Improved method for the determination of triacylglycerols in olive oils by high performance liquid chromatography

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SUMMARY

The analysis of triacylglycerols has great importance as quality control and origin determination (Graciani-Constante 1988a); (Graciani-Constante 1988b); (Aitzetmüller 1993). Thus, the percentage of trilinolein (LLL) was adopted as criterion to detect the presence of seed oils in olive oils (EEC 2568/91). Lately, it was substituted by a new parameter: differences between experimental and theoretical values of triacylglycerols with equivalent carbon number equal to 42 (ΔECN42) (EEC 2472/97). In these official methods, the oil is previously purified by passing through a silica-gel column and then the TAG composition determined by isocratic non-aqueous reversed-phase high-performance liquid chromatography (HPLC) with refractive index (RI) detector using acetone/acetonitrile (1:1) as mobile phase (EEC 2568/91). The TAGs are separated according to their equivalent carbon number (ECN), which is defined by the formula: ECN = CN - 2n, where CN is the acyl carbon number and n the number of double bonds of fatty acids constituting the triacylglycerols (Firestone 1994). The ΔECN42 determination allows the detection of seed oils at detection level of 1-3%, but in the case of hazelnut oil the detection level increases up to 20%, because the TAG composition of the hazelnut oil is similar to that of olive oil.

Recently a new method based in the comparison of mathematical algorithms with those included in a database built with genuine olive oils has been proposed (Cert and Moreda 2000). These algorithms are constituted by parameters obtained from the theoretical TAG composition calculated from the fatty acid composition and the experimental TAG composition determined by HPLC. For a successful application of this method, an accurate determination of the TAGs, in particular LLL and OLLn, is necessary. Nevertheless, the chromatographic resolution achieved with the official analytical method is poor, and therefore, improvements to the TAG analysis by HPLC should be introduced.

Triacylglycerols are effectively separated by HPLC on reversed-phase (RP) columns, containing silica with chemically bonded to octadeyl groups (RP-18) as stationary phase. Using nitriles as mobile phase, separation of TAGs occurs according to the chain length and degree of unsaturation of the fatty acids in the glycerol moiety (Schulte 1981). The

1. INTRODUCTION

The analysis of edible oils by their triacylglycerol (TAG) content has recently assumed a great importance as far as quality control and possible origin determination (Graciani-Constante 1988a); (Graciani-Constante 1988b); (Aitzetmüller 1993). Thus, the percentage of trilinolein (LLL) was adopted as criterion to detect the presence of seed oils in olive oils (EEC 2568/91). Lately, it was substituted by a new parameter: differences between experimental and theoretical values of triacylglycerols with equivalent carbon number equal to 42 (ΔECN42) (EEC 2472/97). In these official methods, the oil is previously purified by passing through a silica-gel column and then the TAG composition determined by isocratic non-aqueous reversed-phase high-performance liquid chromatography (HPLC) with refractive index (RI) detector using acetone/acetonitrile (1:1) as mobile phase (EEC 2568/91). The TAGs are separated according to their equivalent carbon number (ECN), which is defined by the formula: ECN = CN - 2n, where CN is the acyl carbon number and n the number of double bonds of fatty acids constituting the triacylglycerols (Firestone 1994). The ΔECN42 determination allows the detection of seed oils at detection level of 1-3%, but in the case of hazelnut oil the detection level increases up to 20%, because the TAG composition of the hazelnut oil is similar to that of olive oil.

Recently a new method based in the comparison of mathematical algorithms with those included in a database built with genuine olive oils has been proposed (Cert and Moreda 2000). These algorithms are constituted by parameters obtained from the theoretical TAG composition calculated from the fatty acid composition and the experimental TAG composition determined by HPLC. For a successful application of this method, an accurate determination of the TAGs, in particular LLL and OLLn, is necessary. Nevertheless, the chromatographic resolution achieved with the official analytical method is poor, and therefore, improvements to the TAG analysis by HPLC should be introduced.

