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Title: Kinetics of the combustion of olive oil. A semi-global

model

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- 1 Kinetics of the combustion of olive oil. A semi-global model
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- 5 1. INTRODUCTION
- 6 The increasing need of energy by segments of our society, the reduction of petroleum
- 7 reserves and increased environmental concerns have caused biomass materials to gain
- 8 much interest with respect to energy utilization. For example, waste vegetable oils can
- 9 through thermal decomposition be used to directly obtain energy or fuels [1]. It is very
- 10 important to perform thermal analysis of the oils to predict their behavior in real
- 11 combustion systems. The combustion kinetics of these fuels gives relevant information
- on their thermal behavior and on the possible formation of a carbonaceous residue and
- its subsequent oxidation.
- Jansson et al. [2] studied the pyrolysis of olive oils and other vegetable oils, and
- determined the evolved compounds on a Pyrolyzer/GC/MS. Gases such as propene and
- liquids such as oleic acid, docosene and octadecenal, with boiling points at around 360
- 17 °C (633 K), were found. In a combustion process these compounds are oxidized, which
- changes the composition of the gas phase.
- 19 The subject of a previous paper was a study of the pyrolysis kinetics of olive and used
- 20 olive oil [3]. The pyrolytic decomposition was analyzed taking into consideration the
- 21 vaporization process involved, and the results were compared with a number of kinetic
- 22 considerations discussed in other papers [4-6]. The proposed kinetic model considered
- 23 two sequential processes: a first process, considering vaporization and decomposition,
- 24 whose apparent activation energy and reaction order were 112 kJ/mol and 0.606,
- 25 respectively, and a second process, whose apparent activation energy and reaction order
- were 194.6 kJ/mol and 2.274, respectively. The values obtained in both of these
- 27 processes are acceptable; in the first process, the values are between those of the

28 29	values for decomposition processes.
30	Others have also studied the oxidative thermal decomposition in order to characterize
31	vegetable oils [7,8]. Tran et al. [9] examined a number of mechanisms of the
32	combustion of oxygenated compounds of biofuels.
33	Dweck and Sampaio [10] analyzed the thermal decomposition of commercial vegetable
34	oils by TG/DTA and observed four decomposition steps. They proposed that the last
35	one corresponds to the burnout of the residual carbonaceous material.
36	Concerning the global kinetics, Vecchio et al. [7] studied the oxidative thermal
37	decomposition of single-varietal extra olive oil by TG/DSC, and observed a complex
38	multistep decomposition. They attributed the first apparent peak to two different
39	processes for the purpose of relating them to the chemical composition. From the first
40	decomposition step they obtained apparent activation energies for the de-convoluted
41	peaks ranging between 27 and 158 kJ/mol, and 31 and 278 kJ/mol for the first and
42	second peaks, respectively. No other information concerning kinetic parameters was
43	presented.
44	Gouveia de Souza et al. [11] elucidated the oxidation kinetics of sunflower oil by TG,
45	by considering three decomposition steps in which the interaction of the oxidation
46	reactions was important. The first step takes place between 503 and 653 K with reaction
47	order around 1 and activation energy around 90-110 kJ/mol, in which the volatile
48	compounds were removed by the vapor generated during heating. The second is
49	between 653 and 753 K with reaction order around 2 and activation energy of 205-300
50	kJ/mol. The third step takes place between 753 and 823 K and the deduced reaction
51	orders and apparent activation energies were around 2 and 300-400 kJ/mol,
52	respectively.
53	Santos et al. [12] considered three decomposition steps in the oxidative decomposition
54	of a number of edible oils, including olive oil. Similar kinetic parameters were obtained.
55	In the first step, the apparent activation energy was between 78 and 106 kJ/mol and the
56	reaction order was between 0.92 and 1.06. In the second step, the apparent activation
57	energy was between 208 and 349 kJ/mol and the reaction order was between 1.86 and
58	2.11. In the last step, an activation energy between 274 and 370 kJ/mol and a reaction

59	order between 1.87 and 2.13 were obtained. No values were reported for the mass
60	fractions of the volatiles evolved in each step.
<i>(</i> 1	
61	Zhengwen [13] recently studied the combustion of cooking oil tar on a TG apparatus.
62	He observed four DTG peaks after the initial evaporation of the absorbed water, and
63	made several plots for correlating the data which suggest a model of First Order
64	Reaction and Three-dimensional Diffusion Separate-stage. However, values for the
65	apparent activation energy were not reported.
66	Vecchio et al. [14] studied the decomposition of triglycerides contained in olive oil by
67	TG. They observed the presence of four decomposition steps and determined the kinetic
68	parameters of the first two decomposition steps.
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69	More recently, Tomassetti et al. [15] analyzed the thermal decomposition of saturated
70	mono-, di- and tri-glycerides. They also observed four decomposition steps and
71	proposed the kinetic parameters for the two or three first steps.
72	
72 72	The decomposition kinetics of complex materials (synthetic polymers, biomass, oils,
73	etc.) is a subject that deals with the examination and analysis of kinetic parameters, with
74	a view to clarifying their significance [16-18]. Thus, efforts to study the decompositions
75	of substances such as vegetable oils can help to reduce the existing chaos in the field of
76	reaction kinetics of complex materials.
77	In this paper, a kinetic model for the combustion of olive oil at air atmosphere and also
78	in one that is oxygen-poor has been developed by simultaneous determination at each
79	step of the kinetic parameters and the mass fraction of the volatiles. The experimental
80	data are compared with those obtained by simulation using the deduced expressions. We
81	also discuss the possibility of a carbonaceous residue formed during the thermal
82	oxidation of the fuels in question, which has not been considered in previous papers.
83	The kinetic study is analyzed by contrasting with the study on olive oil by Vecchio et al.
84	[7], as well as other studies carried out on other vegetable oils. The kinetic model can be
85	used to characterize certain decomposition steps of the edible oils and/or their
86	corresponding wastes and to analyze the formation of a carbonaceous residue

