Oils rich in alpha linolenic acid: chemical composition of perilla (Perilla frutescens) seed oil

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Abstract – In this paper, the main chemical properties of Perilla seed oil (PO) obtained by mechanical pressing of Perilla seeds are reported. The analysis of fatty acid composition has highlighted a very high amount of (n-3) α-linolenic acid (ALA), more than 60%, higher than other ALA rich oils such as linseed and sacha inchi oils and similar to chia oil. PO has an important sterol (higher than 3000 mg/kg) and vitamin E (approx. 870 mg/kg) content, while biophenols are in quite low concentration. The analyzed sample showed a very low acid and peroxide value and this is the demonstration that, following the proper technological procedures, it is possible to obtain high quality oils even in presence of high α-linolenic acid concentrations.

Keywords: Perilla frutescens / oil composition / omega 3 fatty acids / tocopherols

1 Introduction

Uncommon oils rich in alpha linolenic acid have several interesting properties, as demonstrated during some previous studies (Bondioli, 2006; Bondioli et al., 2006; Bondioli, 2019). Perilla frutescens is an annual aromatic plant belonging to the Lamiaceae family, also known in Japan as egoma or shiso. It is the only species belonging to the genus Perilla and it is widespread in India, Vietnam, Korea, China and Japan.

Both leaves and the oil obtained from the seeds are used for culinary and medical purposes (Yu et al., 1997). Perilla seed oil (PO), on the other hand, can be used as a salad oil or for industrial applications such as paints, dyes and inks.

PO is not an absolute novelty for Italian market, because after the Second World War, in the North of Italy, some facilities for seed crushing and oil refining were installed. At that time, the oil was in use as a drying oil for paints preparation. In fact, some information about its composition can be found in a technical manual distributed until 1988 in Italy by a supplier of oil plants for oil extraction and refining (Agenda Gianazza, 1988). From this source, we can get iodine value (190–208 g J2/100 g), saponification value (180–197 mg KOH/g), melting point (–14/–18 °C), refractive index (1.4800–1.4820 at 20 °C). The seed contains 41–45% of oil. The reported fatty acid composition is palmitic 5–8%, palmitoleic 0.5%, stearic 5–8%, oleic 4–10%, linoleic 34–44%, linolenic 44–49%.
More recently, some researchers investigated the impact of physical pretreatments on the quality of extracted PO (Siri et al., 2016), while others studied the possibility to carry out an aqueous enzyme-assisted process for oil recovery (Li et al., 2014).

During our research, we received a sample of PO for other studies about the preparation of a balanced blend (under the point of view of fatty acid composition), elsewhere published (Torri et al., 2019) and we took the opportunity to carry out a very detailed study on the composition of this interesting vegetable oil.

From the nutritional standpoint, an interest for this vegetable oil stands in its high concentration of α-linolenic acid (approximately 60%) and in the possibility to use it as an integrator contributing to re-balance the fatty acids ω6/ω3 ratio, strongly altered in the Western diet. In fact, our actual lipid nutrition is strongly oriented towards ω6 fatty acids, with a very low amount of ω3. The reason for this comes from their high tendency to oxidation and therefore in the practice to obtain processed food low in α-linolenic acid.

Nowadays, only soy and rapeseed oils, among the common commercial oils contain up to 10% of α-linolenic acid. For this reason, it is suggested to integrate the lipid profile using other oilseeds, such as linseed (Linum usitatissimum), sachie inchi (Plukenetia volubilis), walnut (Juglans regia), hemp (Cannabis sativa), camelina (Camelina sativa) and chia (Salvia hispanica).

In this paper, we describe the fatty acid composition of Perilla oil compared to other linolenic acid rich oils, like linseed, sachie inchi and chia: we also report the sterol and tocopherol composition, together with other relevant chemical parameters of PO.

2 Material and methods

2.1 Materials

Perilla seed oil was obtained from a Japanese factory located in Toyama prefecture. The oil was prepared by simple mechanical pressing using a hydraulic press operating batch wise at ambient temperature, without previous heating, with a seed load of approximately 2 kg of seed for each batch. The maximum operating pressure was 400 kg/cm² and the oil was filtered immediately after extraction to obtain a limpid product to be stored in dark containers.

We received a sample of this oil freshly prepared for our research.