Triacylglycerols are effectively separated by HPLC on reversed-phase (RP) columns, containing silica with chemically bonded to octadeyl groups (RP-18) as stationary phase. Using nitriles as mobile phase, separation of TAGs occurs according to the chain length and degree of unsaturation of the fatty acids in the glycerol moiety (Schulte 1981). The
nitrile most commonly used is acetonitrile with an organic modifier to improve the solubility of TAG bringing about changes in mobile phase polarity, increasing peak selectivity, retention times and selectivity of pairs of groups of TAG with the same ECN. Although, many organic modifiers have been used, the most commonly employed is acetone in different proportions. In the official methods, acetone/acetonitrile (1:1) is used as mobile phase (IUPAC 1987); (EEC 2472/97). However, these methodologies yield only relatively satisfactory separations for critical pairs LLL/OLLn and OLL/OOLn. Using propionitrile, Fiebig 1985, achieved better separations on long HPLC columns (500 x 4 mm, 5 m particle size) at 27.5 °C, and Ollivier et al. 1999 also obtained good separations using shorter columns (250 x 4 mm, 4 m particle size) at 24 °C and linear flow gradient. Mixtures of acetonitrile with tetrahydrofuran, dichloromethane and chloroform in different proportions (Graciani-Constante and Delgado-Noriega 1987), (Bouteiller and Maurin 1991) have been also assayed.

Most analysis of TAG has been done with the column at ambient temperature. However, changes in column temperature resulted in changes in chromatographic resolution, showing that higher column temperature decreased retention times (Fiebig 1985) and selectivity for the TAG, especially for those critical pairs. Although, lower temperatures results in better separations of the TAG, the election of the temperature analysis represent a compromise to ensure good solubility of the saturated TAG, good selectivity of the critical pairs with the same ECN and shorter run time (Frede 1986).

In this work, several variables affecting to TAG determination in olive oils are changed, in order to achieve a better separation of critical pairs than in the official methods. These changes are size of stationary phase particle, column temperature and nature of mobile phase. The triacylglycerol contributing to each HPLC peak are determined. In addition, the application of solid phase extraction (SPE) to the oil purification (Cert et al. 1996) is proposed.

2. MATERIALS AND METHODS

2.1. Materials

All reagents were of analytical grade, except acetone, acetonitrile and propionitrile, which were of super purity grade from Romil (Cambridge, United Kingdom). Silica gel cartridges of 1 g (6 ml) for solid phase extraction were purchased from Waters (Massachusetts, USA).

Reference triacylglycerols (SSS, PPP, PoPoPo and OOO) were supplied by Sigma-Aldrich (St. Louis, MO, USA).

For the assays, virgin olive oils from different varietal origins were chosen in order to have a wide range of fatty acid composition.

2.2. Oil purification

The oil samples were purified by passing the oil through silica SPE. Silica SPE column was placed in a vacuum elution apparatus and washed under vacuum with 6 ml of hexane. The vacuum was released to prevent the dryness of the column and then a solution of the oil (0.12 g) in 0.5 ml of hexane was charged into the column. The solution was pulled through and then eluted with 10 ml of hexane-diethylether (87:13 v/v) under vacuum. The eluted solvents were homogenised and evaporated to dryness in a rotary evaporator under reduced pressure at room temperature. The residue was dissolved, in 2 ml of acetone for triacylglycerol (TAG) analysis.

2.3. HPLC analysis of triacylglycerols

A 10 µl aliquot of the purified oil solution in acetone (5%) was injected onto the HPLC system using an autosampler Beckman Gold 508 (Beckman-Coulter, Fullerton, CA, U.S.A.). The analyses were done on a Lichrosphere 100 RP-18 (4m) column (25 cm x 4 mm I.D.) using a Beckman Gold 126 pumping unit (Beckman-Coulter, Fullerton, CA, U.S.A.), refractive index detector Perkin Elmer 200 (Perkin Elmer, Norwalk, CT, USA) and Beckman Mistral peltier column thermostat unit (Beckman-Coulter, Fullerton, CA, U.S.A.).

Method A: EU official method for olive oils (EEC 2568/91; EEC 2472/97). The mobile phase was acetone/acetonitrile (1:1) at a flow rate to elute trilinolein at 12 min. (1.1 ml/min approximately). Oven temperature 30 °C.

Method B: the mobile phase was acetone/propionitrile (55:45) at a flow rate to elute the trilinolein at 13 min (1.4 ml/min, approximately). Oven temperature gradient: initial temperature 20°C (30 min) and then increased up to 35°C at 0.5°C/min.

