtmánková, A., Hejtmánková, K., 2011. Determination of
702 antioxidant activity and antioxidant content in tomato varieties and evaluation of
703 mutual interactions between antioxidants. LWT - Food Sci. Technol. 44, 1703–
704 1710.
- 705 Li, Y., Fabiano-Tixier, A. S., Ruiz, K., Castera, A. R., Bauduin, P., Diat, O., Chemat, F.
706 (2015). Comprehension of direct extraction of hydrophilic antioxidants using

- 707 vegetable oils by polar paradox theory and small angle X-ray scattering analysis.
- 708 Food Chem. 173, 873–880.
- 709 Marco, G.J., 1968. A rapid method for evaluation of antioxidants. J. Am. Oil Chem.
- 710 Soc. 45, 594–598.
- 711 Michel, T., Destandau, E., Elfakir, C., 2011. Evaluation of a simple and promising
- 712 method for extraction of antioxidants from sea buckthorn (*Hippophaë rhamnoides*
- 713 L.) berries: Pressurised solvent-free microwave assisted extraction. Food Chem.
- 714 126, 1380–1386.
- 715 Murado, M.A., Prieto, M.A., 2013. NOEC and LOEC as merely concessive expedients:
- 716 Two unambiguous alternatives and some criteria to maximize the efficiency of
- 717 dose–response experimental designs. Sci. Total Environ. 461–462, 576–586.
- 718 Naguib, Y.M., 2000. A fluorometric method for measurement of oxygen radical-
- 719 scavenging activity of water-soluble antioxidants. Anal. Biochem. 284, 93–98.
- 720 Niki, E., Komuro, E., Takahashi, M., Urano, S., Ito, E., Terao, K., 1988. Oxidative
- 721 hemolysis of erythrocytes and its inhibition by free radical scavengers. J. Biol.
- 722 Chem. 263, 19809–14.
- 723 Oliveira, M., 2006. The evolution of the portuguese processed tomato sector: situation
- 724 and prospects on the global market. New Medit, Mediterr. J. Econ. Agric. Environ.
- 725 1, 38–46.
- 726 Pinela, J., Barros, L., Carvalho, A.M., Ferreira, I.C.F.R., 2012. Nutritional composition
- 727 and antioxidant activity of four tomato (*Lycopersicon esculentum* L.) farmer'
- 728 varieties in Northeastern Portugal homegardens. Food Chem. Toxicol. 50, 829–34.
- 729 Porter, W.L., 1993. Paradoxical behavior of antioxidants in food and biological systems.
- 730 Toxicol. Ind. Health, 9, 93–122.

- 731 Prieto, M.A., Vázquez, J.A., 2014. *In vitro* determination of the lipophilic and
732 hydrophilic antioxidant capacity of unroasted coffee bean extracts and their
733 synergistic and antagonistic effects. Food Res. Int. 62, 1183–1196.
- 734 Prieto, M.A., Murado, M.A., Vázquez, J.A., Curran, T.P., 2013. A new microplate
735 procedure for simultaneous assessment of lipophilic and hydrophilic antioxidants
736 and pro-oxidants, using crocin and β-carotene bleaching methods in a single
737 combined assay: Tea extracts as a case study. Food Res. Int. 53, 836–846.
- 738 Prieto, M.A., Rodríguez-Amado, I., Vázquez, J.A., Murado, M.A., 2012. β-Carotene
739 assay revisited. Application to characterize and quantify antioxidant and
740 prooxidant activities in a microplate. J. Agric. Food Chem. 60, 8983–8993.
- 741 Prieto, M.A., Vázquez, J.A., Murado, M.A., 2014. A critical point: The problems
742 associated with the variety of criteria to quantify the antioxidant capacity. J. Agric.
743 Food Chemistry 62, 5472–84.
- 744 Prieto, M.A., Vázquez, J.A., Murado, M.A., 2015. Crocin bleaching antioxidant assay
745 revisited: Application to microplate to analyse antioxidant and pro-oxidant
746 activities. Food Chem. 1, 299–310.
- 747 Prikler, S., 2009. Robert de Levie: Advanced Excel for scientific data analysis, 2nd ed.
748 Anal. Bioanal. Chem. 395, 1945.
- 749 Prior, R. L., Hoang, H., Gu, L., Wu, X., Bacchiocca, M., Howard, L., Hampsch-
750 Woodill, M., Huang, D., Ou, B., & Jacob, R. 2003. Assays for hydrophilic and
751 lipophilic antioxidant capacity (oxygen radical absorbance capacity (ORACFL)) of
752 plasma and other biological and food samples. J. Agric. Food Chem. 51, 3273–
753 3279.
- 754 Ranic, M., Nikolic, M., Pavlovic, M., Buntic, A., Siler-Marinkovic, S., Dimitrijevic-
755 Brankovic, S., 2014. Optimization of microwave-assisted extraction of natural

- 756 antioxidants from spent espresso coffee grounds by response surface methodology.
- 757 J. Clean. Prod. 80, 69–79.
- 758 Sashimatsung, Giribabu, M., Lanusunep, 2011. A study on Marketable surplus and
- 759 Price Spread of Tomato in Mokokchung District of Nagaland. Int. J. Humanit. Soc.
- 760 Sci. Invent. 2, 37–42.
- 761 Shahidi, F., Zhong, Y., 2015. Measurement of antioxidant activity. J. Funct. Foods 1–
- 762 25.
- 763 Shi, P., Tsai, C.-L., 2002. Regression model selection: a residual likelihood approach. J.
- 764 R. Stat. Soc. B (Statistical Methodol.) 64, 237–252.
- 765 Sultana, B., Anwar, F., Ashraf, M., 2009. Effect of extraction solvent/technique on the
- 766 antioxidant activity of selected medicinal plant extracts. Molecules 14, 2167–2180.
- 767 Taresco, V., Crisante, F., Francolini, I., Martinelli, A., D'Ilario, L., Ricci-Vitiani, L.,
- 768 Buccarelli, M., Pietrelli, L., Piozzi, A., 2015. Antimicrobial and antioxidant
- 769 amphiphilic random copolymers to address medical device-centered infections.
- 770 Acta Biomater. 22, 131–40.
- 771 Watanabe, J., Oki, T., Takebayashi, J., Takano-Ishikawa, Y., 2014. Extraction
- 772 Efficiency of Hydrophilic and Lipophilic Antioxidants from Lyophilized Foods
- 773 Using Pressurized Liquid Extraction and Manual Extraction. J. Food Sci. 79,
- 774 C1665–C1671.
- 775 Weibull, W., Sweden, S., 1951. A statistical distribution function of wide applicability.
- 776 J. Appl. Mech. 18, 293–297.
- 777 Zhang, P., Omaye, S., 2001. Antioxidant and prooxidant roles for β -carotene, α -
- 778 tocopherol and ascorbic acid in human lung cells. Toxicol. Vitr. 15, 13–24.

