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Volatile profiles of extra virgin olive oil, olive pomace oil, soybean oil and palm oil in different heating conditions Giuffrè A.M.*, Capocasale M., Macrì R., Caracciolo M., Zappia C., Poiana M. Università degli Studi Mediterranea di Reggio Calabria, AGRARIA - Dipartimento di Agricoltura, Risorse forestali, Ambiente Risorse zootecniche, Ingegneria agraria, Alimenti - Contrada Melissari, 89124 - Reggio Calabria, Italia. *Corresponding author: amgiuffre@unirc.it Università degli Studi 'Mediterranea' di Reggio Calabria, Italy - Dipartimento di AGRARIA.

16 ABSTRACT

Extra virgin olive oil from Calabria Region (South Italy), pomace olive oil, palm oil and 17 soybean oil were heated at 180 and 220°C for 30, 60 and 120 mins. Emission of 18 19 volatile organic compounds produced during the heat treatment was evaluated by the 20 solid phase micro-extraction of the head space technique and analysed by gas chromatography with mass spectrometric detector. The results after oil heating showed 21 a significant variation in the volatile composition with temperature and time increasing. 22 23 Twenty-five compounds have been identified, and alkanals, alkenals, and alkadienals 24 were the most represented classes. Results from volatile profile evolution suggest that 25 specific aldehyde compounds can be used as markers both for the evaluation of the 26 organoleptic properties and for the health characteristics of the studied oils.

27

28 Keywords

- 29 Deep-fried oil, volatile organic compounds, SPME, GC-MS.
- 30

32 **1.** Introduction

33 One of the many uses of an edible vegetable oil is in deep-frying processes in food preparation. Deep-frying is one of the oldest cooking methods (Jaarin & Kamisah, 34 35 2012) which give a characteristic taste to the food. It is highly valued by the consumers and easy and fast to do. When a vegetable oil is heated, changes in physical 36 appearance occur such as darkening in colour and increasing in viscosity due to the 37 alteration of the fatty acid composition (Rani et al., 2010). Heating causes some 38 chemical reactions in the oil such as oxidation, hydrolysis and polymerization (Choe & 39 40 Min, 2007). During these processes, many oxidative products such as hydroperoxides and aldehydes are produced, which can be absorbed into the fried food (Choe & Min, 41 2006). 42

43 In Italy, one of the main agricultural crops is represented by the olive tree, mainly for 44 extra virgin olive oil (EVOO) production. The flavour in an olive oil is very important and it was widely studied (Vichi et al., 2003; Cavalli et al., 2004; Baccouri et al., 2007; 45 Baccouri et al., 2008;). It is characterised by a large number of carbonyl substances, 46 47 which includes linear saturated and unsaturated aldehydes and alcohols, esters and 48 hydrocarbons (Flath et al., 1973; Esposto et al., 2009;). This composition can be affected by geographical origin (Ouni et al., 2011), by ripening degree (Konavouras et 49 al., 2005) and by extractive processing method (Dabbou et al., 2012). Olive pomace is 50 51 the most important by-product in the olive oil extraction industry, in particular when the 52 three-phase technology is used (Nasopoulou et al., 2011). The three phases system 53 (oil, olive pomace and olive mill waste water) produces an olive pomace with a low 54 moisture content which facilitates the pomace olive oil (PO) solvent extraction by nhexane (Ciftci et al., 2009; Mele et al., 2018). Palm oil (P) and soybean oil (SO) are 55

56 rich in antioxidants such as tocopherols and tocotrienols (Kamisah et al., 2005; Clemente & Cahoon, 2009) and are the most commonly used vegetable oils in the 57 household and industrial deep frying purposes. Each oil produces different volatile 58 59 emissions in different cooking methods (Peng et al., 2017). Many studies were 60 conducted on these and others vegetable oils about the variation in the chemical composition during heating, in particular on its oxidative stability (Saad et al., 2007; 61 Maggio et al., 2011; Jaarin & Kamisah, 2012; Poyato et al., 2014; Qing et al., 2016; 62 Giuffrè et al., 2017a; Giuffrè et al., 2017b; Giuffrè et al., 2018). A minor number of 63 64 studies exist on their volatile profile after heating (Katragadda et al., 2010; Peng et al., 2017). 65

The most rapid and accurate technique to study the volatile fraction of an oil is the SPME (Song et al., 1997) introduced since 90s for the aroma analysis of various food products (Arthur & Pawliszyn, 1990). It is based both on the capacity of the silica fibre to absorb the analytes released in head space from the sample and to desorb the same molecules in an instrument, such as gas chromatography with mass spectrometric detector (GC-MS).

