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Design and in-house validation of a portable system for the determination of free acidity in virgin olive oil

This is the submitted version (pre peer-review, preprint) of the following publication:

Published Version: Design and in-house validation of a portable system for the determination of free acidity in virgin olive oil / Grossi M.; Palagano R.; Bendini A.; Ricco B.; Servili M.; Garcia-Gonzalez D.L.; Gallina Toschi T.. - In: FOOD CONTROL. - ISSN 0956-7135. - ELETTRONICO. - 104:(2019), pp. 208-216. [10.1016/j.foodcont.2019.04.019]

Availability: This version is available at: https://hdl.handle.net/11585/700389 since: 2019-09-25

Published:

DOI: http://doi.org/10.1016/j.foodcont.2019.04.019

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1	Design and in-house validation of a portable system for the determination		
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## 18 ABSTRACT

Nutritional and healthy values are well known properties of virgin olive oil (VOO). The product quality, in terms of belonging to a specific quality grade (extra virgin, virgin, lampante), is defined by a set of chemical-physical and sensory measurements. According to the official regulation of the European Union (EU Reg. 1348/2013) the free acidity is the first parameter that has to be determined by analysts; it gives information about the quality of the olives used to produce the VOO as well as the hydrolytic state of VOO just produced and stored. The official procedure is based on an acid-base titration that needs to be carried out in a chemical laboratory.

In this paper a portable battery-operated electronic system to measure olive oil free acidity is presented: the system can be used for quick "in situ" tests in a production environment (olive oil mills or packaging centers) by people without particular training. The working principles of the system is based on the creation of an emulsion between oil and a hydroalcoholic solution: the free acidity is estimated on the value of the emulsion electrical conductance.

This new system has been calibrated and in-house validated showing good results in terms of limit of detection and quantification, precision and accuracy, beyond a good correlation with free acidity data obtained applying the official method ( $R^2_{adi} = 0.97$ ).

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Keywords: portable system; free acidity; virgin olive oil; impedance spectroscopy; in-house
 validation; electrical conductance.

37

#### 38 Abbreviations

EIS: electrical impedance spectroscopy, EVOO: extra virgin olive oil, FA: free acidity, LOD:
limit of detection, LOO: lampante olive oil, LOQ: limit of quantification, RSD: relative standard
deviation, TAGs: triglycerides, VOO: virgin olive oil.

#### 42 1. Introduction

Virgin Olive Oil (VOO) is obtained from olives (the fruits of Olea europaea L.) applying only a 43 mechanical-physical extraction process and represents a product highly appreciated for its 44 beneficial effects on human health, mainly due to a high content of oleic acid and minor 45 components such as phytosterols, carotenoids, tocopherols and hydrophilic phenols (Bendini et al., 46 2007). The European Commission Implementing Regulation 1348/2013 (EU Reg. No 1348/2013) 47 defines a decision tree for verifying whether a VOO is consistent with the category declared, and 48 the quality criteria that have to be checked by analysts are: free acidity (FA), peroxide value, 49 50 specific extinctions in UV, sensory characteristics and ethyl esters of fatty acids. The first quality parameter in the above cited decision tree is, therefore, the determination of the free acidity of the 51 52 oil sample; this is defined as the amount of free fatty acids, no longer linked to their parent triglyceride molecules (TAGs), and measured as percentage of oleic acid. Specifically, the top-53 54 quality product, Extra Virgin Olive Oil (EVOO), features a maximum FA of 0.8 g oleic acid/100 g oil, then the VOO features a maximum FA value of 2.0 g oleic acid/100 g oil and, finally, the 55 56 Lampante Olive Oil (LOO), that is not suitable for the commercialization as it is, is characterized by a FA higher than 2.0 g oleic acid/100 g oil. 57

This parameter is especially affected by the quality of the olives used to produce the oil since free 58 fatty acids arise from the separation of fatty acids from TAGs because of the action of enzymes, 59 further stimulated by light, water, and heat. Thus, FA is an indicator of how fresh and how well 60 handled the olives were before being milled (Tena, Wang, Aparicio-Ruiz, García-Gonzaléz, & 61 Aparicio, 2015). To confirm this, different studies reported that geographic and environmental 62 factors (Bustan et al., 2014) and the application of specific technological processes (such as 63 filtration or a cooling treatment of olive paste) (Veneziani et al., 2018 a-b) do not affect 64 significantly this parameter. 65

The official procedure to measure the oil free acidity is defined by the European regulations (EEC Reg. 2568/1991 and following amendments) and consists of an acid-base titration that, albeit simple and quick, must be carried out in a laboratory by trained personnel essentially due to the need of specific solvents and lab equipment.