Figures

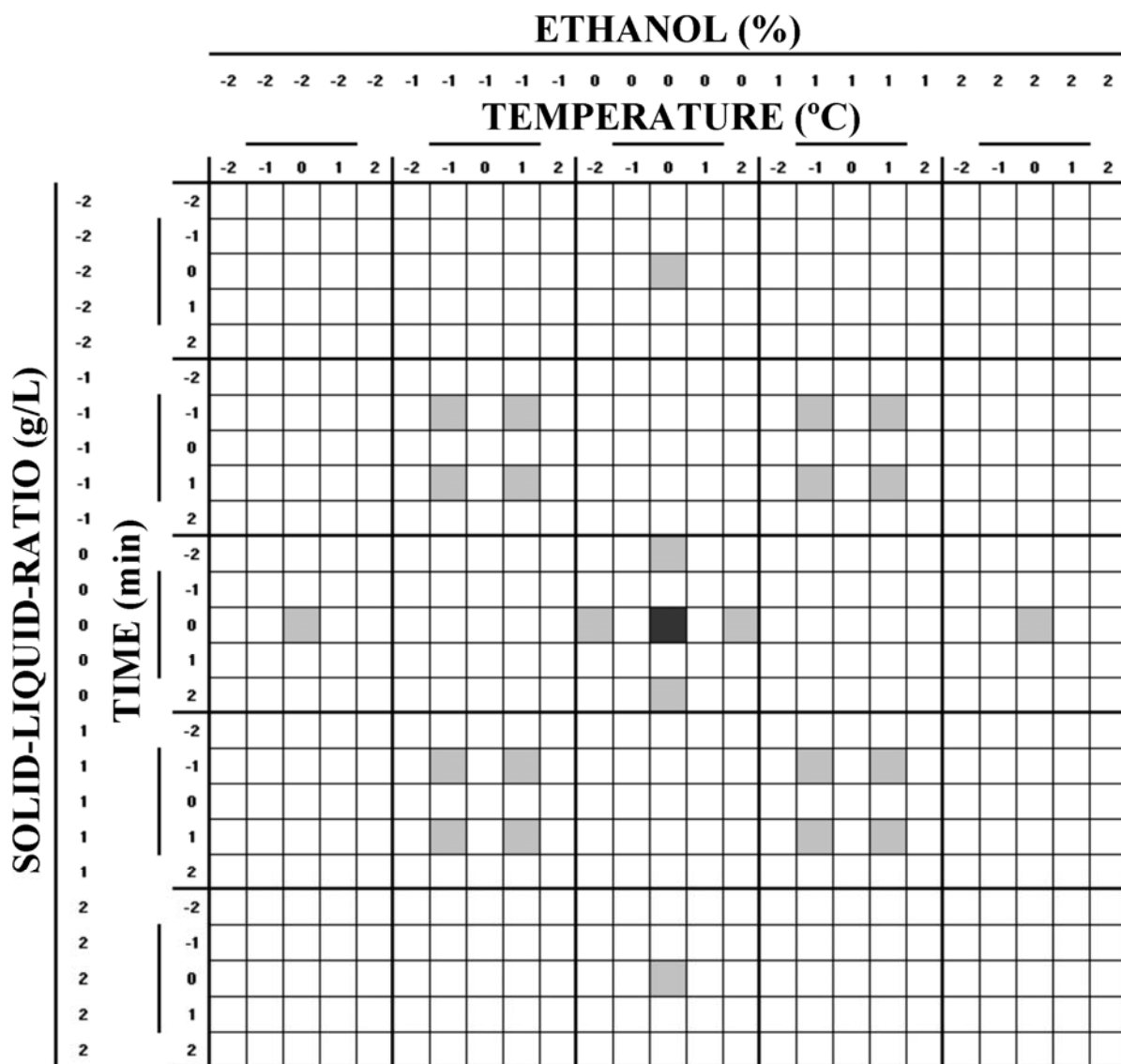


Fig. 2 - Visual representation of the applied experimental RSM design. Four independent variables (extraction time (X_1), temperature (X_2), ethanol concentration (X_3), and solid/liquid ratio (X_4)) were combined in a five-level full factorial design of 25 independent variable combinations (grey grid) and 7 replicates in the centre of the experimental domain (dark grid). Coded values (-2, -1, 0, +1, +2) are in natural values X_1 (t, min: 0, 5, 10, 15, 20), X_2 (T, °C: 60, 90, 120, 150, 180), X_3 (Et, %: 0, 25, 50, 75, 100) and X_4 (S/L, g/L: 5, 15, 25, 35, 45).