87

#### 88 2. EXPERIMENTAL

89	2.1 Raw material
90	Pure olive oil, waste olive oil and waste mixed oil were selected as materials for
91	studying the kinetics. This study employed the same pure olive oil as a previous
92	pyrolysis kinetic study [3]. The waste olive oil was obtained after four/five frying
93	processes, which corresponds to an average use of this oil, and was also the same waste
94	olive oil employed in the previous pyrolysis kinetic study [3]. A waste mixed oil,
95	consisting of a mixture of different used cooking oils, was also utilized to determine
96	whether there are any great differences between the oils. An elemental analysis of the
97	samples was carried out on a Perkin-Elmer 2400 to determine the mass fractions of
98	carbon, hydrogen, nitrogen and sulphur; oxygen content was determined by a direct
99	oxygen analysis carried out on a Flash-2000 Thermo Fisher Scientific; a LECO
100	Instruments AC-350 calorimetric bomb was used to obtain the net calorific value. Table
101	1 shows the results of the elemental analysis and the net calorific values of the three
102	samples tested. As observed, there are no big differences between the samples.
103	Table 1
104	2.2 Apparatus and experimental procedure
105	The combustion runs at air atmosphere were carried out on two different TG apparatus
106	whereas in the $N_2:O_2 = 9:1$ runs only one of them was used:
107	1) A Mettler Toledo Thermobalance model TGA/SDTA851e/LF/1600. This instrument
108	incorporates a horizontal furnace and a parallel-guided balance. In this way, positioning
109	of the sample has no influence on the measurement, and flow gas perturbation and
110	thermal buoyancy are minimized. The sample temperature was measured by a sensor
111	directly attached to the sample holder. Two different atmospheres were used; $N_2:O_2 =$
112	4:1 and $N_2:O_2 = 9:1$ . The crucibles employed in the runs were a nearly cylindrical
113	aluminum crucible of 0.55 cm internal diameter and 0.41 cm height, which is slightly
114	curved at the bottom of the cylinder, and a cylindrical alumina crucible of 0.47 cm
115	internal diameter and 0.42 cm height.
116	2) A Perkin Elmer Thermobalance model TGA/SDTA-6000. This instrument
117	incorporates a vertical furnace and a single beam vertical balance. As in the previous
118	case, positioning of the sample has no influence on the measurement, and flow gas
119	perturbation and thermal buoyancy are minimized. The SaTurnA sensor measures both

120	the sample and reference temperature directly for superb performance. The alumina
121	crucible used in all runs was nearly cylindrical with 0.65 cm internal diameter and 0.42
122	cm height and was slightly curved at the bottom of the cylinder. Synthetic air was used
123	as fluid, so these results can be compared with the results obtained at $N_2:O_2 = 4:1$ using
124	the Mettler Toledo Thermobalance.
125	Dynamic experiments were carried out at heating rates between 5 and 20 K/min, from
126	the initial room temperature up to 850 K, including thus the entire range of
127	decomposition. Isothermal experiments started at a constant heating rate until the
128	desired temperature was reached and then the final temperature was maintained
129	constant. The experiment was considered to have finished when the weight loss rate was
130	negligible (less than 1·10 <sup>-5</sup> s <sup>-1</sup> ). Small size samples, between 1 and 10 mg, were used in
131	the runs.
132	A pyrolysis run at a heating rate of 5 K/min using Avicel PH-105 microcrystalline
133	cellulose was done on each apparatus. The kinetic parameter values obtained showed
134	good agreement with the results reported by Grønli et al. [19] in their round-robin study
135	of cellulose pyrolysis kinetics by thermogravimetry (at 5 K/min and 244 kJ/mol, the
136	experimental and calculated data coincide, obtaining logarithmic values of the pre-
137	exponential factors of around 18.8, a value within the accepted interval). These
138	experiments were useful to check how well the two thermobalances performed.
139	The TG-MS runs were carried out on a Mettler Toledo model TG-ATD
140	TGA/SDTA851e/LF/1600 coupled to a Thermostar GSD301T Pfeiffer Vacuum MS
141	apparatus using $He:O_2 = 4:1$ as carrier gas. The operating conditions were: a mass
142	sample of around 5 mg, a 30 K/min heating rate, a 70 eV ionization energy, and SIR
143	detection of several ions (4, 13-18, 25-32, 35-46 in one run and 4, 32, 43-46, 50-52, 55-
144	58, 60, 65, 68, 73, 78, 91, 96, 105, 106 in another run). The response of each ion was
145	divided by that of helium (m/z= 4) and afterwards the corresponding minimum value
146	was subtracted from each response.
147	The TG-IR runs were carried out on a Perkin Elmer STA6000 and a Nicolet 6700 FT-IR
148	using air as carrier gas, a mass sample of around 12 mg and at 30 K/min heating rate.
149	The transmittance was measured between 4000 and 600 cm <sup>-1</sup> .

