Evaluation of the performance of Frying Oils using an ultrasonic technique

By D. Izbaim*, B. Faiz†, A. Moudden‡, N. Taifi§ and I. Aboudaoud

* Laboratory of Metrology and Information Treatment, Faculty of Sciences, University Ibn Zohr, Agadir, Morocco
† Superior School of Technology, University Ibn Zohr, Agadir, Morocco.
§ Faculty of Sciences, University Chouaib Doukkali, El Jadida, Morocco.
(*corresponding author: dizbaim@gmail.com)

1. INTRODUCTION

Frying is one of oldest and simplest methods of cooking food, due to its flavor and ease of preparation. From the consumer standpoint, fried food palatability is related to unique sensory characteristics. However, the frying process involves physical, chemical and sensory changes in the food which can promote the formation of a variety of decomposition compounds with adverse nutritional implications and risks of being hazardous to human health (Marquez-Ruiz and Dobarganes, 1996). Complex changes occur during deep-fat frying which lead to thermal and oxidative decomposition. The physical changes are mainly increased viscosity, foaming and color changes and decreased smoke-point. The main chemical changes are increased free fatty acids and polar components as well as decreased levels of unsaturation, flavor quality and nutritive value (Warner, 1998). As a consequence, in many countries, the polar compound measurement is considered to be the most important test for the quality degradation of oil and a maximum level of polar compounds has been set at 25%, while in some countries, a polar compound cut-off point has been established between 20% and 27% (Firestone, 1996). Some other countries also use free fatty acids. An eventual decline in food quality is a result of the increase in polar fraction (Pokorny, 1989). Weiss (1983) showed that it is mainly free fatty acids which are produced in the frying operation of which contribute to the smoke haze and therefore has a substantial effect on its smoke point. Orthoefer et al. (1996) has demonstrated that the smoke point of the frying fat affects oil absorption by the fried food.

There are many types of oils available for frying. The choice is difficult, with factors such as stability, price and nutritive value. The thermal stability of the frying oil is related to the level of hydrogenation (Arroyo et al., 1995) and the oxidative stability of the frying oil will increase with an increased degree of hydrogenation. However, these oils are undesirable from a nutrition and health standpoint (Cuesta et al., 1988). So, hydrogenated oil becomes a major concern in frying applications. Recently, partially hydrogenated oils have been increasingly adopted as an alternative to hydrogenated frying oil.

Several methods exist for controlling and assessing oil quality during deep frying. Chemical analyses provide reliable results but are time-consuming, costly, and usually require analytical expertise like...
the use of FTIR spectroscopy (Innawong et al., 2004) or chromametric methods (Xin Qing, 2003). In this respect ultrasonic techniques are fast, non-invasive and inexpensive. Many studies have been conducted to assess the composition and structure of different types of food products using ultrasound (McClements, 1997; Mulet et al., 1999). Therefore, velocity measurements can be used for oil composition and adulteration assessment (McClements and Povey, 1987; Raghupathi et al., 1980).

The objective of this study was the quantitative evaluation of quality parameters of USBO and PHSBO under thermo-oxidation using an ultrasonic technique.

2. MATERIALS AND METHODS

USBO and PHSBO were obtained from Cristal S.A. (Casablanca, Morocco). The oils were heated in a domestic fryer at 180 °C for 8 h/day over 4 days, for a total of 32 h. Samples of 150 mL were periodically removed and kept at −18 °C for further analyses.

2.1. Ultrasonic Measurements

The experimental setup used for the experiments consisted of an ultrasonic transducer (5 MHz, 0.5 in. crystal diameter, A309S-SU Model, Panametrics, Olympus), attached to a cubic container (50 x 50 x 50 mm) where the oil samples were placed. The container was introduced into a temperature-controlled bath to maintain the sample temperature and the oil was moderately stirred to prevent the formation of bubbles. The ultrasonic measurements were carried out while the oil sample was cooled from 50 to 30 °C. The transducer was linked to a pulser-receiver (Sofranel Model 5073PR, Sofranel Instruments) which sent the electrical signal to a digital storage oscilloscope (LeCroy 9310M, LeCroy Cor). For the ultrasonic transducer was linked to a pulser-receiver (Sofranel Instruments) which sent the electrical signal to a digital storage oscilloscope (LeCroy 9310M, LeCroy Cor). For the ultrasonic velocity measurement, eight signal acquisitions (LeCroy 9310M, LeCroy Cor). For the ultrasonic transducer was linked to a pulser-receiver (Sofranel Instruments) which sent the electrical signal to a digital storage oscilloscope (LeCroy 9310M, LeCroy Cor). For the ultrasonic velocity measurement, eight signal acquisitions were taken and averaged. It can be written as:

\[
C_{\text{sample}} = \frac{2\omega L}{\phi_{A4} - \phi_{A2}}
\]

\[
\text{with } \omega = 2\pi v
\]