2.2 Analytical methods

- Free fatty acids were determined by means of volumetric titration using phenolphthalein as an indicator using UNI EN ISO 660:2009 standard method.
- Peroxide value was determined by means of volumetric titration based on the liberation of iodine from potassium iodide in presence of hydroperoxides. A starch aqueous solution was used as an indicator according to UNI EN ISO 3960:2010.
- UV/Vis absorption: samples were dissolved in isoctane (1% concentration) and spectrophotometrically evaluated in the UV wavelength range using a cell having an optical path of 1 cm. The extinction coefficients were calculated respectively for the absorption at 232 and 270 nm, according to ISO 3656:2011.
- Fatty acid composition was determined according to ISO 12966-2:2017 + ISO 12966-4:2015. The analyses were carried out by a gas-chromatograph FOCUS (Thermoquest Instrument, Rodano, Italy) equipped with a flame ionization detector (GC-FID), using a capillary column (CP-Sil 88–1 = 100 m, 0.32 mm ID, film thickness 0.25 μm; Supelco, Bellefonte, PA, USA) after derivatization of fatty acids into the corresponding methyl esters, under the following experimental conditions: carrier gas He at a flow rate of 1.5 mL/min; split injection system with a splitting ratio 1:40; injector and detector temperatures set at 250 and 260 °C respectively; using the following program: 90–240 °C at 7 °C/min; injected quantity 1 μL.
- Sterol content and composition: the procedure, according to NGD 71-1989 + NGD 72-1989, is based on saponification, recovery of unsaponifiable fraction, purification of sterol fraction by preparative TLC, recovery and TMS derivatisation of the sterol, GC-FID evaluation by means of an internal standard (α-cholestanol). The sterol composition was determined using a gas-chromatograph TRACE (Thermoquest Instrument, Rodano, MI, Italy) equipped with a flame ionization detector. Analysis was carried out with a CPSil 8CB (Supelco, Bellefonte, PA, USA) capillary column (l = 30 m, 0.32 mm ID, film thickness 0.25 μm) under the following conditions: carrier gas He at a flow rate of 1.5 mL/min; split injection system with a splitting ratio 1:50; injector and detector temperatures set at 250 and 260 °C, respectively; oven isothermal temperature set at 240 °C; injected quantity 1 μL.
- Unsaponifiable matter, according to ISO 3956:2000: samples were treated with alcoholic KOH solution and the unsaponifiable matter was obtained by means of liquid-liquid extraction using diethyl ether. The quantification of unsaponifiable material was carried out gravimetrically.
- Tocopherols were evaluated according to ISO 9936:2016. A sample amount is simply diluted in hexane and injected in a HPLC system operating in direct phase with a silica column 4.6 mm ID × 250 mm length (Hypersil, Thermofisher, Rodano, Italy) using a mixture of hexane 99.5% and isopropanol 0.5% as isocratic eluting system with a flow rate of 1.0 mL/min; split injection system with a splitting ratio 1:50; injector and detector temperatures set at 250 and 260 °C, respectively; oven isothermal temperature set at 240 °C; injected quantity 1 μL. The quantification of tocopherols was obtained from Sigma Aldrich, Italy.
- Oxidation stability (Rancimat test, ISO 6886:2006): to determine the oxidation stability an accelerated oxidation test was performed. Oxidation induction times were measured by a Rancimat apparatus model 743 (Metrohm, Herisau, CH) using ISO 6886:2006 standard on 3 g of oil heated at 110 °C, under a purified air flow of air flow of 10 L/h.
- Biophenols (internal method based on COI T20 Doc. 29 2009 method). Biophenols were extracted from oil (2 g) using a solution of methanol/water 80/20 v/v (5 mL) and analysed by HPLC-PDA injecting 20 μL in the analytical
system at flow 1.0 mL/min. The analysis was performed using a gradient composed by water containing 0.2% H₃PO₄ (v/v), methanol and acetonitrile with a composition able to resolve most of the peaks present. A reverse phase column was used (Alltima–Grace, Italy 250 mm length × 4.6 mm ID, particle size 5 μm). The chromatogram was recorded at 280 nm and the quantitative content was reported as mg/kg of tyrosol for comparison with olive oil. All tests were carried out in duplicate under repeatability conditions.