150

3. RESULTS AND DISCUSSION

5

151	Most of the 1G runs were carried out on the Mettler Toledo Thermobalance in the
152	aluminium crucible. Figures 1 and 2 show the first experimental TG plots for the
153	combustion of pure olive oil, which must be analyzed in order to understand subsequent
154	runs and the proposed kinetic model. Figure 1a shows the TG runs carried out on the
155	Mettler Toledo (M-T) instrument for combustion and pyrolysis of pure olive oil (data
156	for the latter were obtained elsewhere [3]) at 10 K/min and for a 5 mg initial mass. As
157	observed, the thermal decomposition is faster under oxidative conditions. It is possible
158	that a carbonaceous residue has been formed by oxidation, whose subsequent
159	combustion results in the presence of a fraction in the oxidation run curve on the right
160	of the pyrolysis run curve. That this residue has been possibly formed should be
161	confirmed by means of other techniques, such as TG-MS and TG-IR, since other
162	explanations are also possible. Figure 1b shows the results of three runs carried out at 5,
163	10 and 20 K/min on an initial mass of 5 mg. It can be observed that the curves intersect,
164	which also occurs in other series of runs. The exothermal nature of the combustion run
165	can be confirmed from the variation in the temperature increment of the DTA
166	corresponding to 20 K/min, by noticing that there is an increase in temperature
167	throughout the entire process and a peak that coincides with the weight that has been
168	lost. Figure 1c shows the results of three TG runs carried out under the same operating
169	conditions but varying the initial masses. It can be seen that there is a considerable
170	difference between the experimental curves obtained for 1 and 5 mg on the one hand,
171	and 10 mg on the other. Concerning the experimental data, Figure 1d shows the results
172	of the TG (weight fraction) and DTG (mass fraction increment in volatiles per unit
173	temperature increment, $\Delta V/\Delta T$ ) for a run carried out at 5 K/min on 5 mg of oil. Three
174	peaks are visible at 600, 700 and 800 K (the label "cal" refers to data calculated by
175	means of the proposed model). The results of two other runs carried out under the same
176	operating conditions are shown in Figures 2a and 2b. Small differences between the
177	DTG curves can be observed, which demonstrates again that the runs are not exactly
178	reproducible. Figure 2c shows calculated results for the decomposition steps involved in
179	the reactions that are proposed in the following sections. Figure 2d shows the results of
180	a run carried out at 20 K/min on 5 mg of oil. In light of the previous results, the
181	following aspects deserve comment: a) at least three decomposition steps can be
182	considered based on the presence of three peaks in the DTG runs, b) the run carried out
183	on 10 mg has a curve that is very separated from the curves of the other two runs carried
184	out on 1 mg and 5 mg, probably as a consequence of the large sample mass, which

185	could cause the temperature of the samples to be different to that programmed (the
186	effect of sample mass must be considered to obtain acceptable results) c) there could be
187	a factor that leads to random behavior and provokes crossing of the curves. This can be
188	attributed to a vaporization process, as in the case of the pyrolysis runs [3]. Previous
189	studies have revealed that the vaporization processes exhibit a random variation in
190	weight loss vs. temperature in dynamic TG runs done within the interval of
191	vaporization, which is the result of irregular diffusion of vapours along the length of the
192	crucible [20,21]. No other reasons for the random behavior have been found.
193	Figures 1 and 2
194	Figure 3 shows the results of runs also carried out on the Mettler Toledo
195	Thermobalance, but now on 1 mg of pure olive oil. The DTA in Figure 3a is for the 20
196	K/min run and is indicative of an exothermic process throughout the entire run. The first
197	of the three peaks in the DTG plot in Figure 3c is very broad, so presumably four
198	decomposition steps, two decomposition steps corresponding to the broad peak and two
199	decomposition steps from the following two peaks should be considered when
200	analyzing the experimental data. Figures 3b and 3d show calculated data that will be
201	explained later.
202	Figure 3
203	The analysis of the runs carried out at $N_2:O_2 = 9:1$ atmosphere, the results of the runs
204	carried out with the Perkin Elmer Thermobalance and some results of the dynamic +
205	isothermal runs are presented in Supplementary Material.
206	Figures 4a to c show the TG curves of runs carried out on pure olive oil, waste olive oil
207	and waste mixed oil, respectively. The overall decomposition is similar in all cases in
208	spite of the thermal treatment undergone by the waste oils. Another TG run carried out
209	on waste olive oil at 10 K/min instead of 20 K/min is shown in Figure 4d.
210	Firm 4
210	Figure 4
211	Before turning to a description of the proposed kinetic model, the TG-MS and TG-IR
212	data will be presented and analyzed since they are useful in identifying the different
213	decomposition steps.
214	4 ANALYSIS OF TG-MS DATA