Method C: elution with propionitrile at a flow rate to elute the trilinolein at 15.5 min (0.6 ml/min, approximately). Oven temperature 20°C.

3. RESULTS AND DISCUSSION

Vegetable oils possess a characteristic and more or less unique pattern of TAG that can be used to determine origin and to detect adulteration. The separation of the TAGs according to their ECN is strongly influenced by the mobile phase composition.
and the column temperature. Using the analytical conditions indicated in the official method for TAG determination in olive oils (method A) (EEC 2568/91; EEC 2472/97), the resolution of most TAG was acceptable, but certain critical pairs remained unresolved, such as, LLL/OLLn in the ECN42, OLL/OOLn, PLL/POLn in the ECN44, and OOL/PoOO in the ECN46 groups. Figure 1 shows a HPLC chromatographic profile of a virgin olive oil (“Chamlali” variety) analysed using the three methods. In the profile A, it can be seen that the ECN42 triacylglycerols can be accurately determined, but the critical pair LLL/OLLn is not well separated (Cort and Moreda 2000). In order to improve the resolution, acetonitrile/acetone (55:45) as mobile phase was used and the column maintained at sub-ambient temperature (20 °C) using a Peltier oven to thermostat the column. The results showed good separations in the TAG of ECN42 and 44 but the increase of the retention times because of the low temperatures results in long analysis. To ensure a compromise between the resolution and the run time analysis, the low temperature was maintained until elution of the ECN44 (30 min) and then increased at 0.5 °C/min to reach 35 °C at the end of the analysis (60 minutes) (Figure 1, profile B). The obtained separation of the critical pairs was better than the standard method in the area of the ECN42 and ECN44. Although, this method yield better results, the improvements were not enough to obtain good separations between peaks. Therefore, seeking new improvements in the analysis of TAGs was necessary.

The steps were directed toward the use of new eluting solvents. Several solvents were tested and propionitrile resulted in the most suitable. Although its toxicity is higher than the acetonitrile, the use of a single elution solvent and an autosampler decreases dramatically the risk of vapour inhalation. This method (method C) yielded an HPLC chromatogram of virgin olive oil (“Chamlali” variety) in which the resolution of the critical pairs is almost complete (Figure 1, profile C). Figure 2 shows the different HPLC profiles obtained from an oil with low linoleic content (“Picual variety”), in which it can be seen that the resolution and quantitation of LLL using method C (Figure 2, profile C) is much better than in the other methods; Figure 2, profiles A and B. These results encourage its use and therefore, it was worthwhile to perform a complete study of the assignment of the
peaks and precision of the method. The use of propionitrile instead of acetone/acetonitrile, reduces dramatically the baseline drift due to the volatility of the solvent and, therefore, better quantitations can be obtained.

The first step was towards the assignment of the corresponding peaks. Elution order is determined by calculating equivalent carbon number, ECN, often defined as $CN-2n$. To calculate ECN more accurately, a plot of $\log \alpha$ versus $n$ was depicted for various series ($S - O$, $P - O$, $O - L$, $O - Ln$ and $O - Po$ series) (Figure 3), being $\alpha = \frac{RT_i - RT_s}{RT_{OOO} - RT_s}$ ($RT_i$ is the retention time of the TAG, $RT_s$ the retention time of the solvent and $RT_{OOO}$ the retention time of the triolein). In this plot, it can be seen that the successive replacement of the fatty acid moiety having the same CN results in a change in the $\log \alpha$. Under these chromatographic conditions conditions, ECN approximates: $ECN = CN - (3.15x n_0) - (2.90 X n_{po}) - (2.34 n_{ln})$ where $n_0$, $n_{po}$, $n_{ln}$ are the number of double bonds of oleic, palmitoleic, linoleic and linolenic acids respectively, and the coefficients are calculated from reference triacylglycerols (Firestone 1994).

Using the above equation, it was possible to assign the different peaks in HPLC analysis with propionitrile as mobile phase (Figures 1 and 2, profiles C). It can be seen that the triacylglycerols contributing to the HPLC peaks are different using acetone/acetonitrile and propionitrile elution.