However, in addition to official methods for the quality control of virgin olive oils, there is a strong need for simple, rapid and environmental friendly techniques, also suitable for on-site quality control even for new clients, who are "non-professional analytical skilled" end users (Inajeros-García, Gómez-Alonso, Fregapane, & Salvador, 2013; Valli et al., 2016). This represents an important issue in particular for the small oil mills and packaging centers that cannot afford the cost of external laboratory analysis.

The possibility of simple, quick and in-situ analysis for the food quality control (often implemented 76 in the form of portable electronic systems) has been widely researched in the last years (Oujji et al., 77 2014; Grossi, Di Lecce, Arru, Gallina Toschi, & Riccò, 2015; Sture, Ruud Oye, Skavhaug, & 78 Mathiassen, 2016; Arsalane et al., 2018). Concerning the olive oil free acidity determination, 79 80 different innovative techniques have been presented in literature (Valli et al., 2016). One approach is based on Near-Infrared (NIR) spectroscopy, allowing to estimate acidity by optical spectroscopy 81 analysis of the transmission spectra in the wavenumber range 4541 to 11726 cm<sup>-1</sup> (Armenta, 82 Garrigues, & de la Guardia, 2007). This solution has the advantage that the measurement can be 83 84 carried out on the oil sample without any reagent, but it needs expensive instrumentation (optical spectrophotometer) and requires frequent re-calibration (different calibrations must be carried out 85 86 depending on olives varieties or geographical origin etc.). A pH-metric procedure to measure oil acidity has been also proposed (Tur'yan, Berezin, Kuselman, & Shenhar, 1996): the oil sample is 87 88 extracted with a chemical solvent (0.20 M triethanolamine in a 1:1 solution of water and isopropanol) and the pH measured. This technique is reported to be accurate, but the pH probe 89 90 needs to be used with care and frequent cleaning and re-calibration are needed. Moreover, the used 91 chemicals add costs for disposal after use.

92 The aim of this work is to present a portable electronic system to measure the free acidity in olive 93 oil samples: the instrument is battery-operated and can be used for quick "in-situ" measurements in the oil production or bottling site. The working principle is based on the estimation of the oil FA 94 from the electrical conductance measured by Electrical Impedance Spectroscopy (EIS) of the oil 95 emulsion with a hydroalcoholic solution. EIS is used in a wide range of applications (Grossi, & 96 Riccò, 2017a), such as to estimate the ripening degree of fruits (Harker, & Maindonald, 1994), to 97 characterize plant tissues (Lin, Chen, & Chen, 2012; Ben Hamed, Zorrig, & Hichem Hamzaoui, 98 99 2016), to characterize and to detect the freezing end point of ice cream mixes (Grossi, Lanzoni, 100 Lazzarini, & Riccò, 2012a), to estimate the water content in extra virgin olive oil (Ragni et al., 101 2013), to investigate the corrosion of metal surfaces exposed to acid electrolytes (Ansari, Quraishi, & Singh, 2015; Hussin, Rahim, Nasir, Ibrahim, & Brosse, 2016), to analyse human body 102 103 composition (Khalil, Mohktar, & Ibrahim, 2014; Bera, 2014), to detect the end-point in a titration assay (Grossi, & Riccò, 2017b). 104

The system described in this paper represents a considerable improvement over previous studies (Grossi, Di Lecce, Gallina Toschi, & Riccò, 2014a-b) since it has been completely re-designed with focus on low-cost electronics and small dimensions. To make the system completely portable and reliable for on-site measurements, a temperature sensor has been included to compensate for conductance variations linked to changes in environmental temperature. Moreover, the presented instrument has been in-house validated evaluating its performances in terms of correlation between the concentration of the analyte and the instrument response, limit of detection (LOD) and quantification (LOQ), precision and accuracy. The validation step, in fact, is one of the measures universally recognized as a necessary part of a comprehensive system of quality assurance in analytical chemistry and is an essential component of the measures that a laboratory should implement to allow it to produce reliable analytical data (Thompson, Ellison, & Wood, 2002).

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#### 117 **2. Material and methods**

#### 118 **2.1 Reagents and chemicals**

119 Diethyl ether (ACS reagent, purity  $\geq 99.8\%$ ), ethanol (ACS reagent, purity  $\geq 96\%$ ) and 120 phenolphtalein solution (indicator, 1% in ethanol) were supplied from Sigma-Aldrich, Inc. (St. 121 Louis, MO, USA). Sodium hydroxide 0.1 mol/L (N/10) was purchased from Carlo Erba Reagents 122 S.r.l. (Milan, Italy).

Distilled water was produced by Elix Essential system (Millipore, Molsheim, France). Oleic acid (CAS Number 112-80-1, assay 90%) used as standard for building the calibration curve was supplied from Sigma-Aldrich, Inc. (St. Louis, MO, USA).