VOCs produced by cooking were also found to be an indoor pollutant both in household and in food factory, in particular if the ventilation is not enough (Jones, 1999). Outdoor cooking is a problem in volatile organic compounds (VOCs) emission even if lower than for indoor cooking (Edwards et al., 2017). Studies on Chinese women have demonstrated that the exposure to fumes of cooked oils is associated with the lung cancer (Zhao et al. 2018). For this reason is important to know the emission composition in VOCs during oil frying.

79 The aim of this work was to study the effect of temperature and heating duration on the

- volatile organic compound (VOC) composition of extra virgin olive oil, olive pomace oil,
- 81 soybean oil and palm oil.
- 82 2. Materials and method

83 **2.1. Materials**

EVOO and PO were produced in the Calabrian region (South of Italy) in the harvest
year 2016-2017. SO and P were purchased in a supermarket EVOO was obtained by a
mechanical extraction with a three phase apparatus. PO, SO and P were obtained by
solvent (*n*-hexane) extraction.

Nine 1L bottles of the each oil were randomly taken for analysis. After this, three 88 representative batches each one composed by the mixed content of nine bottles were 89 90 prepared for each oil to analyse VOCs in triplicate. At this point, from each batch were 91 prepared 7 glass pyrex beakers, containing 150 g oil each one; 6 out of 7 contained the oil to be heated for the six different treatments: 180 °C (30, 60 and 120 min), 220°C 92 (30, 60 and 120 min) and 1 out of 7 contained the oil to be analysed as a fresh oil 93 94 (control). Beakers containing oils were placed on a heating plate and temperature was adjusted and controlled by a thermometer. 95

96 **2.2.** Solid phase micro-extraction of the head space (HS-SPME)

The most rapid and accurate technique to study the volatile fraction of an oil is the HS-SPME (Song et al., 1997) introduced since 90s for the aroma analysis of various food products (Arthur & Pawliszyn, 1990). A SPME fibre coated with poly-dimethylsiloxane (PDMS, 100 μ m) from Supelco (Bellafonte, PA, USA) was used to absorb the VOC's with the procedure described by Temime et al. (2006) modified as follow. A 10 g aliquot of each heated and unheated (fresh) oil was placed in a 20 ml glass vial with septum

103 cup and provided with a magnetic stirrer. The fibre was conditioned before using at 104 270°C for 30 mins as recommended by the manufacturer. Sample was placed in a 105 thermostatic bath at 45°C in stirring (500 rpm). The fibre was inserted into the vial's HS. 106 Absorption was stopped after 20 mins and fibre was thermally desorbed for 5 mins in a 107 gas chromatographer in split-less mode. All SPME operations were conducted 108 manually.

109 2.3. GC-MS analysis

A GC Thermo Trace 1310 apparatus (Waltham, MA, USA) equipped with Single 110 111 Quadrupole Mass Spectrometer ISQ LT system and a fused-silica capillary columns (30 m length, 0.25 mm i.d., 0.25 µm film thickness, Thermo Scientific, Whaltam, MA, 112 USA), TG-5MS 5% phenyl phase was used for GC-MS analysis. The oven 113 114 temperature was programmed at 60°C held for 10 mins, then from 60°C to 220°C at 115 5°C/mins, and held isothermally at 220°C for 10 mins. Injector, MS transfer line and ion source temperatures were respectively 250, 270 and 260°C. Mass range was from 45 116 to 500 atomic mass unit (amu). Gas carrier was helium at 1.5 mL/mins flow rate. The 117 tentative identification of the VOCs was based on the comparison of spectra with those 118 119 of NIST/EPA/NIH Mass spectral library Version 2.0.

120 2.4. Statistical analyses

Each sample was analysed in triplicate and volatile profiles aroma data were submitted to analysis of variance (ANOVA) using SPSS 20.0 software for Windows (SPSS Inc., Chicago, IL, U.S.A.). Wherever F values were significant, Tukey's test was used to separate the mean effects (different temperatures for each oil type). Significance was defined at P < 0.001. Graphics were done using Excel 2010 software for Windows (Microsoft Corp., Redmond, WA, USA).

127 **3.** Results and discussion

128 **3.1.** Volatile organic compounds profile

129 The highest number of compounds, i.e., twenty-one was found in EVOO (Table 1). A similar result was obtained both by Temime et al. (2006) in Tunisian extra virgin olive 130 131 oils, in which were found twenty-five compounds and by Cavalli et al. (2004) in nine 132 virgin olive oils from France and Spain, in which were found a number of VOCs ranging 133 between 23 and 25. A lower number of compounds (twenty) was found in three Algerian EVOOs (Cherfaoui et al., 2018). It has to be pointed out that other authors, in 134 135 different cultivars, have identified a greater number of compounds with the same 136 technique (Hammouda et al., 2017) or with a different technique (Lukić et al., 2019).