To obtain the phase \( \phi \) experimentally, the FFT of signals A2 and A4 were calculated (Bakkali et al., 2001).

The attenuation coefficient (\( \alpha \)) was computed by fitting the experimental data to equation.

\[
\ln A = \ln A_0 - \alpha d
\]

Where \( d \) is the distance travelled by the wave, \( A_0 \) is the initial amplitude of the signal measured as the peak-to-peak voltage and A is the amplitude of the signal at distance \( d \). Four ultrasonic echoes were considered to compute attenuation.

2.2. Chemical Analyses

Fatty acids

The FA profile analysis was performed by deriving their corresponding methyl esters (Hartman and Lago, 1973) prior to the analysis by GC. Oil samples (50 mL) were diluted in the hexane to obtain a solution to be analyzed (about approximately 0.1%). A VARIAN 3800 chromatograph on a CP Select CB (VARIAN) capillary column (50 m x 0.25 mm i.d., film thickness 0.25 µm) was used under the following temperature program: 185 °C (40 min), 15 °C/min to 250 °C (10 min). Sample injection was 1 µL (split ratio 1:100) at 250 °C and the flow rate of Helium, used as carrier gas, was 1.2 mL/min. The temperature of both the split injector and flame ionization detector was 250 °C.

Free fatty acids

The FFA contents, as the percentage of oleic acid, were determined using AFNOR NF T 60-204 standard method. Acid value was defined as the amount (mg) of KOH required to neutralize FFA in 1 g of oil sample dissolved in a mixture of diethyl ether and ethanol in the presence of phenolphthalein.

Polar compounds

The contents of total polar compounds were determined following the method proposed by the IUPAC (IUPAC, 1992).

3. RESULTS AND DISCUSSION

3.1. Chemical Characteristics

Tables 1 and 2 show the results for the Fatty Acid (FA) profiles of the USBO and PHSBO carried out on the oil samples after each heating period. The FA composition of an oil has marked effects on its frying performance as well as on its physical and chemical behavior. The FA profile of the frying oils changed as a result of cyclization, polymerization and hydrolytic, oxidative and other chemical reactions promoted by frying conditions (Nawar, 1996). The linoleic acid level in deep-frying oils appears to be an obviously negative factor in

<table>
<thead>
<tr>
<th>Table 1</th>
<th>Soybean oil FA compounds at several frying periods (%)</th>
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<tbody>
<tr>
<td></td>
<td>0h</td>
</tr>
<tr>
<td>C16:0</td>
<td>10.75</td>
</tr>
<tr>
<td>C18:0</td>
<td>3.32</td>
</tr>
<tr>
<td>C18:1</td>
<td>22.30</td>
</tr>
<tr>
<td>C18:2</td>
<td>54.21</td>
</tr>
<tr>
<td>C18:3</td>
<td>5.88</td>
</tr>
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</table>
Partial hydrogenation soybean oil FA compounds at several frying periods (%)

<table>
<thead>
<tr>
<th></th>
<th>0h</th>
<th>8h</th>
<th>16h</th>
<th>24h</th>
<th>32h</th>
</tr>
</thead>
<tbody>
<tr>
<td>C16:0</td>
<td>13.23</td>
<td>13.35</td>
<td>13.28</td>
<td>13.24</td>
<td>13.25</td>
</tr>
<tr>
<td>C18:0</td>
<td>7.40</td>
<td>8.29</td>
<td>9.07</td>
<td>9.46</td>
<td>9.77</td>
</tr>
<tr>
<td>C18:1</td>
<td>56.10</td>
<td>57.15</td>
<td>57.91</td>
<td>59.10</td>
<td>60.08</td>
</tr>
<tr>
<td>C18:2</td>
<td>20.60</td>
<td>19.08</td>
<td>17.75</td>
<td>16.21</td>
<td>14.98</td>
</tr>
<tr>
<td>C18:3</td>
<td>0.56</td>
<td>0.62</td>
<td>0.47</td>
<td>0.36</td>
<td>0.32</td>
</tr>
</tbody>
</table>