#### 126 **2.2 Samples**

127 In order to calibrate the system, 5 samples of refined sunflower oil (with free acidity and peroxide

- value under the limits for vegetables oils according to CODEX STAN 210-1999) added with oleic
- acid were prepared. To cover the range of free acidity of the different quality grades (EVOO, VOO,
- LOO) different concentrations of oleic acid (from 0% to 3.75%) were added (Table 1).
- 131 The in-house validation study was carried out on a set of 30 olive oil (numerically coded from 1 to
- 132 30) with a different quality grade: 20 EVOOs, 7 VOOs and 3 LOOs. An aliquot of 50 mL for each
- sample was stored in PET dark bottles at 11-12  $^{\circ}$ C before the analysis.

# 134 **2.3 Determination of free acidity**

# 135 2.3.1 Official method

136 The determination of FA by titration was applied according to the EU Reg. 2016/1227. An aliquot

- 137 of the oil sample (the weight depends on the presumed free acidity) was dissolved in 100 mL of a
- mixture of diethyl ether and ethanol (1:1 v/v, previously neutralized) and the free fatty acids present
- 139 were neutralized using sodium hydroxide solution (0.1 M). Phenolphthalein was used as indicator.

#### 140 **2.3.2 Portable system**

The working principle of the portable system (Figure 1) is based on the measure of the electrical conductance of an emulsion between a hydroalcoholic solution and the oil sample. In presence of the hydroalcoholic solution, the free fatty acid molecules RCOOH, where R is the hydrocarbon chain, dissociate in the ionic compounds  $H_3O^+$  and  $RCO_2^-$  that contribute to the increase of the emulsion electrical conductance. Consequently, the higher the free fatty acid molecules concentration, the higher ions concentration and the higher the electrical conductance.

A 50 mL round bottom polypropylene tube (Falcon) modified to feature a couple of cap-shaped stainless-steel electrodes (6 mm in diameter, spaced by 12 mm one from the other) required for the electrical characterization is used as sensor to realize the measurement.

The first step to realize the measurement is a "reagent test" to check the electrical properties 150 (electrical conductance) of the hydro-alcoholic solution in order to avoid an overestimation of the 151 152 oil free acidity linked to the conductance of the solution. For this purpose, the tube (sensor) is filled with 9 mL of ethanol and 6 mL of distilled water and its conductance is measured. If the reagent 153 154 passes the test and is suitable for the measure (conductance not higher than 0.6 µS), 1 mL of the oil sample to be tested is added and the obtained mixture is manually shaken vigorously for about 20 155 156 seconds to create an emulsion, whose conductance is measured to estimate the oil acidity. A video showing the procedure to realize the analysis is available in the online version of the paper. 157

The instrument is also equipped with a temperature sensor to measure the room environmental temperature during the analysis. The emulsion electrical conductance and the temperature are measured and these values are used to calculate the emulsion conductance at the calibration temperature (23.5 °C). Then the free acidity of the sample is estimated using the calibration function stored inside the microcontroller non-volatile memory.

#### 163 **2.4 In-house method validation**

164 The following parameters were evaluated in order to ensure the method quality: limit of detection165 (LOD) and quantification (LOQ), intra-day and inter-day precision and accuracy.

Before the application of the portable system on olive oil samples and in order to verify the correlation between the concentration of the analyte and the response of the instrument, a calibration using samples of refined sunflower oil added with oleic acid previously described (see paragraph 2.2) was carried out. All the samples were analyzed in triplicate by both the official method to determine the reference value of FA, and by the portable system to measure the conductance value. Then, the values were plotted in order to study the correlation between the electrical conductance and the free acidity of the samples.

The limit of detection (LOD) and quantification (LOQ) were evaluated considering the linear relation between the conductance measured by the portable system and the square root of FA. The following formulas were applied:

$$176 \quad LOD = 3\sigma/m \tag{1}$$

$$177 \quad LOQ = 10\sigma/m \tag{2}$$

where  $\sigma$  is the standard deviation of the FA square root measured for the refined sunflower oil without addition of oleic acid (sample A) and *m* is the slope of the curve.

To evaluate the inter-day precision of the method, two virgin olive oil samples for each type of quality grade (samples EVOO\_2 and EVOO\_19 for EVOOs, samples VOO\_1 and VOO\_7 for VOOs and samples LOO\_1 and LOO\_3 for LOOs) were analyzed in triplicate for three different days and values were statistically evaluated by the Student's Test (p < 0.05).

Accuracy and intra-day precision were studied analyzing the set of 30 olive oils with a different quality grade. Precision were expressed in terms of RSD (Relative Standard Deviation) measured for the free acidity values obtained with the portable system. For the accuracy, all the samples were also analyzed following the official method (acid-base titration) and the comparison between the values obtained by the two methods was performed (two-tailed paired t-test with p < 0.05).

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### 190 **3. Results and discussion**

## **3.1 Design of the portable system**

A portable battery-operated electronic system for in-situ measurements of olive oil free acidity has
been built to allow olive oil quality assurance directly in the oil mills or packaging centers.