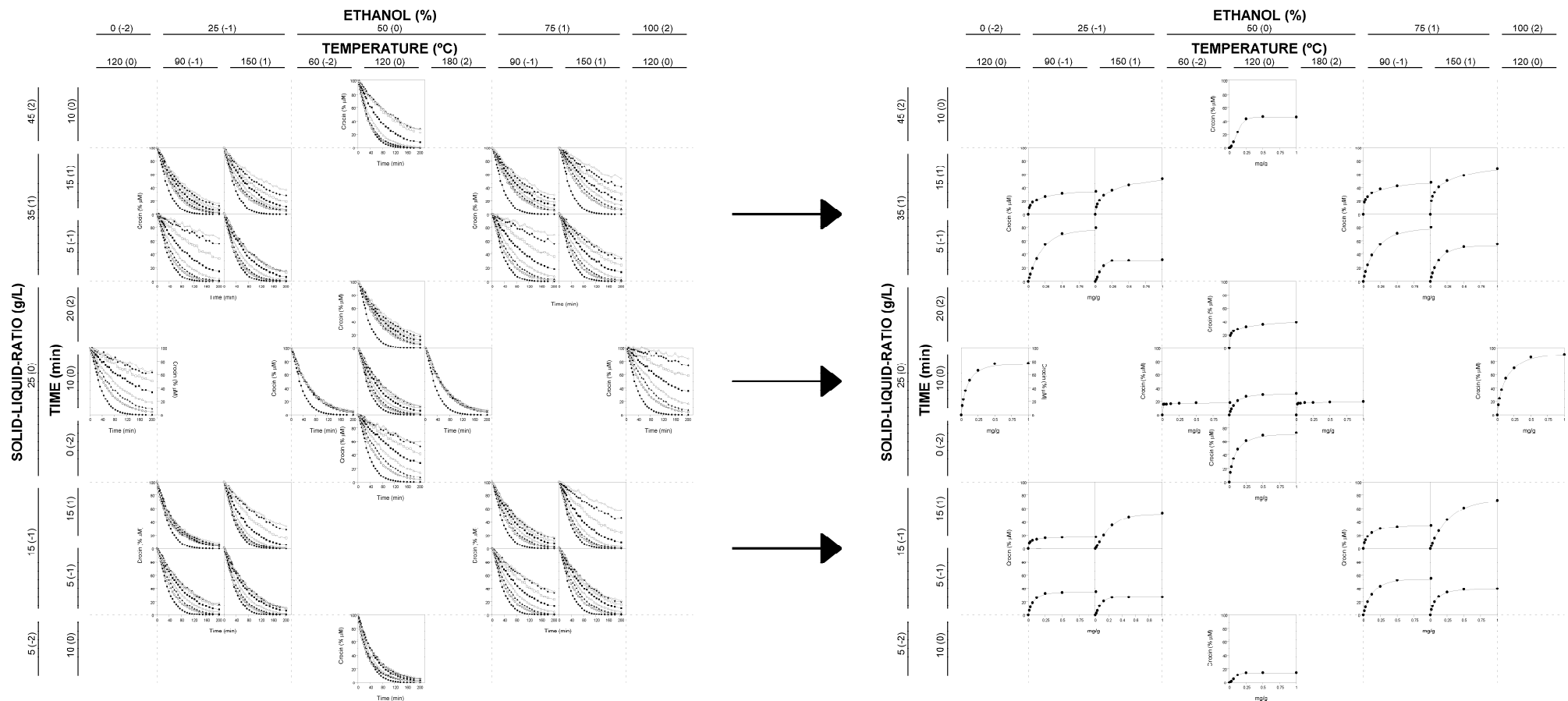


Fig. 3 - Illustration of the H responses obtained for the CM under the experimental RSM design presented in Fig. 2. On the left-hand side, each graph illustrates one of the 25 independent variable combinations and inside each graph can be seen the concentration-time responses of seven serial dilutions ($\circlearrowleft: 1/1$, $\blacktriangle: 1/2$, $\triangle: 1/4$, $\blacksquare: 1/8$, $\square: 1/16$, $\blacklozenge: 1/32$, $\lozenge: 1/64$) and the control (\bullet) of the extracted material. On the right-hand side, each graph shows: 1) Dots (\bullet), which represents the standardized \bar{P} (protected percentage of Cr) values in a concentration-response format obtained by applying Eq. (2) to the concentration-time responses presented in the left-hand side; and 2) Lines (—), fitted responses to the mathematical model of Eq. (3). The obtained parametric fitting values are presented in Table 1.

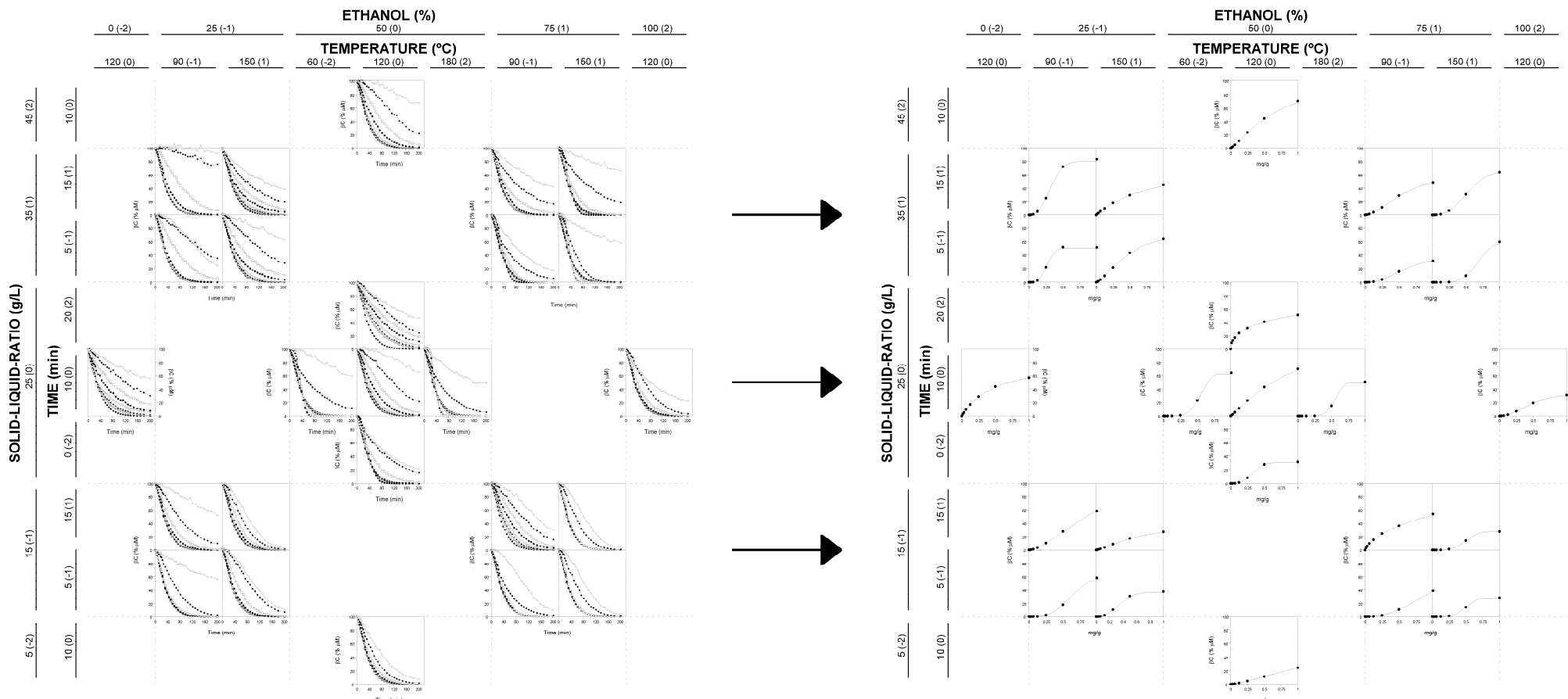


Fig. 4 - Illustration of the L responses obtained for the β CM under the experimental RSM design presented in Fig. 2. On the left-hand side, each graph illustrates one of the 25 independent variable combinations and inside each graph can be seen the concentration-time responses of seven serial dilutions (\circ : 1/1, \blacktriangle : 1/2, \triangle : 1/4, \blacksquare : 1/8, \square : 1/16, \blacklozenge : 1/32, \lozenge : 1/64) and the control (\bullet) of the extracted material. On the right-hand side, each graph shows: 1) Dots (\bullet), which represents the standardized \bar{P} (protected percentage of β C) values in a concentration-response format obtained by applying Eq. (2) to the concentration-time responses presented in the left-hand side; and 2) Lines (—), fitted responses to the mathematical model of Eq. (3). The obtained parametric fitting values are presented in Table 1.