215	Figure 5 shows the results of the TG-MS run carried out on a 5 mg initial mass of pure
216	olive oil at 30 K/min and at a $He:O_2 = 4:1$ atmosphere (the high heating rate was
217	required to obtain acceptable signals for the evolved ions). The intensity of a number of
218	ions have been measured: water (18), carbon monoxide (28, including ethylene), carbon
219	dioxide (44) and methane (15). It can be seen that the ion corresponding to water
220	appears in the interval 500-750 K, coinciding with the thermal degradation of the olive
221	oil, except in the last step. By contrast, ions of both carbon oxides appear throughout the
222	entire decomposition process, from 500 to 850 K, indicating that the last decomposition
223	step corresponds to the combustion of a carbonaceous residue with formation of carbon
224	oxides and very little or no formation of water.
225	Considering the remaining TG-MS data, it seems that formaldehyde (ions 29 and 30),
226	acetaldehyde (ions 29 and 43), ethylene (ions 27 and 26, because ion 28 also
227	corresponds to carbon monoxide), acetylene (ions 26 and 25), other hydrocarbons (ions
228	25, 26, 27, 39, 40 41 and 42) and other oxygenated compounds (ion 57) are formed
229	inside the interval 500-750 K, including methane, as a consequence of the thermal
230	decomposition. The study has been extensive considering all the ions listed in the
231	experimental section. The emission of benzene, toluene or xylenes – ions 78, 91 or 106
232	– was not observed.
233	Figure 5
234	
235	5. ANALYSIS OF THE TG-IR DATA
236	Figure 6 shows the results obtained in the dynamic run carried out on a 12 mg initial
237	mass of pure olive oil at 30 K/min in air, on the Perkin Elmer TG and the Nicolet FT-IR
238	apparatus. Figure 6a shows the transmittance at time 15.5 min, when the weight loss
239	rate is high. The following peaks have been identified in accordance with NIST data
240	base and Vlachos et al. [22]: 3400-4000 cm <sup>-1</sup> and 1300-1600 cm <sup>-1</sup> due to water vapor,
241	2300-2400 cm <sup>-1</sup> and 700 cm <sup>-1</sup> due to CO <sub>2</sub> , 2100-2200 cm <sup>-1</sup> due to CO, 2850-3000 cm <sup>-1</sup>
242	due to C-H bonds, 1700-1800 cm <sup>-1</sup> due to C-O bonds and 1150-1250 cm <sup>-1</sup> due to C-O
243	ester groups.
244	Figure 6

8

- 245 Figure 6b shows the variation of transmittance vs. time corresponding to a wavelength
- of 2930 cm<sup>-1</sup> (C-H bond). Only one broad peak is observed. Similar trends occur in the
- 247 case of the other wavelengths, except for those corresponding to CO and CO<sub>2</sub> (see
- Figure 9c where two broad peaks can be observed). This fact confirms the conclusion
- drawn based on the TG-MS results. The first broad peak in Figure 9c corresponds to
- decomposition reactions, whereas the second broad peak, only observed for CO<sub>2</sub> in
- 251 Figure 6c and CO at its characteristic wavelength, correspond to the combustion of a
- 252 carbonaceous residue accompanied by little or no formation of water and organic
- compounds. Similar trends were obtained in the case of the waste olive oil and waste
- 254 mixed oil.
- 255 6. KINETIC MODEL
- 256 Several kinetic models were considered for the purpose of reproducing the experimental
- 257 results. Since at least four decomposition steps must be taken into account, the
- 258 following scheme of four parallel reactions has been proposed (reactions 1A and 1B
- corresponding to the first broad peak, reactions 2 and 3 for the following peaks in DTG
- 260 runs):

261 Oil 
$$+O_2 \xrightarrow{1A} v_{1A\infty}$$
 volatile fraction + carbonaceous fraction (R1A)

262 Oil 
$$+O_2 \xrightarrow{1B} v_{1B\infty}$$
 volatile fraction + carbonaceous fraction (R1B)

263 Oil 
$$+O_2 \xrightarrow{2} v_{2\infty}$$
 volatile fraction + carbonaceous fraction (R2)

- 264 Carbonaceous fraction +  $O_2 \xrightarrow{3} V_{3\infty}$  volatile fraction (R3)
- where  $v_{1A\infty}$ ,  $v_{1B\infty}$ ,  $v_{2\infty}$  and  $v_{3\infty}$  are the maximum mass fractions of volatile products of
- 266 reactions 1A, 1B, 2 and 3, respectively, having been produced long after the reactions
- 267 had gone to completion. Taking into account that the final residue is negligible, the sum
- of these maximum mass fractions of volatiles must equal 1. The carbonaceous fraction
- in question is the one produced by any of the reactions 1A, 1B or 2.
- 270 This kinetic model is based on the decomposition steps observed and must be
- considered as a simplification of the complicated network of reactions that take place.
- This kinetic model must be considered as a correlation of the experimental data
- obtained.

- For every reaction, the conversion degree is calculated as the ratio of the mass fraction
- of volatiles obtained at any instant during the reaction (V<sub>i</sub>) to the corresponding yield
- coefficient or the mass fraction of volatiles at time infinity  $(v_{i\infty})$ , or

277 
$$\alpha_i = V_i / V_{i\infty}$$
  $i = 1A, 1B, 2 \text{ and } 3$  (1)

278 The kinetic equation of each reaction i can be expressed as

$$d(V_{i}/V_{i\infty})/dt = d\alpha_{i}/dt = k_{i}(1-\alpha_{i})^{n_{i}} = k_{i}(1-(V_{i}/V_{i\infty}))^{n_{i}}$$
 i = 1A, 1B, 2 and 3 (2)

- For reaction 3, the same kinetic model is assumed to apply on the grounds that the
- carbonaceous residue is formed at low temperatures prior to combustion.
- The kinetic constants are obtained from the Arrhenius equation, or

283 
$$k_i = k_{oi} \exp(-E_i / RT)$$
  $i = 1A, 1B, 2 \text{ and } 3$  (3)

- 284 By integrating the above equations, the conversion degrees can be calculated at every
- instant from a knowledge of the temperature program. The weight or mass fraction
- measured in the thermobalance (w) is related to the volatiles obtained (V) by:

287 Mass fraction = 
$$1 - V = 1 - (V_{1A} + V_{1B} + V_2 + V_3) =$$

$$= 1 - (v_{1A\infty}\alpha_{1A} + v_{1B\infty}\alpha_{1B} + v_{2\infty}\alpha_2 + v_{3\infty}\alpha_3)$$
(4)