194 The dimensions of the instrument are  $11 \times 15 \times 5$  cm and some pictures are presented in Figure 2a.

The instrument can be powered by USB port or using batteries (3 AAA alkaline batteries 1.5 V). The system is composed of an electronic board designed ad-hoc that performs all the operations to measure the free acidity, a 2 rows 16 columns LCD screen to output the measure results, four buttons for user interaction and the sensor previously described (see paragraph 2.3.2). The electronic board is based on the microcontroller STM32L152RCT6A and its schematic is presented

- in Figure 2b.
- To determine the FA of an oil sample the electrical conductance of the emulsion is measured by Electrical Impedance Spectroscopy (EIS). In the proposed approach the sample under investigation is stimulated with a 1 V 200 Hz sine-wave voltage signal  $V_{in}(t)$  generated by the built-in 12-bits DAC inside the microcontroller and applied to the sensor electrodes.
- The current drawn by the sensor is converted to a sine-wave voltage  $V_{out}(t)$  by means of a currentto-voltage converter. Given

$$207 V_{in}(t) = V_{M,in} \times sen(2\pi f t) (3)$$

208 it is

209 
$$V_{out}(t) = V_{M,out} \times sen(2\pi f t + \varphi)$$
(4)

- where  $V_{M,in}$  and  $V_{M,out}$  are the amplitude of the corresponding signals,  $\varphi$  is the phase difference between the current through the sample and  $V_{in}(t)$ , while *f* is the frequency of the test signal (200 Hz).
- Both  $V_{in}(t)$  and  $V_{out}(t)$  are acquired by the built-in 12-bits ADC inside the microcontroller using a sampling frequency of 50 kHz and the sine-wave parameters are calculated using the algorithm previously presented by Grossi et al. (Grossi, Lanzoni, Lazzarini, & Riccò, 2012b).
- The emulsion in direct contact with the electrodes can be modelled as the parallel of an electrical conductance (accounting for the conductance of the emulsion) and a capacitance (accounting for the emulsion dielectric properties): while the emulsion conductance (that dominates at low frequency) is affected by the sample acidity due to the variation of the ions concentration, the dielectric properties are almost independent.
- 221 The electrical conductance  $(G_m)$  is thus calculated as:

222 
$$G_m = \frac{1}{R_F} \times \frac{V_{M,out}}{V_{M,in}} \times cos(\varphi)$$
(5)

- where  $R_F$  is the feedback resistance (470 k $\Omega$ ) of the current-to-voltage converter.
- However, the relation between  $G_m$  and the free acidity is non-linear and it can be modelled with the function:

$$226 \qquad G_m = \alpha + \beta \, \times \, \sqrt{FA} \tag{6}$$

where  $\alpha$ ,  $\beta$  are empirical parameters that must be determined by a suitable calibration procedure and are also function of the calibration temperature.

Then, the sample free acidity can be estimated from the measured electrical conductance of the emulsion with the following formula:

$$FA = \left( \left( G_m - \alpha \right) / \beta \right)^2 \tag{7}$$

Since the system must be operated "in the field" and the environmental temperature is not a parameter under control, the environmental temperature (*T*) is measured by means of a MCP9700A analog temperature sensor integrated on the electronic board: the output voltage (that has a sensitivity of 10 mV/°C) is acquired by a channel of the microcontroller ADC and converted to the temperature value.

The system works as follows: the emulsion electrical conductance registered at a certain temperature ( $G_{m,T}$ ) and the temperature (T) are measured and these values are used to calculate the emulsion conductance at the calibration temperature ( $G_{m,Tcalib}$ ). Then the free acidity of the oil sample is estimated from the calculated value of  $G_{m,Tcalib}$  using the calibration function stored inside the microcontroller non-volatile memory. A more detailed description of the system by an electronical and mathematical point of view is presented in Appendix A, while a discussion on the influence of the environmental temperature on the measured conductance and the method used for compensation is presented in Appendix B.

## 245 **3.2 In-house method validation**

Before being used with real olive oil samples, the portable instrument has been calibrated. For this 246 purpose, all the samples of refined sunflower oil added with oleic acid (samples A-E) were analyzed 247 by both the official method to determine the reference value of free acidity and by the portable 248 system to measure the conductance value. The data obtained are shown in Table 1. Then the values 249 250 were plotted (Figure 3): in all cases the electrical conductance measured for the samples increases with its free acidity with a non-linear relation, confirming what previously presented (Grossi, Di 251 252 Lecce, Gallina Toschi, & Riccò, 2014b). Starting from these data and applying a nonlinear regression, the equation of the calibration curve was defined as follow: 253

254 
$$FA = \left(\frac{G_{m,23.5^{\circ}C} + 0.0678}{2.7877}\right)^2$$
 (8)

where  $G_{m,23.5^{\circ}C}$  is the emulsion electrical conductance at the calibration temperature of 23.5 °C.