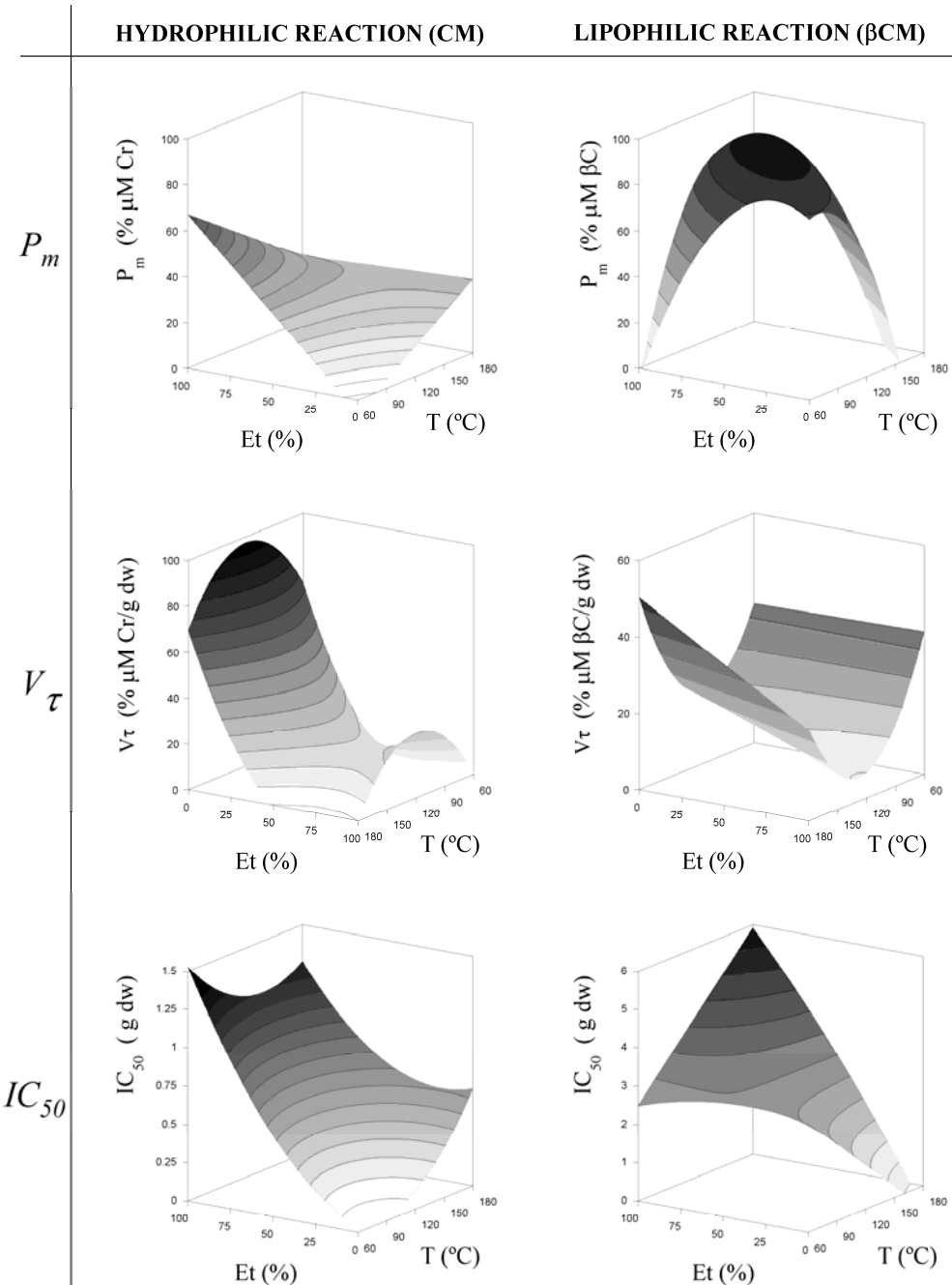


Fig. 5 - Response surfaces of the effects of ethanol concentration (Et) vs temperature (T) on H and L antioxidant reactions for the parametric response criteria P_m , V_m and IC_{50} . For representation purposes, the variables t and S/L were positioned at the centre of their experimental domain ($t=10$ min and $S/L=25$ g/L). The obtained parametric fitting values are presented in Table 2.