- Assuming initial values for all the kinetic constants (k<sub>oi</sub>, E<sub>i</sub>, n<sub>i</sub>) and maximum mass
- fractions,  $v_{i\infty}$ , we calculated the conversion degrees by integrating the differential
- 290 equations in Eq. (2) above, using Euler's method and small time intervals, as well as
- optimization with the Solver function in an Excel spreadsheet. We subsequently
- 292 checked that integration by Euler's method was accurate by decreasing the time
- interval, which gave the same results. It has also been confirmed that the kinetic
- 294 parameters obtained by applying the iso-conversional method [23] to a reaction,
- 295 coincide with those employed in the simulations using Euler's method, for small time
- intervals of the same order as those used in this work. The objective function (OF) to
- 297 minimize was the sum of the square differences between the experimental and
- 298 calculated mass fractions:

OF = 
$$\sum_{m=1}^{M} \sum_{j=1}^{N} (mass fraction_{m,j}^{exp} - mass fraction_{m,j}^{cal})^{2}$$
(5)

- where M is the number of runs and N is the number of points in each run.
- The validity of the model has been established by calculating the variation coefficient
- 302 (VC):

$$VC = 100\sqrt{(OF/(N_{total} - P))} / \text{mass fraction}_{exp}$$
(6)

- where  $N_{\text{total}}$  and P are the number of data values and parameters fitted, respectively, and
- mass fraction exp is the average mass fraction that remains inside the crucible, which is
- close to 0.5. In accordance with the approach proposed in Martín-Gullón et al. [24], the
- optimization was performed with respect to a 'comparable kinetic constant', K<sub>i</sub>\*,
- instead of optimizing k<sub>oi</sub> directly. This constant was calculated at a reference
- temperature (T<sub>ref</sub>) around the maximum decomposition rate, after the inclusion of a
- 310 factor  $(0.64)^{ni}$ , as:

311 
$$K_i^* = k_i (0.64)^{n_i} = (k_{oi} \exp(-E_i / RT_{ref}))(0.64)^{n_i}$$
 (7)

- The number 0.64 was introduced to weaken the dependence of the reaction order and
- the other kinetic parameters on each other [24]. From the optimized parameters  $K_i^*$ ,  $E_i$
- and  $n_i$ , the values of  $k_{0i}$  can be deduced. Note that the parameter  $K_i^*$  is only used and
- valid for correlation purposes, since it facilitates optimization and decreases the
- 316 computational time.
- The optimization parameters for reactions 1A, 2 and 3 were  $K_i^*$ ,  $E_i$ ,  $n_i$  and  $v_{i\infty}$ . As for
- reaction 1B, they were  $E_{1B}$ ,  $n_{1B}$ ,  $v_{1B\infty}$  and the value of  $K_{1B}^*$  in each run. The fact that
- $K_{1B}$  varies between runs can be justified if a vaporization process takes place during the
- devolatilization process; it has been established that the pre-exponential factor depends
- 321 on the initial mass in the vaporization process, and that it can vary between similar runs
- due to a random process that depends on the heating rate [20,21].
- To deduce the best kinetic parameters that minimize the objective function so that the
- 324 experimental and calculated TG curves match, the data obtained with the Mettler

325	Toledo 1G apparatus at a $N_2:O_2 = 4:1$ atmosphere were used as initial values. The same
326	set of parameters was used in the runs carried out on the Perkin Elmer TG.
327	Table 2 shows the optimized pre-exponential factors $k_{\text{ol}B}$ obtained in each run and
328	Table 3 shows the kinetic parameters obtained for each reaction. With the optimized
329	parameters, the mass fraction curves were calculated and plotted together with their
330	experimental values, both TG and DTG, in Figures 1 to 4. The same was done in the
331	case of the other tests. It can be seen that the calculated results agree well with the
332	experimental ones in most cases (reason why the small differences cannot be observed),
333	demonstrating that the proposed model is useful for correlating the data. Figures 2c and
334	3d show the variation in volatile mass fraction for the four reactions: reactions 1A and
335	1B take place in the interval of 500-750 K, whereas reaction 2 and 3 take place in 700-
336	800 K and 750-850 K, respectively.
227	The varieties coefficient of each my was relevated value a moon value for the mass
337	The variation coefficient of each run was calculated using a mean value for the mass
338	fraction of 0.5 in all the runs. Table 2 shows these results, where it can be seen that in
339	the $N_2:O_2 = 4:1$ runs the variations are smaller than 10 %, except in the case of the two
340	runs carried out on the Perkin Elmer TG at 5 K/min. This makes us confident about the
341	ability of our kinetic model to correlate the experimental results obtained from the two
342	different TG apparatus. It is worth noting, however, that in the two runs where the VC
343	exceeds 10 %, the experimental conversions are greater than those predicted by the
344	model, and therefore the model is useful to check that a conversion is obtained or
345	surpassed.
346	The following analysis can be done based on the obtained kinetic parameters:
347	- Reaction 1A is the most important and contributes up to 50.2 % of the initial mass,
348	whereas reaction 1B contributes only 18.7%. These two reactions have similar apparent
349	activation energies – around 125 kJ/mol – and their reaction orders are 1.73 for reaction
350	1A, and 1.07 for reaction 1B. The obtained parameters are the result of the best
351	correlation of the data, and consequently they have no clear physical meaning. This fact
352	may indicate that the proposed scheme is an over-simplification of the real process. The
353	relatively low apparent activation energy of reaction 1A and its reaction order of 1.73
354	indicate that there are many consecutive and parallel reactions giving rise to these
355	correlation values