It allows to obtain, directly on the display of the portable system, the free acidity value of the testedsample estimated starting from its electrical properties.

Subsequently, some parameters for the in-house validation of the instrument were evaluated inorder to check its performances and applicability on real olive oil samples.

LOD and LOQ were measured considering the FA of the refined sunflower oil and the slope of the linear correlation between the free acidity square root and the conductance. The values obtained were 0.02% and 0.06% of oleic acid for LOD and LOQ respectively.

Next, a set of 30 olive oil samples with a different quality grade (20 Extra Virgin Olive Oils –
EVOO; 7 Virgin Olive Oils – VOO; 3 Lampante Olive Oils – LOO) were analyzed in triplicate
applying this new analytical approach in order to study its accuracy and intra-day precision
(expressed in terms of RSD).

The portable instrument showed good precision results since all the obtained values were under the 267 15%. All the samples were also analyzed following the official method (acid-base titration) and the 268 comparison between the values obtained by the two methods was performed to measure the 269 accuracy of the system. The differences between the two series of results (official method vs 270 271 portable system) were evaluated by using the two-tailed paired t-test (p < 0.05). The two approaches did not give statistically differences for the mean values (t<t (critical value)). Moreover, the 272 regression between the two series of values provided a coefficient  $R^2_{adi}$  of 0.97 (Figure 4) in 273 agreement with data previously presented by Grossi et al. (Grossi, Di Lecce, Gallina Toschi, & 274

Riccò, 2014). Considering the commercial categories of the samples analyzed, all of them, with the only exception of the sample VOO\_7, were classified in the same way by both approaches. Finally, considering the inter-day precision of the instrument, no significative differences (Student's Test, p < 0.05) were found among the results obtained for each analyzed sample in the three different days.

280

## 281 **4.** Conclusions

The design and in-house validation of a portable battery-operated electronic system suitable for in-282 283 situ measurements of olive oil free acidity has been presented. The system is built with low cost 284 electronics and embeds a temperature sensor to compensate variations of the measured electrical 285 parameters with the environmental temperature, thus making it suitable for on-site free acidity measurements outside a laboratory. Its working principle is based on the estimation of the olive oil 286 287 FA from the measure of the conductance of an emulsion between a hydro-alcoholic solution and the sample to be tested. When the free fatty acids present in the sample comes in contact with the 288 289 hydro-alcoholic solution, a dissociation occurs, leading to the formation of ions that produce an 290 increase of the measured electrical conductance.

The system has been calibrated and in-house validated. The data obtained showed good performances of the instrument in terms of LOD and LOQ, intra-day and inter-day precision. Moreover, it showed a good correlation ( $R^2_{adj} = 0.97$ ) with the FA evaluated applying the official method demonstrating a satisfactory accuracy.

The procedure to realize the analysis is very quick and easy. This make the system suitable also for 295 people without specific training. The application of this analytical device is addressed, in particular, 296 to the estimation of free acidity of just produced virgin olive oils in oil mills. A next study will be 297 focused on the evaluation of the performance of this portable system when applied to bottles stored 298 299 for several months in dark/light conditions simulating possible different commercial conservation/exposure on shelves of groceries. This study will give indications regarding the 300 maximum oxidation state of olive oil sample tolerable to use this portable system as accurate and 301 302 reliable.

303

#### 304 Funding

This work was supported by the Horizon 2020 European Research project OLEUM "*Advanced* solutions for assuring the authenticity and quality of olive oil at a global scale", which has received

funding from the European Commission within the Horizon 2020 Programme (2014–2020), grant