The E-2-hexenal has been found to be the most represented constituent (28.3% in 137 138 EVOO) (Table 1), in fact it is considered as a quality fresh-marker of an olive oil 139 (Cavalli et al., 2004), it gives the characteristic sensory properties of sweet, fragrant, almond, green and leafy to the olive oil (Kanavouras et al., 2005). During heating a 140 very high significant decrease in E-2-hexenal content was observed to the lowest value 141 (1.7%) in the EVOO 220°C-120 min sample. The same decreasing trend in E-2-142 143 hexenal was found in almost all the studied oils with the exception of PO 180°C-30 min 144 (Tables 1-4). The higher the heating temperature and the longer the heating duration, 145 the lower the E-2-hexenal content. Other compounds characterising the volatile profile 146 of EVOO were α -farnesene (15.3%), α -cubebene (14.4%) and nonanal (8.9%) (Table 147 1). They are characteristic for an EVOO composition and it was demonstrated to be 148 affected by the olive cultivar (Reboredo-Rodríguez et al., 2012) and the olive growing 149 area (Temime et al., 2006). Some aldehyde compounds such as Z-2-heptanal, E-2octenal, 2-nonenal and 2-decenal (Z), which were not in EVOO, were formed during 150

151 heating, due to oleic acid oxidation (Belitz et al., 2009), and could be used as indicators of the oil heating treatment. The Z-2-decenal alkenal was not detected in EVOO, 152 153 however it was found in high quantity in all the heated samples, and increased 154 constantly with heating to 16.3% in EVOO 180°C-30 min and to 30.8% in EVOO 155 220°C-30 min (Table 1). Other minor compounds were not in fresh EVOO but they were identified after heating: butyl-cyclopentane (found in EVOO heated at 220°C), Z-156 157 2-heptenal, 1-heptanol, 1-octen-3-ol (in EVOO 220°C-60 min), 5-hepten-2-one, 6methyl- (2.0 and 2.2% in both EVOO heated at 180°C-60 min and 180°C-120 min), E-158 159 2-octenal (2.5 and 2.6% in EVOO 180°C-120 min and 220°C-120min respectively), 1octanol (in highest quantity, 1.6%, in EVOO 220°C-60 min and in EVOO 220°C-120 160 min), 2-nonenal which was mainly produced at 220°C after 120 min heating. This was 161 162 due to the oxidation mechanism of the oil, accelerated by the heating (Fullana et al., 163 2004; Lin & Liou, 2000), that causes a chemical primary oxidation and the consequent production of hydroperoxides (Choe & Min, 2006), and its successive degradation to 164 different class of compounds (aldehyde, ketons, alcohols, etc.). 165

Aldehydes can possess toxic properties, in particular, many studies have been conducted on E,E-2,4 decadienal (Chang et al., 2005; Chang & Lin, 2008; Halvorsen & Blomhoff, 2011), considered either carcinogens or suspected carcinogens. It was categorized into group 2A by the International Agency of Research on Cancer, which estimated a daily limit in human diet to 5 mg/kg of body weight (Conklin et al., 2010; Wang et al., 2008).

The major constituent detected in the PO volatile fraction was decanal (42.54%), followed by E-2-hexenal (16.01%), 2-undecenal (15.70%) and nonanal (12.43%) (Table 2). In PO there was a smaller decrease in E-2-hexenal content than in EVOO, which

was lowest (7.76%) in PO180°C-120 min. Interesting was the highly significant
decrease in decanal in all the heated samples, which achieved a value of 2.24% in PO
220°C-120 min and which can be taken into account as the oil temperature stress
index parameter. A similar conclusion, but in reverse, can be done for Z-2-decenal
passed from 5.02% in PO to 21.67% in PO 220°C-120 min.

In P were found seven volatiles compounds: the highest was nonanal (32.75%) and the 180 lowest was Z-2-decenal (6.82%) (table 4). In P heated samples, heptanal, octanal and 181 decanal showed a highly significant decrease. These compounds passed respectively 182 183 from 14.41%, 13.12% and 10.84% to 1.37% (P 180°C-120 min), 1.92% (P 220°C-60 min and P 220°C-120 min) and 1.47% (P 220°C-60 min). Nine new compounds were 184 formed during P heating. The E,E-2,4-decadienal was the most represented in all the 185 186 heated samples, mainly in P180°C-120 min (32.35%), followed by Z-2-heptenal in P 187 180°C-30 min (8.68%) and E,E-2,4-decadienal in P 180°C-120 min (6.44%), (Table 3). The formation of these compounds is due to the oxidation after heating of the 188 unsaturated fatty acids: the higher the unsaturated fatty acid content, the higher the 189 aldehydes formation (Peng et al., 2017). This because poly-unsaturated fatty acids 190 191 such as linoleic and linolenic acids are more susceptible to oxidation because of presence of multi double-bonds (Schauer et al., 2002). In fact, SO, considered a 192 193 linolenic acid-rich oil showed higher concentration in the aldehyde compounds, with 194 E,E-2,4-dodecadienal found as the highest (58.73%) in SO 180°C-120 min (Table 4). 195 Nonanal (34.97%) and decanal (34.19%) were the most represented VOCs in fresh 196 SO. Decanal constantly decreased with heating and with heating duration, whereas 197 nonanal showed an initial increase at 180°C-30 min and a subsequent decrease with heating and with heating duration (Table 4). Furthermore, fresh SO was the one with 198