356	-Where reaction 1B is concerned, the vaporization effect together with consecutive and
357	parallel reactions give rise to a reaction order close to 1, a value between zero for
358	vaporization processes and orders greater than unity that can be found in literature for
359	chemical reactions. The activation energy also has a low value, but is greater than that
360	of a volatilization process (30-70 kJ/mol). It is curious that in the pyrolysis of olive oil
361	[3] using the same thermobalance, there was also a first vaporization + reaction process,
362	with a reaction order of 0.606 and an apparent activation energy of 112 kJ/mol, whereas
363	in the case of reaction 1B in the combustion process, the reaction order is 1.07 and the
364	apparent activation energy is 124 kJ/mol, a value close to 112 kJ/mol. Perhaps there is a
365	similarity between the processes of pyrolysis and combustion, with the difference being
366	that the oxidation of the reacting mass gives rise to an increase in the overall reaction
367	rate because of oxygenated radicals.
368	The kinetic parameters of reactions 1A and 1B are comparable with those (reaction
369	order around 1 and activation energy around 85-100 kJ/mol) obtained in the first step of
370	the decomposition proposed by Santos et al. [12] and also comparable to those obtained
371	by Gouveia de Souza et al. [11].
3 / 1	by Gouveia de Bouza et al. [11].
372	- For reaction 2, the apparent activation energy is high, 389 kJ/mol, and so is the
373	reaction order, 3.31. These results have opposing effects: high activation energies mean
374	sharp peaks in a DTG run, whereas high reaction orders mean broad peaks. Perhaps
375	lower activation energies and reaction orders are also acceptable. Nevertheless, the
376	obtained correlation values are optimal – also upon taking into account the other three
377	decomposition steps and all the dynamic and dynamic + isothermal runs. Gouveia de
378	Souza et al. [11] proposed activation energies of 205-300 kJ/mol and reaction orders of
379	2.0 or 2.1.
380	- Reaction 3 corresponds to the combustion of a carbonaceous residue, which is in
381	keeping with the comparison between pyrolysis and combustion runs, and considering
382	the TG-MS and TG-IR results. The activation energy and reaction order are 240 kJ/mol
383	and 1.04, respectively, which are acceptable values for combustion processes. Gouveia
384	de Souza et al. [11] proposed activation energies of 300-380 kJ/mol and reaction orders
385	of around 1.9-2.1 which are similar to those proposed by Santos et al. [12]

- Vecchio et al [14] presented DTG data of triglycerides: tristearate, trioleate, trilinoleate
- and trilinolenate. They observed three steps of decomposition: a first wide one, which
- can be decomposed in two for trioleate, a second step with an acute peak and a third
- step, which corresponds to the burnout of the carbonaceous residue.
- The data presented in Table 3 correspond to a  $N_2:O_2 = 4:1$  and  $N_2:O_2 = 9:1$  atmosphere.
- The same set of parameters was used in the correlation of runs carried out at  $N_2:O_2 =$
- 392 9:1 and  $N_2:O_2 = 4:1$ . However, several aspects are worth commenting:
- The pre-exponential factor k<sub>o1B</sub> of each run has been optimized, as was done in N<sub>2</sub>:O<sub>2</sub>
- 394 = 4:1 runs.
- It seems that less of the carbonaceous residue forms than in the case of  $N_2: O_2 = 4:1$
- runs, so that the mass fractions of the other reactions (1A, 1B and 2) increase, as shown
- 397 in Table 3.
- For the runs carried out on 5 mg samples, the pre-exponential factor of reactions 1A, 2
- and 3 decrease with respect to  $N_2:O_2 = 4:1$  runs by a factor of 0.32, which is obtained
- 400 experimentally by optimization when the corresponding experimental data are
- 401 correlated. This factor corresponds to a reaction order of 1.64 with respect to the oxygen
- 402 partial pressure, and was calculated as follows:

403 
$$\frac{\log 0.32}{\log \left[ \frac{P_{O2} \text{ for } N_2 : O_2 = 9 : 1}{P_{O2} \text{ for } N_2 : O_2 = 4 : 1} \right]} = \frac{\log 0.32}{\log \left[ \frac{0.1}{0.2} \right]} = 1.64$$
 (8)

- This reaction order is greater than unity probably as a consequence of diffusion of
- 405 oxygen inside the crucible, which causes the oxygen concentration in the surface of the
- oil to be less than the external oxygen concentration.
- For the runs carried out on 1 mg samples, the pre-exponential factors of reactions 2
- and 3 decrease by the same factor, 0.32. However, for reaction 1A, the pre-exponential
- factor is the same as in the  $N_2:O_2 = 4:1$  run. This would indicate that in the case of
- reaction 1A, the oxygen is probably required as an initiator in oxygenated radical
- 411 formation and as a reactant. For the runs carried out on 1 mg and 5 mg samples at N<sub>2</sub>:O<sub>2</sub>
- 412 = 4.1, and on 1 mg at  $N_2:O_2 = 9.1$ , there is sufficient oxygen present to achieve the
- 413 maximum degradation rate, whereas for 5 mg at  $N_2:O_2 = 9:1$  there is not, and thus the