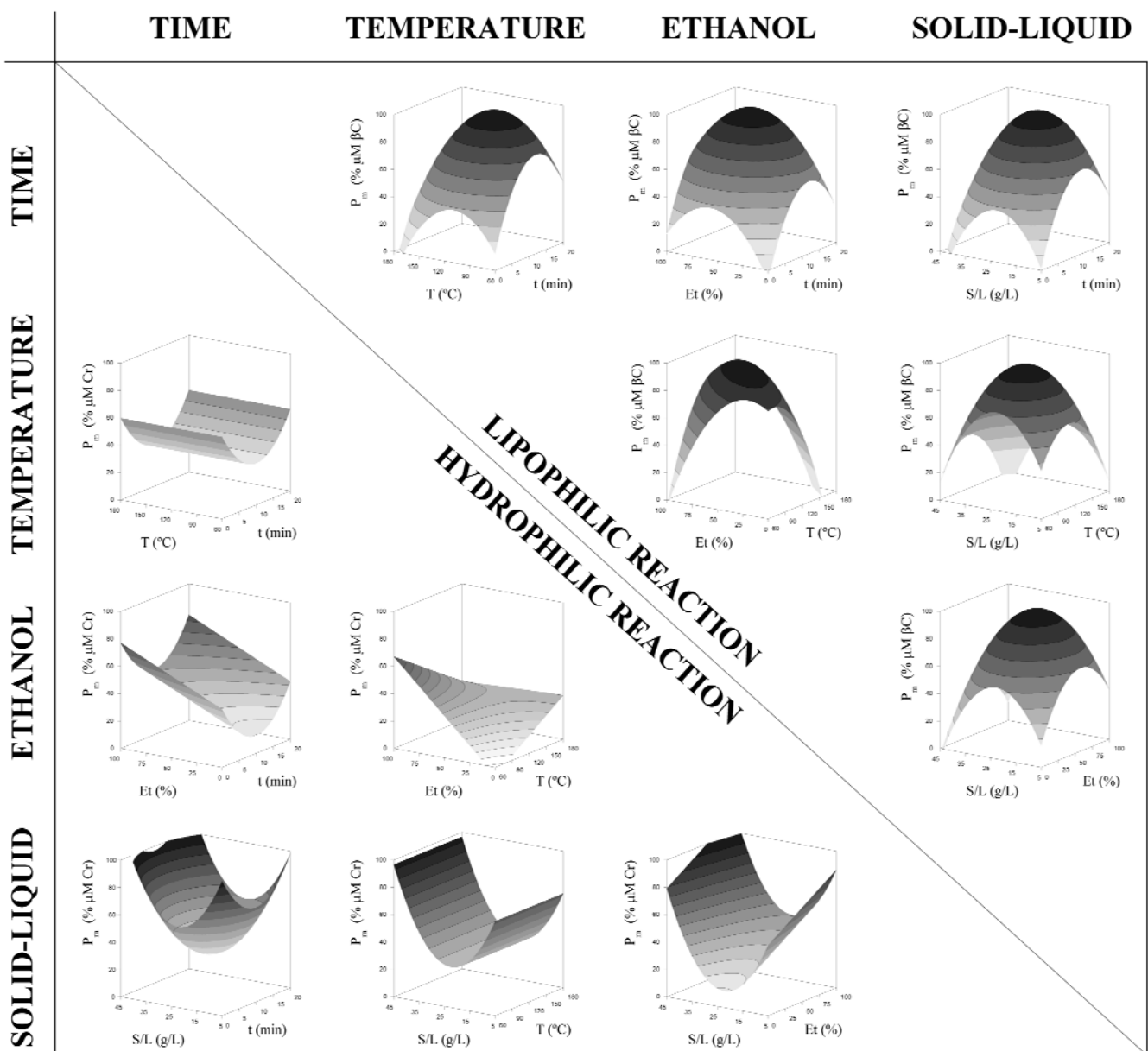


Fig. 6 - Matrix combination of the response surfaces of the *H* and *L* antioxidant reactions obtained for the parametric coefficient P_m (% μM of Cr or βC), which is organized as follows: a) in the top diagonal part is presented the response surface of the *L* reaction; and b) in the bottom diagonal part is presented the response surface of the *H* reaction. For representation purposes, the variables excluded in each 3D graph were positioned at the centre of their experimental domain ($t=10$ min; $T=120$ °C; $Et=50$ %; and $S/L=25$ g/L). The obtained parametric fitting values are presented in Table 2.

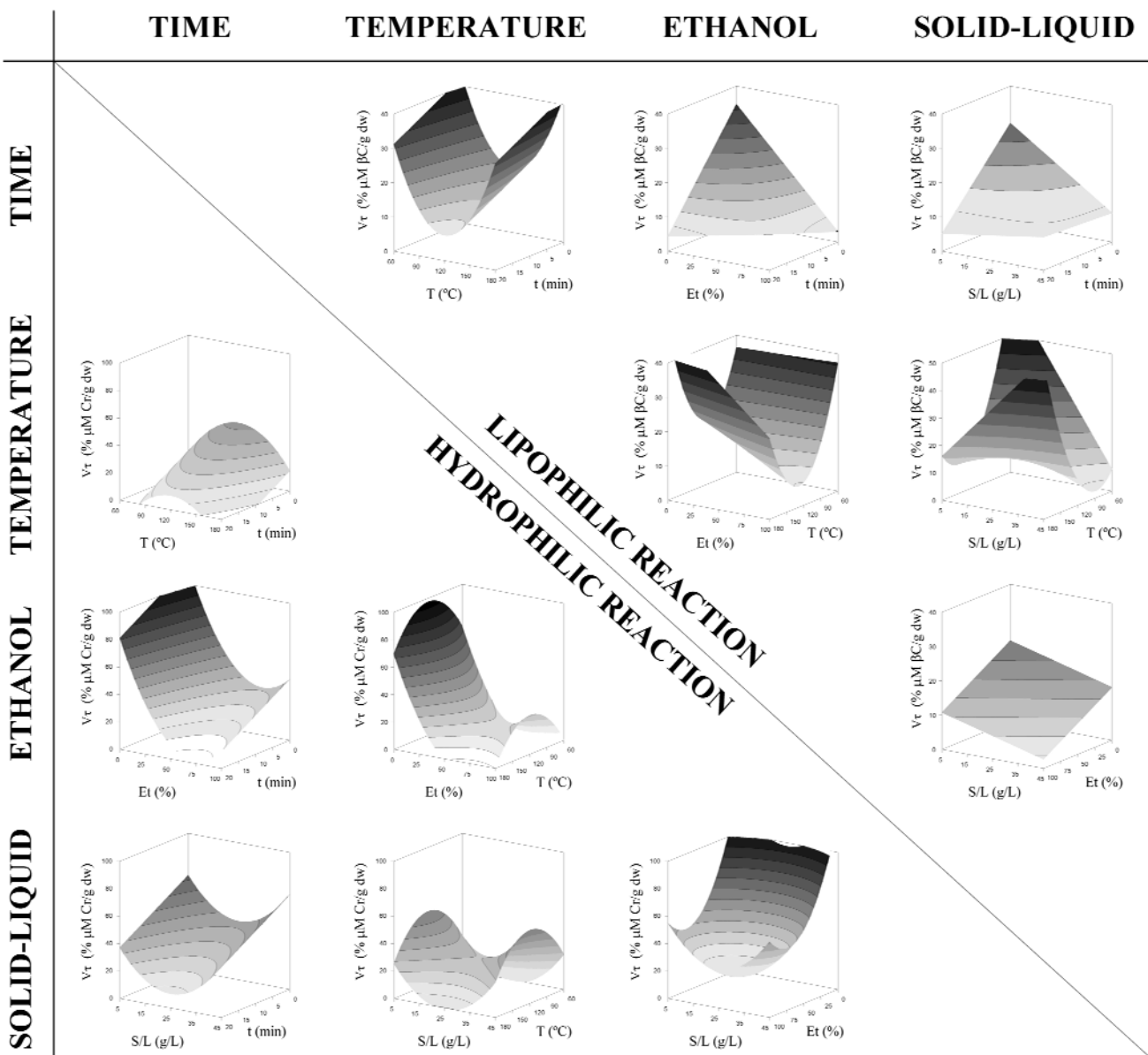


Fig. 7 - Matrix combination of the response surfaces of the *H* and *L* antioxidant reactions obtained for the parametric coefficient V_m (% μM of Cr or $\beta\text{C}/\text{g}$ extract), which is organized as follows: a) in the top diagonal part is presented the response surface of the *L* reaction; and b) in the bottom diagonal part is presented the response surface of the *H* reaction. For representation purposes, the variables excluded in each 3D graph were positioned at the centre of their experimental domain ($t=10$ min; $T=120$ °C; $Et=50$ %; and $S/L=25$ g/L). The obtained parametric fitting values are presented in Table 2.