308	agreement no. 635690. The information expressed in this article reflects the authors' views; the
309	European Commission is not liable for the information contained herein.
310	
311	Declaration of interest
312	None.
313	
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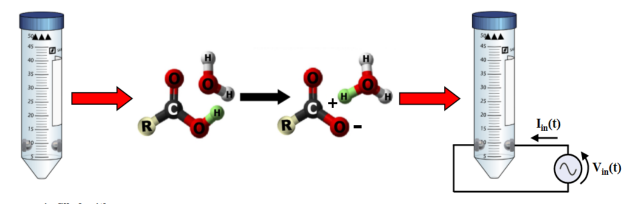
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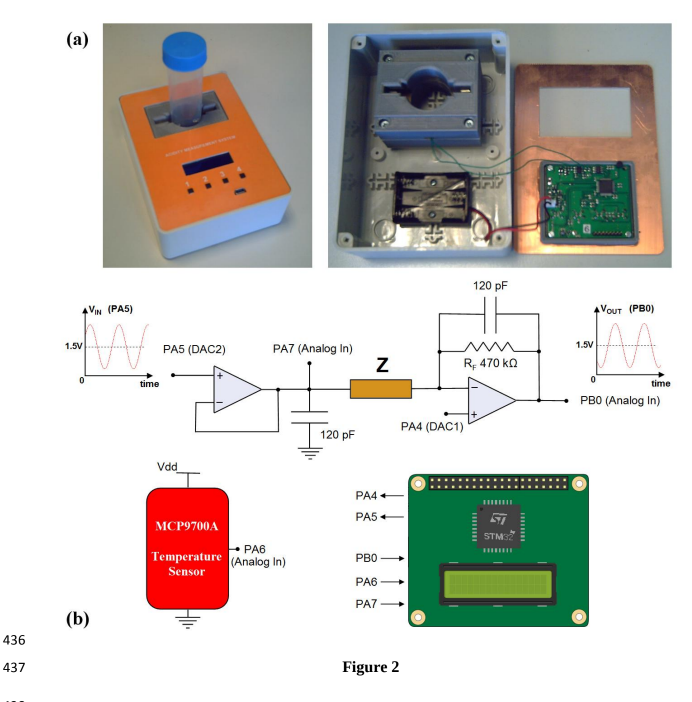


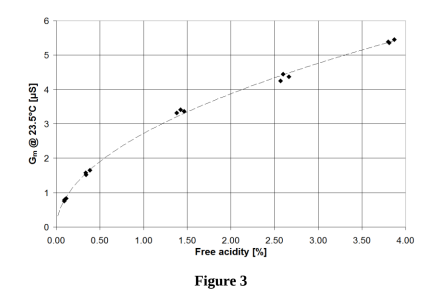
The sensor is filled with the hydroalcoholic solution (40% distilled water / 60% ethanol) and, after the addition of 1 ml of the oil sample, is manually shaken to create the emulsion

The free fatty acid moleculesA sinddissociate in presence of the200 Hhydroalcoholic solution and the H3O+sensions contribute to the increase of theelectricatemulsion electrical conductancethe oil s

A sine-wave signal (1V amplitude, 200 Hz frequency) is applied to the sensor electrodes. The emulsion electrical conductance is measured and the oil sample free acidity is calculated

Figure 1





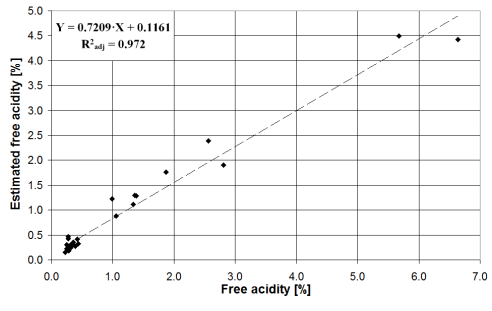


Figure 4

## 444 **Figure captions**

- Figure 1. Scheme of the working principle of the proposed electronic system for the determination
  of free acidity in virgin olive oil (Grossi, & Riccò, 2017a).
- 447 Figure 2. Pictures of the outside and inside of the electronic system and the sensor for olive oil free
- 448 acidity analysis (a); schematic of the system hardware and of its electronical functioning (b).
- 449 Figure 3. Measured electrical conductance  $(G_m)$  at 23.5°C vs free acidity for the sunflower oil
- 450 calibration set.
- 451 **Figure 4.** Scatter plot of the estimated free acidity vs the free acidity measured by titration for a set
- 452 of 30 olive oil samples.

# 453 **Table 1**

are mean of three replicates.

454 Codes and description of the samples used for the calibration of the system, data of free acidity 455 determined by the official method and conductance measured with the portable system. The values

Sample Free acidity Conductance Sample description code (% oleic acid) (**µS**)  $0.79\pm0.03$ А Refined sunflower oil (without addition of oleic acid)  $0.10\pm0.01$ В Refined sunflower oil + 0.25% oleic acid  $0.36\pm0.03$  $1.58\pm0.06$ С Refined sunflower oil + 1.25% oleic acid  $1.42\pm0.04$  $3.36\pm0.04$ D Refined sunflower oil + 2.50% oleic acid  $2.61\pm0.05$  $4.35\pm0.10$ E Refined sunflower oil + 3.75% oleic acid  $3.83\pm0.04$  $5.40\pm0.05$ 