the lowest number of identified compounds, moreover it showed a greater number of VOCs in the heated samples than in the fresh ones, confirming the high instability of this vegetable oil under heating (Table 4). The SO instability was also confirmed by the lowest total phenolic content (15 mg gallic acid/kg) and by the lowest antioxidant activity quantified with both the lowest values in the ABTS assay (51.8 μ M TE/100g) and the lowest values in the DPPH hydrophilic assay (6.6 μ M TE/100g) measured in the same samples studied by Giuffrè et al., (2018).

206 **3.2.** Total alkanals, alkenals and alkadienal

The identified aldehyde compounds have been grouped as alkanals, alkenals and alkadienals to study the relationship between the heating temperatures, the heating time and their production in the studied vegetable oils.

210 The total aldehydes rate varied significantly during heating of all of the four studied oils 211 both at 180°C and 220°C (Figure 1a,b). This was due to the presence of some alkans (α -cubebene, α -muurolene and α -farnesene) (Table 1) in the EVOO, the only oil 212 obtained from a mechanical process which minimally alters the olive original volatile 213 214 profile, while they are totally absent in all the other three studied unheated vegetable oils obtained by solvent extraction. In figures 1a and 1b a significant increment is 215 216 shown by the total alkenals, which increased from 32.1% in EVOO to 60.9% and 68.8% respectively in EVOO 180°C-120 min and EVOO 220°C-120 min, confirming how the 217 218 formation of these compounds is strictly correlated to the oleic acid oxidation, the major 219 acid constituent in olive oil.

The PO heated at 180°C showed a significant difference in the total aldehydes with respect to PO control with a decrease at 180°C-30 min and a tendency increase at 180°C-60 min and 180°C-120 min (Figure 2a), this was probably due to an initial

degradation of some aldehydes and a subsequent formation of new ones, while the 224 220°C heated samples showed a significant difference after 30 mins heating with a 225 decrement in total alkanals (-57.1%) and an increment in total alkenals (+58.6%) after 226 120 mins heating (Figure 2b).

The total aldehyde profile of P showed a similar trend in both heating temperature, with a significant increment in total alkadienal compounds, from 0% in P to 38.79% and 25.40% respectively in P 180°C-120 min and P 220°C-120 min, and a decrement in total alkanal compound, from 71.13% in P to 29.81% and 28.16% respectively in P 180°C-120 min and P 220°C-120 min (Figure 3a,b). Total alkanals and total alkenals of P varied significantly with heating and with heating duration (Figure 3a,b).

The total aldehydes in SO showed a trend similar to P, in fact total alkanals decreased when total alkadienal increased during heating time at each heating temperature (figure 4a,b). A very significant difference was found in the sample before heating and after heating at 180°C for 60 mins, with a decrement in alkanals (-80.0%) and an increment in alkadienal of +867.2%; while the sample heated at 220°C showed the highly difference from its unheated oil immediately after 30 mins, -90.1% and +916.2% respectively in total alkanals and total alkadienals.

240 **4. Conclusion**

Results showed that temperature and heating duration have a very high significant influence in the volatile organic compounds content of edible vegetable oils. **EVOO** showed the highest number of components with E-2-hexenal as the highest in quantity in fresh oil and Z-2-decenal and 2-undecanal as the highest in quantity in heated extra virgin olive oils. **PO** (control) contained decanal in higher quantity and after heating Z-2decenal and 2-undecanal prevailed as for extra virgin olive oil in addition to nonanal.

Nonanal prevailed in both fresh P and SO whereas E,E-2,4-dodecadienal prevailed in both P and SO oils after their heating. Heating completely varied the volatile organic compounds content of four of the edible vegetable oils mostly used in the world. By the results of this study, the oil user can decide the temperature and the heating duration to be applied to preserve flavours and to reduce off-flavours which are produced during heating in the EVOO, PO, P and SO.

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