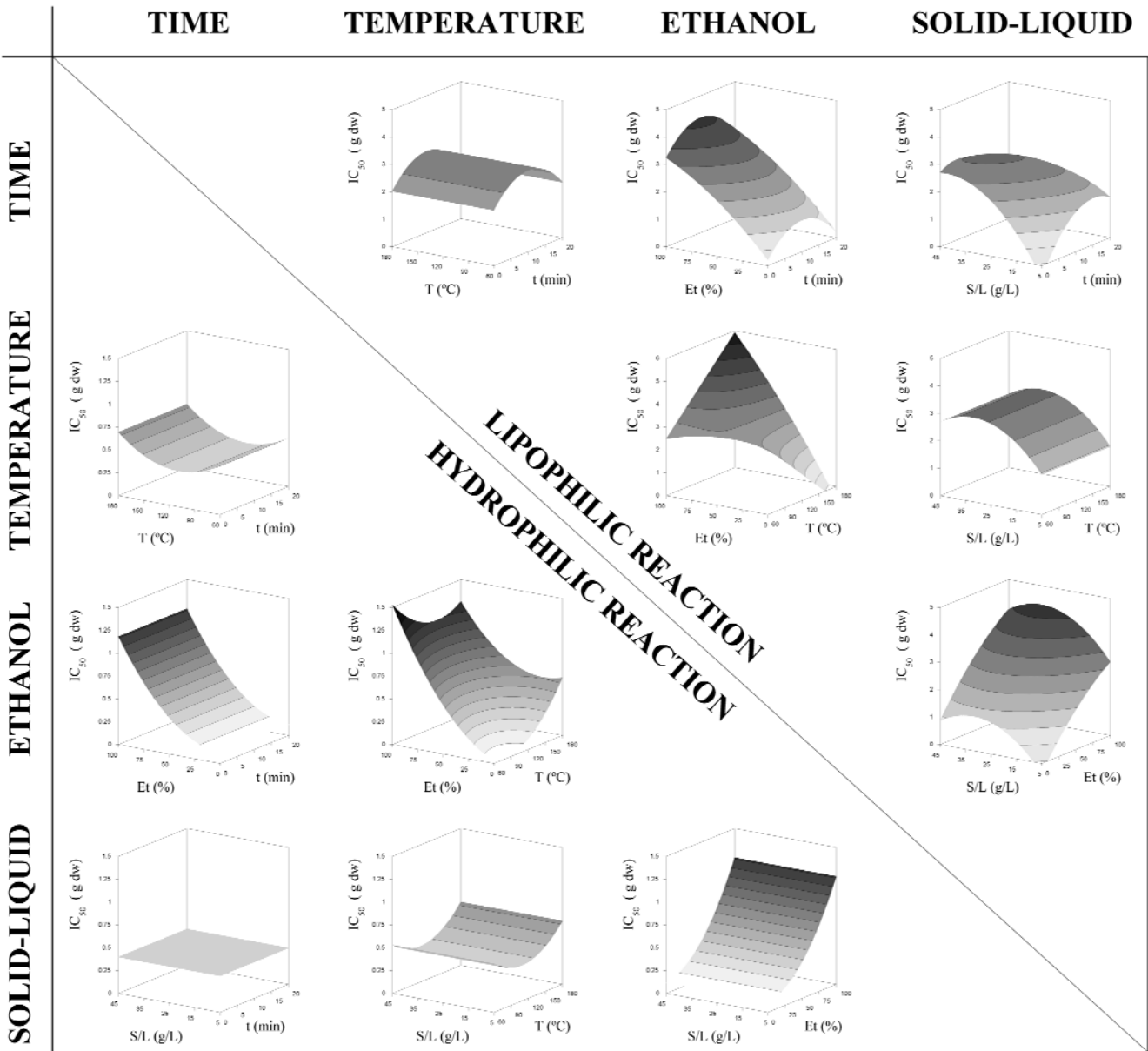


Fig. 8 - Matrix combination of the response surfaces of the *H* and *L* antioxidant reactions obtained for the parametric coefficient IC_{50} (g extract), which is organized as follows: a) in the top diagonal part is presented the response surface of the *L* reaction; and b) in the bottom diagonal part is presented the response surface of the *H* reaction. For representation purposes, the variables excluded in each 3D graph were positioned at the centre of their experimental domain ($t=10$ min; $T=120$ °C; $Et=50$ %; and $S/L=25$ g/L). The obtained parametric fitting values are presented in Table 2.

Tables

Table 1 - Estimated numerical values of the parameters (P_m , V_m and IC_{50}) of Eqs. (3) and (4), after fitting the concentration-response values presented in the right-hand side of Fig. 3 and Fig. 4 (for CM and β CM, respectively) of the tomato extracts obtained under the experimental RSM design presented in Fig. 2.

