Alma Mater Studiorum Università di Bologna Archivio istituzionale della ricerca

Characterization of 'Olivastro di Bucchianico cv' extra virgin olive oils and its recognition by HS-GC-IMS

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

Published Version:

Characterization of 'Olivastro di Bucchianico cv' extra virgin olive oils and its recognition by HS-GC-IMS / Di Serio M.G.; Giansante L.; Del Re P.; Pollastri L.; Panni F.; Valli E.; Di Giacinto L.. - In: JOURNAL OF THE SCIENCE OF FOOD AND AGRICULTURE. - ISSN 0022-5142. - ELETTRONICO. - 101:14(2021), pp. 6074-6082. [10.1002/jsfa.11264]

Availability:

This version is available at: https://hdl.handle.net/11585/839880 since: 2021-11-29

Published.

DOI: http://doi.org/10.1002/jsfa.11264

Terms of use:

Some rights reserved. The terms and conditions for the reuse of this version of the manuscript are specified in the publishing policy. For all terms of use and more information see the publisher's website.

This item was downloaded from IRIS Università di Bologna (https://cris.unibo.it/). When citing, please refer to the published version.

(Article begins on next page)



Analytical study of cultivar "Olivastro di Bucchianico" extra virgin olive oils and its recognition by HS-GC-IMS

Journal:	Journal of the Science of Food and Agriculture
Manuscript ID	JSFA-20-4749
Wiley - Manuscript type:	Research Article
Date Submitted by the Author:	01-Dec-2020
Complete List of Authors:	Di Serio, Maria Gabriella; CREA, Research Centre for Engineering and Agro-Food Processing Giansante, Lucia; CREA, Research Centre for Engineering and Agro-Food Processing Del Re, Paolo; CREA, Research Centre for Engineering and Agro-Food Processing Pollastri, Luciano; Regione Abruzzo, Department of Agriculture Panni, Filippo; Università di Bologna, Department of Agricultural and Food Sciences, Alma Mater Studiorum Valli, Enrico; Università di Bologna, Department of Agricultural and Food Sciences and Interdepartmental Centre for Industrial Agro-food Research, Alma Mater Studiorum Di Giacinto, Luciana; CREA, Research Centre for Engineering and agro-food processing
Key Words:	Olea europaea L., single cultivar, characterization, sensory profile, volatile compounds, chemometric analysis

SCHOLARONE™ Manuscripts

- 1 Analytical study of cultivar "Olivastro di Bucchianico" extra virgin olive oils
- 2 and its recognition by HS-GC-IMS
- 3 Running Title: Cv Olivastro di Bucchianico recognition
- 4 Authors: Maria Gabriella Di Serio¹, Lucia Giansante¹, Paolo Del Re¹, Luciano Pollastri²,
- 5 Filippo Panni³, Enrico Valli⁴, Luciana Di Giacinto*,¹
- 6 ¹ CREA Research Centre for Engineering and Agro-Food Processing, I-65012 Cepagatti, PE, Italy
- Regione Abruzzo, I-65012 Cepagatti, PE, Italy
- ³Department of Agricultural and Food Sciences, Alma Mater Studiorum—Università di Bologna,
 47521 Cesena, Italy
- ⁴ Department of Agricultural and Food Sciences and Interdepartmental Centre for Industrial Agrofood
 Research, Alma Mater Studiorum—Università di Bologna, 47521 Cesena, Italy
- *Correspondence: Dr Luciana Di Giacinto (CREA Research Centre for Engineering and Agro-Food
 Processing (CREA-IT), I-65012 Cepagatti, PE, Italy Email: luciana.digiacinto@crea.gov.it)

20 ABSTRACT

- 21 Background
- 22 Single olive cultivar "Olivastro di Bucchianico" extra virgin olive oils, obtained from olives
- 23 cultivated in a restricted area of the Abruzzo region, Italy. Principally is present in the
- 24 municipality of Bucchianico and in some neighbouring municipalities in the province of
- 25 Chieti. There are very few research works in literature describing the morphological and
- 26 chemical characteristics of this cultivar.
- 28 Results

- 29 A morphological characterization of the plant and the fruit was carried out. In addition, extra
- 30 virgin olive oil was chemical, physical-chemical and sensory characterized. The conducted

analyses were as follows: free acidity, peroxide value, UV spectrophotometric indices, contents in fatty acid ethyl esters, waxes, tocopherols, fatty acids, triglycerides, sterols, alcohols, phenolic substances, volatile compounds and sensory profile. The analysis of the volatile compounds was performed using a HS-GC-IMS instrument connected to a nitrogen

Conclusion

The results of the chemical analyses showed good levels of nutraceutical components in the oils that resulted organoleptically well balanced with medium values of fruity, bitter and pungent. The HS-GC-IMS method based on the analysis of 15 volatile molecules might be a useful tool for a chemometric discrimination of the varietal origin for the oils under

- **Keywords:** Olea europaea L., single cultivar, characterization, sensory profile, volatile
- 45 compounds, chemometric analysis

generator for carrier/drift gas production.