3 Results and discussion

In Table 1, the fatty acid composition of Perilla frutescens oil is reported, in comparison with some of the possible competitors, linseed, sacha inchi and chia oils.

Looking at fatty acid composition we can say that PO contains the highest concentration of α-linolenic acid, if compared with linseed or sacha inchi oils, very close to the one of chia oil (Rodriguez et al., 2020), with a ratio between ω6 and ω3 fatty acids of 0.30. From the nutritional standpoint, ratios between 5 and 10 are recommended as optimal for human health (WHO Interim Summary, 2008). So, the value of 0.30 is quite far from the optimal one, but the use of PO for culinary or integration purposes could represent a way to contribute to re-equilibrate the ratio ω6/ω3 in the diet. Common vegetable oils such as olive, sunflower, corn, peanut, palm, are characterized by very high ω6/ω3 ratios, higher than the nutritionally optimal value. If we consider classical vegetable oils in use for human consumption only soybean and rapeseed oil demonstrate a correct ω6/ω3 ratio. However, all linolenic acid rich oils cannot be regarded as a substitute of (n-3) acids from fish oils such as eicosapentaenoic acid (EPA) and docosahexaenoic acid (DHA), but a very important complement (Cassiday, 2017; Stanley, 2009).

Like in all vegetable oils, the sterol composition is peculiar to each botanical family and in the case of Perilla frutescens is reported in Table 2. The most abundant sterol in PO sterol fraction is represented by beta-sitosterol, with important amounts of campesterol, delta 5 and delta 7 avenasterol, delta 7 stigmastenol. Very interesting is the high total sterol content that exceeds 3000 mg/kg. It is well known that phytosterols counteract cholesterol adsorption at intestinal level and this could represent an additional reason for the food use of this highly unsaturated oil.

The main chemical properties of the evaluated sample, representing a simply cold pressed and filtered oil, are resumed in Table 3.
We can observe that the sample was in good hydrolytic and oxidative conditions, as confirmed by the UV absorption data. Unsaponifiable and total sterols contents are in good agreement, while only few mg/kg of biophenols have been detected. The oxidation stability, as evaluated by means of Rancimat, is very low, because of the presence of high concentration of di- and triunsaturated fatty acids.

In Table 4, the results of tocopherols analysis are reported. The total content demonstrates that Perilla oil could represent a good source of vitamin E with a level similar to sunflower oils.

In practice, gamma-tocopherol represents more than 95% of this fraction, the remaining amount being covered by alpha and delta tocopherol. No tocotrienols were detected in the evaluated sample.

Moreover, free fatty alcohols (20.8 mg/kg), free sterols (2052.0 mg/kg), waxes C_{36}-C_{46} (26 mg/kg) concentrations were measured to complete the chemical characterisation, using the procedure suggested by Mariani et al. (1999).

Chlorophylls are only present in trace amounts as Phaeophytin B + B1 (0.1 mg/kg) and A + A1 (0.2 mg/kg). The evaluation was carried out using ISO 29841:2009 standard.

### 4 Conclusions

In this short paper, the main chemical properties of Perilla (Perilla frutescens) seed oil, obtained by mechanical pressing of Perilla seeds are reported. The analysis of fatty acid composition allowed to detect a huge amount of (n-3) α-linolenic acid (ALA), in concentration of approximately 60%, higher than other ALA rich oils such as linseed and sacha inchi oils, comparable with the one of chia. Perilla oil demonstrated an important sterol and vitamin E content, while biophenols are in a very low concentration. The analyzed sample showed very low acid and peroxide value and this is the demonstration that, following the proper technological procedures, it is possible to obtain good quality oils, even in presence of high α-linolenic acid concentrations. The flavor was evaluated as acceptable, even though this category of oil has not to be used as gourmet oils but as a supplement to cover the low intake of (n-3) fatty acids. The very high content of tocopherols (mainly in the γ form) allows a good antioxidative protection of the oil, but we must remember that highly unsaturated oils are very sensitive towards oxidation and must be in every case protected from the action of air and light, even using special containers or inert gases in packaging. Also preparing small packs may encourage a quick consumption once opened. Consuming highly oxidized oils may represent a hazard for human health.

### Conflict of interest

The authors declare that they have no conflicts of interest in relation to this article.

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