RUN	EXPERIMENTAL DOMAIN				PARAMETRIC ANTIOXIDANT RESPONSES								STATISTICS	
					CM (HYDROPHILIC REACTION)				β CM (LIPOPHILIC REACTION)				R^2_{adj}	
	$X_1: t$ min	$X_2: T$ °C	$X_3: Et$ %	$X_4: S/L$ g/L	P_m % $\mu M Cr$	IC_{50} g extract	a --	V_m % $\mu M Cr/g$ extract	P_m % $\mu M \beta C$	IC_{50} g extract	a --	V_m % $\mu M \beta C/g$ extract	CM	βCM
1	-1(5)	-1(90)	-1(25)	-1(15)	34.08±0.47	0.17±0.01	0.84±0.14	58.24±4.75	59.64±1.52	1.88±0.17	3.06±0.32	33.69±1.39	0.9884	0.9569
2	1(15)	-1(90)	-1(25)	-1(15)	17.50±1.42	0.06±0.00	0.42±0.03	44.24±2.52	94.39±2.35	2.43±0.22	1.56±0.13	20.99±3.93	0.9567	0.9906
3	-1(5)	1(150)	-1(25)	-1(15)	26.63±1.81	0.19±0.02	1.16±0.03	56.07±0.36	36.28±1.70	1.02±0.03	2.35±0.00	28.87±4.35	0.9587	0.9727
4	1(15)	1(150)	-1(25)	-1(15)	51.12±3.09	0.51±0.04	1.18±0.16	41.33±4.78	31.79±1.05	1.41±0.04	1.25±0.04	9.79±1.25	0.9760	0.9896
5	-1(5)	-1(90)	1(75)	-1(15)	53.46±0.31	0.29±0.03	0.90±0.01	57.66±6.18	58.77±1.60	2.56±0.05	2.38±0.41	18.96±1.15	0.9744	0.9641
6	1(15)	-1(90)	1(75)	-1(15)	35.08±3.07	0.18±0.02	0.64±0.11	43.38±0.02	86.41±3.34	2.09±0.02	0.75±0.15	10.73±1.64	0.9706	0.9766
7	-1(5)	1(150)	1(75)	-1(15)	38.52±1.12	0.19±0.00	0.80±0.09	57.32±9.27	28.00±1.27	1.50±0.07	4.02±0.09	26.00±1.72	0.9549	0.9827
8	1(15)	1(150)	1(75)	-1(15)	72.90±5.31	0.59±0.01	0.98±0.09	41.77±4.87	27.31±1.19	1.50±0.10	3.81±0.23	24.01±4.50	0.9752	0.9847
9	-1(5)	-1(90)	-1(25)	1(35)	76.55±3.65	1.06±0.04	0.99±0.09	24.92±4.07	49.85±2.10	1.90±0.14	3.22±0.25	29.27±4.71	0.9726	0.9654
10	1(15)	-1(90)	-1(25)	1(35)	37.83±2.73	0.59±0.02	0.47±0.01	10.34±1.78	79.68±2.70	2.24±0.16	2.52±0.03	31.09±4.89	0.9921	0.9787
11	-1(5)	1(150)	-1(25)	1(35)	30.50±2.12	0.53±0.03	1.23±0.18	24.75±1.05	69.58±3.48	2.84±0.09	1.36±0.19	11.52±1.89	0.9828	0.9787
12	1(15)	1(150)	-1(25)	1(35)	61.51±5.82	1.18±0.12	0.53±0.09	9.55±1.57	57.48±3.64	3.44±0.06	0.98±0.07	5.70±0.25	0.9787	0.9531
13	-1(5)	-1(90)	1(75)	1(35)	78.81±2.66	0.97±0.07	0.88±0.13	24.92±4.79	33.10±2.59	3.81±0.27	2.15±0.08	6.48±0.24	0.9886	0.9919
14	1(15)	-1(90)	1(75)	1(35)	58.61±2.29	0.68±0.05	0.36±0.06	10.65±0.27	53.69±1.90	3.42±0.09	1.63±0.20	8.87±0.38	0.9677	0.9802
15	-1(5)	1(150)	1(75)	1(35)	52.40±4.11	0.67±0.04	0.91±0.02	24.62±1.73	57.62±5.40	4.76±0.10	4.54±0.18	19.03±3.35	0.9621	0.9740
16	1(15)	1(150)	1(75)	1(35)	80.03±7.95	0.82±0.01	0.44±0.01	14.79±2.66	64.42±4.82	3.65±0.14	2.65±0.26	16.21±1.22	0.9884	0.9920
17	-2(0)	0(120)	0(50)	0(25)	70.12±4.78	0.35±0.04	0.77±0.15	53.75±2.98	30.00±2.69	1.65±0.03	2.85±0.31	17.97±0.72	0.9742	0.9818
18	2(20)	0(120)	0(50)	0(25)	51.04±1.47	0.37±0.03	0.27±0.03	12.73±1.61	75.06±5.76	2.01±0.15	0.52±0.08	6.74±1.17	0.9840	0.9787
19	0(10)	-2(60)	0(50)	0(25)	32.22±2.89	0.48±0.03	0.05±0.01	1.20±0.10	63.15±4.18	2.77±0.05	4.68±0.57	36.97±5.78	0.9570	0.9858
20	0(10)	2(180)	0(50)	0(25)	34.89±0.90	0.63±0.04	0.06±0.01	1.15±0.15	49.13±3.50	2.78±0.11	6.15±0.14	37.60±1.55	0.9832	0.9728
21	0(10)	0(120)	-2(0)	0(25)	75.52±6.22	0.35±0.03	0.82±0.10	60.82±6.17	64.81±5.05	1.57±0.05	0.94±0.09	13.44±0.53	0.9586	0.9685
22	0(10)	0(120)	2(100)	0(25)	91.54±3.46	0.47±0.02	0.75±0.06	50.61±1.06	31.65±0.18	2.21±0.04	1.90±0.18	9.46±1.02	0.9710	0.9626
23	0(10)	0(120)	0(50)	-2(5)	14.31±0.87	0.08±0.01	1.67±0.29	104.79±7.95	41.18±3.07	0.84±0.08	1.44±0.03	24.35±0.89	0.9569	0.9628
24	0(10)	0(120)	0(50)	2(45)	45.20±0.57	1.11±0.03	1.82±0.33	25.84±0.25	80.94±0.64	4.13±0.51	1.23±0.04	8.37±0.10	0.9551	0.9755
25	0(10)	0(120)	0(50)	0(25)	30.39±1.56	0.38±0.04	0.91±0.02	25.43±0.89	97.33±6.29	3.03±0.31	1.11±0.18	12.35±0.56	0.9728	0.9592
26	0(10)	0(120)	0(50)	0(25)	30.39±2.04	0.38±0.02	0.91±0.00	25.43±4.30	97.33±8.12	3.13±0.11	1.26±0.11	13.63±1.08	0.9585	0.9870
27	0(10)	0(120)	0(50)	0(25)	30.39±0.50	0.38±0.04	0.91±0.01	25.43±2.09	97.33±2.22	3.05±0.16	1.14±0.14	12.58±1.01	0.9700	0.9843
28	0(10)	0(120)	0(50)	0(25)	30.39±1.15	0.38±0.03	0.91±0.03	25.43±0.14	97.33±4.47	3.05±0.16	1.12±0.01	12.34±0.38	0.9637	0.9834
29	0(10)	0(120)	0(50)	0(25)	30.39±1.50	0.38±0.01	0.91±0.08	25.43±0.27	97.33±0.94	3.04±0.28	1.10±0.15	12.21±1.68	0.9725	0.9575
30	0(10)	0(120)	0(50)	0(25)	30.39±2.12	0.38±0.04	0.91±0.11	25.43±5.02	97.33±3.58	2.80±0.28	1.08±0.04	13.03±0.14	0.9888	0.9782
31	0(10)	0(120)	0(50)	0(25)	30.39±1.83	0.38±0.02	0.91±0.00	25.43±2.82	87.46±3.60	2.66±0.30	1.43±0.17	16.30±2.64	0.9886	0.9823
32	0(10)	0(120)	0(50)	0(25)	30.39±0.57	0.38±0.01	0.91±0.13	25.43±1.54	97.33±8.41	3.00±0.18	1.16±0.19	13.02±2.59	0.9827	0.9906

Table 2 - Estimated coefficient values of Eq. (8), parametric intervals and numerical statistical criteria for each parametric response criteria of the H and L reactions.