457

Sample code	Free acidity	Free acidity
Sample code	(portable system)	(official method)
EVOO_1	0.18	0.25
EVOO_2	0.31	0.34
EVOO_3	0.25	0.28
EVOO_4	0.43	0.27
EVOO_5	0.30	0.25
EVOO_6	0.22	0.25
EVOO_7	0.18	0.25
EVOO_8	0.29	0.37
EVOO_9	0.29	0.34
EVOO_10	0.47	0.27
EVOO_11	0.32	0.33
EVOO_12	0.41	0.42
EVOO_13	0.24	0.28
EVOO_14	0.27	0.28
EVOO_15	0.32	0.44
EVOO_16	0.35	0.35
EVOO_17	0.18	0.28
EVOO_18	0.15	0.22
EVOO_19	0.27	0.39
EVOO_20	0.24	0.31
VOO_1	1.30	1.36
VOO_2	1.29	1.38
VOO_3	0.88	1.05
VOO_4	1.11	1.33
VOO_5	1.22	0.99
VOO_6	1.76	1.87
VOO_7	1.90	2.81
LOO_1	2.39	2.56
LOO_2	4.42	6.64
LOO_3	4.49	5.67

Table 2. Values of free acidity for all samples measured by the portable system and the official
method. Results are expressed as % of oleic acid.

#### 461 Appendix A

- According to the working principle of the portable system, a 50 mL polypropylene tube (Falcon) vial modified to feature a couple of stainless-steel electrodes to measure the emulsion conductance (hereafter the sensor) is filled with 15 mL of hydro-alcoholic solution (40% distilled water/60% ethanol), then 1 mL of the olive oil sample is added and all is stirred to create an emulsion.
- In presence of the hydroalcoholic solution, the free fatty acid molecule RCOOH, where R is the hydrocarbon chain, dissociates in the ionic compounds  $H_3O^+$  and  $RCO_2^-$  that contribute to the increase of the emulsion electrical conductance. In the end, the higher the free fatty acid molecules concentration, the higher ions concentration and the higher the electrical conductance.
- 470 The emulsion electrical conductance is measured by Electrical Impedance Spectroscopy (EIS).
- 471 In the proposed approach the sample under investigation is stimulated with a sine-wave voltage 472 signal  $V_{in}(t)$ :

473 
$$V_{in}(t) = V_{M,in} \times \sin(2\pi f t)$$
(A1)

474 and the current  $I_{in}(t)$  through the sample is measured:

475 
$$I_{in}(t) = I_{M,in} \times sin \left(2\pi f t + \varphi\right)$$
(A2)

- 476 where  $V_{M,in}$  and  $I_{M,in}$  are the amplitudes of the corresponding signals, *f* is the frequency of the test 477 signal and  $\varphi$  is the phase difference between  $I_{in}(t)$  and  $V_{in}(t)$ .
- 478 The sample electrical admittance is then expressed as:

479 
$$Y = \frac{I_{in}(j2\pi f)}{V_{in}(j2\pi f)} = \frac{I_{M,in}}{V_{M,in}} \times (\cos\phi + j \times \sin\phi) = Re(Y) + j \times Im(Y)$$
(A3)

- The emulsion in direct contact with the electrodes can be modelled as the parallel of an electrical conductance (accounting for the conductance of the emulsion) and a capacitance (accounting for the emulsion dielectric properties): while the emulsion conductance (that dominates at low frequency) is affected by the sample acidity due to the variation of the ions concentration, the dielectric properties are almost independent. Thus, the emulsion electrical conductance  $G_m$  can be estimated with the real component of the emulsion admittance  $Re(Y) = |Y| \cdot cos(\phi)$  where |Y| is the admittance modulus.
- In Figure A1 (a) the admittance modulus (measured with the commercial LCR meter Agilent E4980A) is plotted vs. the frequency of the applied test signal for different samples featuring different acidity in the frequency range 20 Hz – 2 MHz. As can be seen, |Y| increases with sample acidity for test signals with a frequency lower than 20 kHz while it is almost independent on sample acidity for higher frequencies. This is the reason why the designed electronic system measures  $G_m$ with a single frequency measurement at 200 Hz.
- 493 The relation between  $G_m$  and the free acidity is non-linear and it can be modelled with the function:

494 
$$G_m = \alpha + \beta \times \sqrt{FA} + \gamma \times \sqrt[4]{FA^3}$$
(A4)

495 where  $\alpha$ ,  $\beta$  and  $\gamma$  are empirical parameters that must be determined by a suitable calibration 496 procedure and are also function of the calibration temperature.

In Figure A1 (b) the qualitative plot of  $G_m$  vs FA is shown: as can be seen, the non-linear function results in better accuracy for the estimated free acidity for lower acidity levels. This has been taken in account by using a suitable amount of sample to create the emulsion to obtain good accuracy in the acidity range of interest.

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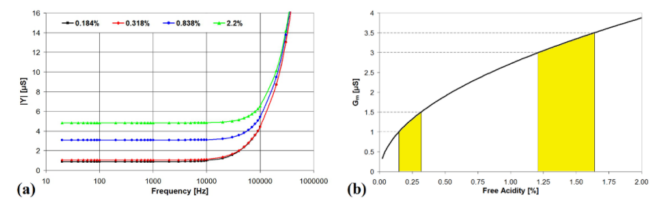


Figure A1 (a) admittance modulus plotted *vs* frequency for olive oil samples featuring different free
acidity; (b) qualitative plot of the electrical conductance as function of sample acidity.