![Figure 3](plot.png)

Figure 3
Plot of the $\log \alpha$ versus number of double bonds for the series $S - O$, $P - O$, $O - L$, $O - Ln$ and $O - Po$ using method C.

| ECN | HPLC peaks | TAGs | Sample 1 | Sample 2 | Sample 3 | Sample 4 | Sample 5 |
|-----|------------|------|----------|----------|----------|----------|----------|
|     |            | Mean (%) | RSD (%) | Mean (%) | RSD (%) | Mean (%) | RSD (%) | Mean (%) | RSD (%) |
| 42  |            | 0.020 | 7.23 | 0.086 | 5.18 | 0.095 | 4.10 | 0.113 | 0.95 | 0.34 | 1.05 |
| 44  |            | 0.085 | 7.44 | 0.24 | 1.78 | 0.26 | 2.25 | 0.35 | 2.02 | 0.50 | 2.83 |
| 46  |            | 0.023 | 15.74 | 0.039 | 5.51 | 0.057 | 5.62 | 0.082 | 4.35 | 0.12 | 6.15 |
| 48  |            | 0.47 | 1.52 | 1.53 | 0.42 | 2.62 | 0.98 | 3.35 | 1.05 | 4.37 | 1.13 |
| 50  |            | 1.07 | 2.01 | 1.54 | 0.46 | 1.61 | 0.71 | 1.72 | 1.07 | 1.77 | 2.40 |
| 11  |            | 0.11 | 12.86 | 0.24 | 4.37 | 0.65 | 1.32 | 1.35 | 0.73 | 2.28 | 1.24 |
| 15  |            | 0.085 | 7.44 | 0.24 | 1.78 | 0.26 | 2.25 | 0.35 | 2.02 | 0.50 | 2.83 |
| 17  |            | 0.023 | 15.74 | 0.039 | 5.51 | 0.057 | 5.62 | 0.082 | 4.35 | 0.12 | 6.15 |
| 21  |            | 0.47 | 1.52 | 1.53 | 0.42 | 2.62 | 0.98 | 3.35 | 1.05 | 4.37 | 1.13 |
| 25  |            | 1.07 | 2.01 | 1.54 | 0.46 | 1.61 | 0.71 | 1.72 | 1.07 | 1.77 | 2.40 |

| Sample 1 | Sample 2 | Sample 3 | Sample 4 | Sample 5 |
|----------|----------|----------|----------|----------|
| Mean (%) | RSD (%)  | Mean (%) | RSD (%)  | Mean (%) | RSD (%)  | Mean (%) | RSD (%)  | Mean (%) | RSD (%)  |
| 0.020 | 7.23 | 0.086 | 5.18 | 0.095 | 4.10 | 0.113 | 0.95 | 0.34 | 1.05 |
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| 1.07 | 2.01 | 1.54 | 0.46 | 1.61 | 0.71 | 1.72 | 1.07 | 1.77 | 2.40 |
| 0.11 | 12.86 | 0.24 | 4.37 | 0.65 | 1.32 | 1.35 | 0.73 | 2.28 | 1.24 |
| 0.42 | 5.11 | 0.49 | 2.89 | 0.55 | 2.01 | 0.85 | 1.83 | 1.09 | 1.96 |
| 6.72 | 0.63 | 8.79 | 0.31 | 11.21 | 0.42 | 13.25 | 0.33 | 15.24 | 0.23 |
| 1.24 | 2.68 | 1.49 | 0.35 | 1.63 | 0.85 | 2.12 | 0.45 | 2.52 | 0.56 |
| 2.70 | 0.65 | 4.05 | 0.70 | 6.02 | 0.65 | 9.86 | 0.53 | 11.53 | 0.31 |
| 0.64 | 4.42 | 0.69 | 3.02 | 0.79 | 1.23 | 1.53 | 0.89 | 1.70 | 1.66 |

Table I
Repeatability data of the determination of virgin olive oil TAGs by HPLC using propionitrile as eluent

* $n = 3$ replicates
* RSD = Relative Standard Deviations of the repeatability
In order to evaluate the repeatability of the determination of the TAGs by HPLC using propionitrile as eluent, analysis of virgin olive oils having different fatty acid compositions was performed 3 times. The results obtained (Table I) indicate that the relative standard deviation of the method is good.

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