47 INTRODUCTION

investigation.

- 48 The olive tree (Olea europaea L.) is the oldest fruit tree plant present in the environment
- 49 surrounding the Mediterranean and thanks to its nutritional properties and its high
- 50 economic value is rapidly expanding into new areas of cultivation.
- 51 The high environmental variability of the Italian territory allows the presence of many
- 52 cultivars of olive trees, estimated about 800 cultivars throughout in all the national
- 53 territory.^{1,2} Most probably, this number is even greater because there are many local
- 54 varieties of which few information is available. The enhancement and preservation of such a

great germplasm was very important in order to contain genetic erosion and diversify production on the territory with single cultivar EVOOs.³⁻⁷

A regular consumption of extra virgin olive oil has been widely demonstrated to prevent many diseases, not only due to its fatty acid composition abundant of monounsaturated fatty acids (MUFA), but also for minor components with health-related properties that can be defined as nutraceuticals: phenolic compounds, tocopherols, sterols.8-11 The concentration of these compounds is variable and strongly influenced by many factors such as cultivars, pedo-climatic environment, cultivation techniques, time and harvesting system, extraction technology. 12-14 Several studies have been carried out to correlate chemical composition of olive oil with the geographical origin.^{3, 4, 15} At the same time, identification of olive germplasm is complicated because no references and numerous cases of synonymy and homonymy for the same cultivars exist. To preserve this genetic diversity within the European Union, EVOOs have been "linked" to their territory of origin through the creation of standardized protocols of various types such as PDO (Protected Designations of Origin), PGI (Protected Geographical Indications) and finally TAF (Traditional Agricultural Food Products). In this framework, the production of single EVOO cultivars, with relative high consumption and market relevance, is very important for the protection of the typical cultivar of a specific area.

Morphological descriptors, even if for some traits influenced by external factors, was the first and principal step for study the genetic diversity within a cultivated plant species and they represent the phenological traits normally used in taxonomic classification.^{6,16} The study and recovery of minor cultivars present in particular cultivation areas, is very interesting and could be a resource to expand the offer of products to consumers. The cultivars are linked to specific environmental conditions, together with the continuous

extraction technology, which has a strong influence on the chemical characteristics, oxidative stability and organoleptic characteristics of extra virgin olive oils. 17,18

Recently an interesting analytical approach has been proposed, based on the determination

82 of volatile compounds using HS-GC-IMS (Gas Chromatography-Ion Mobility Spectrometry),

to support the organoleptic determination of virgin olive oils by panel test. This method can

84 realize a fingerprint of the aroma for a possible discrimination of the samples with respect to

85 the quality grade in a relatively simple, fast and economical way. 19,20

This work was made to analytical study native Olivastro di Bucchianico cultivar from a very restricted area of Abruzzo, Italy (Figure 1). There are very few research works in literature describing the morphological and chemical characteristics of this cultivar.^{2,21,22} So, the aim of this work was to do i) a morphological study of this underexploited local cultivar and chemical characterization of the produced extra virgin olive oils, ii) to use the HS-GC-IMS

MATERIALS AND METHODS

technique for varietal traceability.

95 Morphological description

The Olivastro di Bucchianico olive cultivar belongs to the municipality of Bucchianico and neighbouring municipalities in the province of Chieti. Bucchianico, with an altitude of 330 m, located in the northern part of the provincial territory close to the seaside resorts of Chieti and Villamagna. Its territory is mainly arable land, but with a large presence of olive groves and vineyards. The olive-growing area has a temperate hilly climate with winter temperatures around 6 °C in winter and around 23.5 °C in summer and relatively abundant rainfall. The Olivastro di Bucchianico olive trees have an assurgent bearing, an average

foliage and high vigour; the tree is of medium size, the fruiting branches have long internodes. The leaf is a shiny dark green colour, it is wide and short and has no curvature of the blade. The drupe is ellipsoidal with truncated base and sub-conical apex, of small to medium size ranging from about 1.89 to about 2.31 g, with pulp in percentage varying from 77.4 to 82.7 %. The drupes at the harvest, usually conducted in October, are never completely invaded, with prevalent green and vinous red colours. The drupes show high detachment resistance. The productivity of Olivastro di Bucchianico is medium-high but altering. These cultivars have a good resistance to parasites, other than caries.²²