414	reaction proceeds more slowly. All these considerations highlight the complexity of the
415	process.
416	Figure 4 shows the experimental and calculated results obtained for pure olive oil, waste
417	olive oil and waste mixed oil by means of the same correlation procedure. This means
418	that approximately the same kinetic model can be applied, although for certain waste
419	mixed oils several runs should be done to confirm or modify the kinetic parameters and
420	the mass fraction of volatiles involved in each reaction.
421	The values of the pre-exponential factor $k_{o1B}$ have been correlated roughly by means of
422	a parameter P, which is defined as:
423	P = (initial mass in mg) <sup>a</sup> ·(heating rate in K/min) <sup>b</sup> ·(height in cm) <sup>c</sup> ·(diameter in cm) <sup>d</sup>
424	The optimal values of a, b, c and d that obtain the best correlation between $k_{o1B}$ and P
425	are as follows: $a=-2.3$ , $b=-0.68$ , $c=-3.5$ and $d=16.0$ . A logarithmic plot of $k_{o1B}$ vs. P is
426	shown in Figure 7. The values of exponents a,b,c,d reveal a trend in the variation of
427	process 1B, when vaporization is included. If the process were only vaporization of a
428	pure substance in a pure molecular diffusion process, the expected values would be the
429	following: a= -1; b=0; c=-1; d=2. The obtained values differ from these, but the positive
430	and negative values follow the expected trend. The convective phenomena produced by
431	temperature gradients, the formation of small drops at the end of the run and the shape
432	of the crucible, which is not exactly cylindrical, can alter molecular diffusion inside the
433	crucible, which is one of the factors controlling the vaporization rate [20,21].
434	Figure 7
435	
436	7. APPLICATION OF THE KINETIC MODEL
437	The system of equations that we have deduced is useful to characterize the
438	decomposition of olive oil, waste olive oil and more approximately the decomposition
439	of waste mixed oil. However, there are some aspects that merit consideration:
440	1. The mass fraction of the last reaction (combustion of the carbonaceous material) can
441	depend on the combustion conditions and vary between 0.13 for $N_2:O_2=4:1$ and 0.08
442	for $N_2:O_2 = 9:1$ . For intermediate conditions, an interpolation can be done. The mass

- fraction of the other reactions must be recalculated so that the sum of all fractions is
- 444 equal to unity.
- 2. The value of the pre-exponential factor  $k_{o1B}$  depends on operating conditions, so
- extrapolation of the TG data to industrial conditions may be risky. A first approximation
- would imply assuming that reactions 1A and 1B are similar, so that the kinetic
- parameters of reaction 1A can be used for the sum of the mass fractions of reactions 1A
- and 1B. An analysis of the operating conditions of the industrial process can also be
- done to estimate the equivalent diffusion length (mass transfer coefficient/diffusivity).
- 451 This can be compared with the height of the crucibles that are used, in order to establish
- 452 whether the vaporization process implied in reaction 1B is faster or slower than reaction
- 1A. In any case, the proposed approximation may be valid.
- 454 8. CONCLUSIONS
- Four decomposition steps have been suggested for correlating the complex system of
- reactions involved in the combustion of olive oil. Reactions 1A and 1B take place at
- 457 500-750 K, reaction 2 at 700-800 K and reaction 3 at 750-850 K. In reaction 1B, which
- 458 corresponds to a vaporization + reaction process, the observed random behavior is
- deduced to be the result of the vaporization process. The last reaction corresponds to the
- 460 combustion of a carbonaceous residue.
- The obtained kinetic parameters have been instrumental to satisfactorily simulating the
- 462 experimental results.
- The kinetic model might also be roughly applicable to waste olive oil and waste mixed
- oil, although where waste mixed oil is concerned, more runs should be done to confirm
- or vary the kinetic parameters.
- 466 The kinetic study carried out on different initial masses using two distinct TG apparatus
- 467 together with TG-MS and TG-IR data, is useful for analyzing the thermal behavior of
- 468 liquids and for explaining a number of random results in the TG and DTG data.
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557	initial masses of pure olive oil at different heating rates.
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560	pure olive oil at different heating rates (in Figure 2c: 1A, 1B, 2 and 3 is the weight
561	fraction of volatiles evolved in reactions 1A, 1B, 2 and 3).
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563	Figure 3. Variation of weight fraction vs. temperature for runs carried out on 1 mg of
564	pure olive oil at different heating rates (in Figure 3d: 1A, 1B, 2 and 3 is the weight
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566	
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570	carried out on pure olive oil.
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573 574 575 576 577	<b>Figure 7.</b> Variation of k <sub>o1B</sub> vs. parameter P.

A kinetic model, including vaporization and reaction, is proposed

Two TG apparatus have been used, so the results can be compared

The kinetic model is supported by TG-MS and TG-IR runs.

	Carbon	Hydrogen	Oxygen	Nitrogen	Sulfur	Net calorific value	
	(%)	(%)	(%)	(%)	(%)	(kcal/kg)	
Olive oil	77.5	11.6	10.9	0.0	0.0	8884	
Waste olive oil	77.2	11.6	11.2	0.0	0.0	8859	
Waste Mixed oil	76.9	11.6	11.5	0.0	0.0	8784	