505

502

However, eq. A4 needs a computation intensive iterative algorithm to extract the estimated acidity from the measured value of  $G_m$  as well as manual input of starting point to avoid failing in algorithm convergence. Thus, a simpler model has been chosen (obtained by neglecting the molar conductivity dependence on the H<sub>3</sub>O<sup>+</sup> ions concentration) that is much more suitable to be implemented in a low-cost microcontroller and can be described by the following function:

511 
$$G_m = \alpha + \beta \times \sqrt{FA}$$
 (A5)

The two models of eq. A4 and A5 have been tested with the data set of Grossi et al. (Grossi et al., 2014b) and the results have shown how the accuracy in estimating free acidity is only marginal higher for model A4 than model A5.

515 The sample free acidity can thus be estimated from the measured electrical conductance of the 516 emulsion with the following formula:

517 
$$FA = \left(\frac{G_m \cdot \alpha}{\beta}\right)^2$$
 (A6)

518

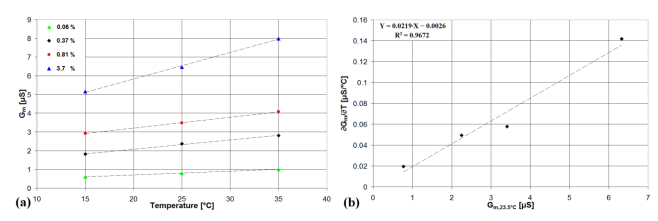
519 Appendix B

The oil sample free acidity can be estimated by measuring the emulsion electrical conductance at the temperature of calibration and then calculating the free acidity using equation A6. However, since the system must be operated "in the field" and the environmental temperature is not a parameter under control, there is the need to investigate how the emulsion conductance varies with the temperature so that the oil free acidity can be estimated by the measure of the emulsion conductance and the temperature.

Four different olive oil samples featuring different free acidity values (0.06% sample A, 0.37%
sample B, 0.81% sample C and 3.7% sample D) were tested inside a Binder APT KB 53 thermal
incubator for different temperatures between 15°C and 35°C.

In Figure B1 (a) the measured emulsion conductance is plotted vs the incubation temperature for each sample. In all cases the  $G_m$  is a linear function of the temperature with determination coefficients  $\mathbb{R}^2 > 0.99$ . The calculated linear regression lines allow to determine the conductance variation with temperature (i.e.  $\partial G_m / \partial T$ ) for all samples: 0.0196 for sample A, 0.0495 for sample B, 0.0579 for sample C and 0.1417 for sample D.  $\partial G_m / \partial T$  is thus found to increase with the sample free acidity. Since the sample free acidity is also a function of the emulsion electrical conductance, the relation between  $\partial G_m / \partial T$  and  $G_{m,23.5^{\circ}C}$  has been plotted in Figure B1 (b).





**Figure B1** (a) measured emulsion conductance vs the incubation temperature for olive oil samples featuring different free acidity; (b)  $\partial G_m / \partial T$  plotted vs the electrical conductance at 23.5 °C.

540

537

541 As can be seen, a linear relation gives a good approximation ( $\mathbb{R}^2 = 0.9672$ ) of the function between 542  $\partial G_m / \partial T$  and  $G_{m,23.5^\circ C}$  thus:

543 
$$\frac{\partial G_m}{\partial T} = 0.0219 \times G_{m,23.5^{\circ}C} - 0.0026$$
 (B1)

where  $\partial G_m / \partial T$  and  $G_{m,23.5^{\circ}C}$  are expressed as  $\mu S / \circ C$  and  $\mu S$  respectively. The emulsion electrical conductance at the environmental temperature *T* can thus be expressed as:

546 
$$G_{m,T} = G_{m,23.5^{\circ}C} + \frac{\partial G_m}{\partial T} \times (T-23.5)$$
(B2)

547 and

548 
$$G_{m,T} = G_{m,23.5^{\circ}C} + (0.0219 \times G_{m,23.5^{\circ}C} - 0.0026) \times (T-23.5)$$
 (B3)

The emulsion electrical conductance at  $T_{calib} = 23.5^{\circ}C$  can thus be estimated from the electrical conductance at temperature *T* and the measured value of *T* using the following formula:

551 
$$G_{m,23.5^{\circ}C} = \frac{G_{m,T} + 0.0026 \times (T-23.5)}{1 + 0.0219 \times (T-23.5)}$$
 (B4)

552 Thus, by measuring  $G_{m,T}$  and T, the value of  $G_{m,23.5^{\circ}C}$  can be calculated using equation B4 and, from

this value, the sample acidity can be estimated using equation A6.

Video Click here to download Video: acidity\_measure\_system\_english.wmv