		HYDROPHILIC REACTION			LIPOPHILIC REACTION		
		P_m	IC_{50}	V_m	P_m	IC_{50}	V_m
<i>Fitting coefficients obtained from Eq. (8) and showed in Eqs. (9)-(14)</i>							
Intercept	b_0	30.50±2.74	0.40±0.05	26.49±3.10	96.10±0.05	2.97±0.37	13.09±0.01
Linear effect	b_1	ns	ns	-8.10±1.79	8.02±0.05	ns	-2.87±0.01
	b_2	ns	0.04±0.04	ns	-7.13±0.03	ns	ns
	b_3	8.70±2.09	0.27±0.04	-17.22±1.79	5.10±0.03	0.76±0.17	-3.20±0.01
	b_4	6.92±2.09	ns	ns	-5.65±0.04	0.31±0.11	-2.02±0.01
Quadratic effect	b_{11}	7.29±1.87	ns	ns	-10.72±0.07	-0.24±0.10	ns
	b_{22}	ns	0.05±0.03	-6.96±1.79	-9.82±0.03	ns	5.94±0.01
	b_{33}	ns	0.06±0.03	9.08±1.79	-8.59±0.06	-0.07±0.03	ns
	b_{44}	13.02±1.87	ns	6.68±1.61	-11.80±0.08	-0.22±0.10	ns
Interactive effect	b_{12}	13.21±2.56	0.16±0.05	ns	-7.70±0.03	ns	ns
	b_{13}	ns	ns	ns	ns	ns	2.35±0.02
	b_{14}	ns	ns	ns	ns	-0.24±0.02	1.57±0.02
	b_{23}	-4.78±2.56	-0.05±0.05	ns	13.04±0.02	0.43±0.12	-1.72±0.02
	b_{24}	ns	ns	ns	ns	ns	6.21±0.02
	b_{34}	ns	ns	ns	ns	0.27±0.07	ns
<i>Statistical information of the fitting analysis</i>							
Observations		32	32	32	32	32	32
R^2		0.9526	0.9236	0.9743	0.9422	0.9437	0.9223
$R^2\text{adj}$		0.9331	0.9136	0.9612	0.9067	0.9058	0.9019
MSE		767.23	0.15	854.45	1127.20	1.50	152.70
RMSE		27.70	0.38	29.23	33.57	1.22	12.35
MAPE		5.99	12.63	22.22	8.14	7.97	9.72
DW		2.39	2.20	1.43	2.32	1.36	2.32

ns: no significant coefficient; **R^2 :** Correlation coefficient; **$R^2\text{adj}$:** The adjusted coefficient of determination for the model; **MSE:** The mean squared error; **RMSE:** The root mean square of the errors; **MAPE:** The mean absolute percentage error; and **DW:** The Durbin-Watson statistic.

Table 3 - Operating conditions that maximize the extraction of *H* and *L* antioxidants from tomato and optimal response values for the parametric response criteria (P_m , V_m and IC_{50}) and antioxidant reactions (*H* or *L*). The independent variables t , T , Et and S/L are presented in natural values.

OPTIMAL EXTRACTION CONDITIONS				RESPONSE OPTIMUM
$X_1: t$ (min)	$X_2: T$ (°C)	$X_3: Et$ (%)	$X_4: S/L$ (g/L)	
For <i>H</i> antioxidants				
P_m (<i>H</i>)	18.7	180.0	56.8	45.0 100 % $\mu M Cr$
V_m (<i>H</i>)	2.5	120.0	0.0	5.0 136.11 % $\mu M Cr/g extract$
IC_{50} (<i>H</i>)	14.5	90.0	44.0	17.0 0.051 g extract
For <i>L</i> antioxidants				
P_m (<i>L</i>)	13.4	93.6	44.0	21.3 100 % $\mu M \beta C$
V_m (<i>L</i>)	2.2	180.0	100.0	5.0 78.70 % $\mu M \beta C/g extract$
IC_{50} (<i>L</i>)	2.6	169.1	91.7	10.9 0.025 g extract
For each response criteria of both <i>H</i> and <i>L</i> antioxidants				
P_m (<i>H</i>)	15.4	127.6	93.2	33.8 100.0 % $\mu M Cr$
P_m (<i>L</i>)				43.4 % $\mu M \beta C$
V_m (<i>H</i>)	3.9	63.3	0.0	5.0 108.93 % $\mu M Cr/g extract$
V_m (<i>L</i>)				74.79 % $\mu M \beta C/g extract$
IC_{50} (<i>H</i>)	13.9	112.7	89.0	5.3 0.06 g extract
IC_{50} (<i>L</i>)				0.05 g extract
For <i>H</i> and <i>L</i> antioxidants based on all the response criteria				
P_m (<i>H</i>)				100 % $\mu M Cr$
V_m (<i>H</i>)	2.25	149.2	99.1	45.0 60.2 % $\mu M Cr/g extract$
IC_{50} (<i>H</i>)				0.09 g extract
P_m (<i>L</i>)				92.4 % $\mu M \beta C$
V_m (<i>L</i>)	15.4	60.0	33.0	15.0 42.9 % $\mu M \beta C/g extract$
IC_{50} (<i>L</i>)				0.38 g extract
For both <i>H</i> and <i>L</i> antioxidants based on all the response criteria				
P_m (<i>H</i>)				100.0 % $\mu M Cr$
P_m (<i>L</i>)				39.1 % $\mu M \beta C$
V_m (<i>H</i>)	12.1	122.3	100.0	27.2 46.39 % $\mu M Cr/g extract$
V_m (<i>L</i>)				9.64 % $\mu M \beta C/g extract$
IC_{50} (<i>H</i>)				0.47 g extract
IC_{50} (<i>L</i>)				0.47 g extract