	N <sub>2</sub> :O <sub>2</sub> ratio	M (mg)	Run	Heating rate (K/min)	TG Apparatus	Crucible Material	Height (cm)	Int. Diam. (cm)	k <sub>01B</sub> (s <sup>-1</sup> )	Parameter P	VC (%)
Olive oil	4:1	4.791	D	20	Mettler Toledo	Aluminium	0.41	0.55	$7.324 \cdot 10^8$	2.18·10 <sup>-7</sup>	4.1
Waste olive oil	4:1	6.187	D	20	Mettler Toledo	Alumina	0.425	0.47	$3.481 \cdot 10^7$	$1.56 \cdot 10^{-7}$	7.1
Waste mixed oil	4:1	5.643	D	20	Mettler Toledo	Alumina	0.425	0.47	$1.361 \cdot 10^7$	1.96·10 <sup>-7</sup>	7.9
Olive oil	4:1	5.024	D	10	Mettler Toledo	Aluminium	0.41	0.55	$1.354 \cdot 10^7$	$2.19 \cdot 10^{-7}$	4.1
Waste olive oil	4:1	5.020	D	10	Mettler Toledo	Aluminium	0.41	0.55	$2.513 \cdot 10^7$	$2.19 \cdot 10^{-7}$	4.7
Olive oil	4:1	4.996	D	10	Mettler Toledo	Aluminium	0.41	0.55	$2.468 \cdot 10^7$	$2.22 \cdot 10^{-7}$	4.7
Olive oil	4:1	5.242	D	5	Mettler Toledo	Aluminium	0.41	0.55	$7.257 \cdot 10^7$	$2.22 \cdot 10^{-7}$	5.1
Olive oil	4:1	1.120	D	20	Mettler Toledo	Aluminium	0.41	0.55	$4.682 \cdot 10^9$	$7.96 \cdot 10^{-6}$	4.7
Olive oil	4:1	1.095	D	10	Mettler Toledo	Aluminium	0.41	0.55	$5.945 \cdot 10^9$	$9.49 \cdot 10^{-6}$	7.2
Olive oil	4:1	1.047	D	5	Mettler Toledo	Aluminium	0.41	0.55	$7.508 \cdot 10^9$	$1.20 \cdot 10^{-5}$	6.9
Olive oil	4:1	5.025	D + I	10	Mettler Toledo	Aluminium	0.41	0.55	$2.709 \cdot 10^7$	$2.18 \cdot 10^{-7}$	4.8
Olive oil	air	5.205	D	20	Perkin Elmer	Alumina	0.645	0.42	$1.104 \cdot 10^{10}$	$4.29 \cdot 10^{-5}$	9.3
Olive oil	air	5.051	D	10	Perkin Elmer	Alumina	0.645	0.42	$5.418 \cdot 10^9$	$5.21 \cdot 10^{-5}$	6.8
Olive oil	air	5.138	D	5	Perkin Elmer	Alumina	0.645	0.42	$1.889 \cdot 10^{10}$	5.63·10 <sup>-5</sup>	19.3
Olive oil	air	10.096	D	20	Perkin Elmer	Alumina	0.645	0.42	$8.099 \cdot 10^8$	$8.32 \cdot 10^{-6}$	4.4
Olive oil	air	10.808	D	10	Perkin Elmer	Alumina	0.645	0.42	$1.788 \cdot 10^8$	$7.92 \cdot 10^{-6}$	7.5
Olive oil	air	10.855	D	5	Perkin Elmer	Alumina	0.645	0.42	$7.504 \cdot 10^9$	$8.83 \cdot 10^{-6}$	7.7
Olive oil	air	10.115	D+I	20	Perkin Elmer	Alumina	0.645	0.42	$7.804 \cdot 10^8$	$8.28 \cdot 10^{-6}$	2.9
Olive oil	air	10.966	D+I	10	Perkin Elmer	Alumina	0.645	0.42	$7.466 \cdot 10^9$	$7.64 \cdot 10^{-6}$	8.8
Olive oil	air	5.152	D + I	5	Perkin Elmer	Alumina	0.645	0.42	$4.058 \cdot 10^{10}$	5.59·10 <sup>-5</sup>	21.5
Olive oil	9:1	5.081	D	20	Mettler Toledo	Aluminium	0.41	0.55	$5.728 \cdot 10^7$	$1.89 \cdot 10^{-7}$	8.5
Olive oil	9:1	5.006	D	10	Mettler Toledo	Aluminium	0.41	0.55	$4.634 \cdot 10^7$	$2.20 \cdot 10^{-7}$	4.2
Olive oil	9:1	4.954	D	5	Mettler Toledo	Aluminium	0.41	0.55	$4.237 \cdot 10^7$	$2.55 \cdot 10^{-7}$	3.9
Olive oil	9:1	1.113	D	20	Mettler Toledo	Aluminium	0.41	0.55	$1.300 \cdot 10^8$	$8.09 \cdot 10^{-6}$	9.3
Olive oil	9:1	1.062	D	10	Mettler Toledo	Aluminium	0.41	0.55	$1.273 \cdot 10^8$	1.02·10 <sup>-5</sup>	4.9
Olive oil	9:1	1.156	D	5	Mettler Toledo	Aluminium	0.41	0.55	$2.781 \cdot 10^9$	$9.35 \cdot 10^{-6}$	14.0

Reaction	Temperature Interval (K)	k <sub>i0</sub> (s <sup>-1</sup> ) for N <sub>2</sub> :O <sub>2</sub> =4:1 and air	$k_{i0} (s^{-1})$ for $N_2: O_2 = 9:1$	E <sub>i</sub> (kJ/mol)	$\mathbf{n_i}$	for N2:O2 = 4:1	$v_{i\infty}$ for $N_2$ : $O_2$ = 9:1
1A	520-700	5.979·10 <sup>8</sup>	$0.32 \cdot 5.979 \cdot 10^{8}$ (for 5 mg initial mass) $5.979 \cdot 10^{8}$ (for 1 mg initial mass)	127.3	1.73	0.502	0.532
1B	500-750 (depending on operating conditions)	see Table 2	see Table 2	124.2	1.07	0.187	0.198
2	700-800	$3.788 \cdot 10^{27}$	$0.32 \cdot 3.788 \cdot 10^{27}$	389.3	3.31	0.178	0.188
3	750-850	$5.040 \cdot 10^{13}$	$0.32 \cdot 5.040 \cdot 10^{13}$	240.7	1.08	0.132	0.080