Many researchers consider the measurement of Total Polar Components (TPC) to be the most reliable indicator of oil degradation (Fritch, 1981; Gere, 1982). Polar compounds include all oxidized and dimerized triglycerides, FFAs, monoglycerides and diglycerides, sterols, carotenoids, antioxidants, antifoamers, hydrogenation catalyst residues and soaps (Blumenthal 1996). According to Billek et al. (1978) and Paradis and Nawar (1981), polar compounds indicate the degradation of oils and the breakdown of triglycerides. Table 4 shows that TPC in USBO and PHSBO increased significantly during frying. Initially, the TPC contents of USBO and PHSBO were similar. The increase in the rate of TPC formation in the USBO was different from PHSBO. After 32 h of frying, the final TPC levels were 25% in USBO and 21% in PHSBO. Not to exceed the established limit (around 25%) on frying fats and oils, the TPC-based stability is 32 h for frying for USBO and 42 h for PHSBO. This would have occurred in an even shorter time if the oil had been used for frying foods.

These parameters are indicators of the state of oil deterioration and all of them indicated that PHSBO was more stable chemically than USBO and that hydrogenation increased soybean oil stability.

### 3.2. Ultrasonic Measurements

The ultrasonic velocity and attenuation depend on the physico-chemical properties of the oil. Ultrasonic velocity decreases with the temperature in fat (McClements, 1997).

Figures 1a and b show the evolution of ultrasonic velocity with temperature for USBO and PHSBO respectively, fried for 8, 16, 24 and 32 h. As expected, as the temperature increases, the ultrasonic velocity decreases in line. After each frying period, the velocity increases with an almost constant value leading to straight parallel curves. The average velocity temperature coefficient is $-3.78 \text{ m s}^{-1} \text{ °C}^{-1}$ and $-3.57 \text{ m s}^{-1} \text{ °C}^{-1}$ for USBO and PHSBO respectively. Figure 2 shows the evolution of the velocity difference at 40 °C with duration of frying. Comparisons of velocity curves for each frying period of USBO and PHSBO show that velocity difference increases as duration of frying increases. Therefore, ultrasonic velocity is a parameter which allows assessing oil quality and
The ultrasonic velocity depends on time of cooling as it increases as the time of cooling increases (Figure 3). Furthermore, the velocity evolutions are very similar for both types of oils. After each frying period the velocity increases and thus it is possible to distinguish the two oils without using temperature. This shows that time of cooling seems to be an alternative to temperature.

Attenuation is also affected by temperature. As previously reported by Benedito et al. (2002), the ultrasonic attenuation decreases as temperature increases due to changes in the physico-chemical properties of the oil, although it is not so widely reported in the literature.

Relationships between the ultrasonic velocity and both TPC and FFA for USBO and PHSBO at 30 °C are shown in Figures 4 and 5, respectively. These relationships are almost the same for both types of oils, although for the same percentage of TPC and FFA. However, significant polynomial fits were found when relating the velocity and these chemical parameters, widely considered as oil quality indicators. Therefore, the single measurement of velocity permits an estimation of the percentages of polar compounds and FFA independently of the type of oil.
EVALUATION OF THE PERFORMANCE OF FRYING OILS USING AN ULTRASONIC TECHNIQUE

Figure 4
Relationship between ultrasonic velocity and the percentage of polar compounds for USBO and PHSBO at 30 °C.

Figure 5
Relationship between ultrasonic velocity and the percentage of free fatty acids for USBO and PHSBO at 30 °C.

4. CONCLUSIONS

Frying duration, frying temperature and degree of hydrogenation are important factors in influencing oil lifetime and its frying properties. The ultrasonic velocity is related to the free fatty acid level and total polar compounds, showing that USBO performance was slightly lower than that of PHSBO. The ultrasonic measurement results were easily differentiated between USBO and PHSBO. Ultrasonic velocity could be used to evaluate the degradation state of oil and monitoring changes in oil during deep-frying; and ultrasonic techniques might be adapted for real-time quality control purposes of frying operations in the food industry.

REFERENCES


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