111 Plant material

The olive fruits were handpicked in perfect sanitary conditions at the green stage of ripening at the mid and late October and stored in boxes and bins. Harvesting was performed with use of mechanical facilitators. Seven samples of EVOOs were produced by an extraction system that used two and two half phase centrifugation. The processing temperature was lower than 25° and the average crushing time lower than 45 minutes. The average oil yield was 15%. The oil samples have been preserved in low temperature (15 – 18 °C) in dark-green glass bottles without headspace before the analysis.

Methods

Free acidity, peroxide value, UV spectrophotometric indices, content in fatty acid ethyl esters (FAEEs), waxes, tocopherols, alcohols, phenolic substances, and volatile compounds, fatty acid and triglyceride compositions, sterol composition and content, and sensory profile were performed on single cultivar EVOO "Olivastro di Bucchianico" samples have undergone analyses. Duplicate analyses were performed on each sample.

Analytical parameters

Free acidity (Annex II), peroxide value (Annex III), ultraviolet spectrophotometric indices

(Annex IX), ethyl esters and waxes contents (Annex XX), fatty-acid composition (Annex X.B),

triglyceride composition (Annex XVIII), sterol composition and content (Annex V), and

alcohols content (Annex XIX). were performed according to the official methods of the

European Union Commission Regulation EEC/2568/91 and its subsequent modifications.²³

Tocopherols

For the HPLC determination of the tocopherols²⁴, a solution of 1 g olive oil in 10 mL acetone was prepared. α -Tocopherol (0.32 $\mu g/10 \mu L$) was used as the external standard. The HPLC analysis was conducted using a high-resolution LC 200 liquid chromatograph equipped with a Series 200 UV/Vis detector (Perkin Elmer, Waltham, Massachusetts, USA), a 7725 Rheodyne injector, a 20 μL sample loop, and a Totalchrom workstation for data acquisition (Perkin Elmer, Waltham, Massachusetts, USA). Separation on a Spherisorb ODS2 column (250 x 4.6 mm I.D., 5µm; Waters, Milford, MA, USA) was performed at 25 °C under a constant flow rate of 1 mL/min with a mobile phase of 0.2% (v/v) H₃PO₄ in water/ methanol/ acetonitrile (2/49/49, v/v/v). The eluted compounds were detected at 292 nm.

Phenols composition

The phenolic compounds were extracted from the olive oil according to the following method: a 500 μ L volume of internal standard solution (0.015 mg/mL syringic acid in methanol) was added to 2.500 g of oil sample. After removal of the methanol under reduced pressure at <35 °C, the samples were dissolved in 6 mL hexane and loaded onto solid-phase extraction columns (Discovery DSC-DIOL 500 mg, 3 mL; Supelco, Bellefonte, PA, USA), which were previously conditioned with 6 mL methanol and 6 mL hexane. The samples were then washed with 2 x 3 mL hexane and 4 mL hexane/ ethyl acetate (90/10, v/v), and then eluted

with 10 mL methanol. After removal of the methanol in a rotary evaporator at a temperature < 35 °C, the dry residue was taken up with 1 mL methanol/ water (1:1, v/v), filtered through a 0.45 µm PVDF membrane, and then injected into the HPLC system. The HPLC analysis of the phenolic extracts was carried out using a high-resolution LC 200 liquid chromatograph, which was equipped with a Series 200 UV/Vis detector (Perkin Elmer, Waltham, Massachusetts, USA), a 7725 Rheodyne injector, and a 20 µL sample loop, using a Totalchrom workstation for data acquisition (Perkin Elmer, Waltham, Massachusetts, USA). Separation on a Spherisorb ODS2 column (250 x 4.6 mm I.D., 5 μm; Waters, Milford, MA, USA) was performed at 25 °C at a constant flow rate of 1 mL/min, with mobile phase mixtures composed of A (0.2% [v/v] H₃PO₄ in water), B (methanol) and C (acetonitrile), at the following ternary gradient program (as A/B/C, v/v/v): 0 min, 96/2/2; 40 min, 50/25/25; 45 min, 40/30/30; 60 min, 0/50/50; 70 min, 0/50/50; 72 min, 96/2/2; 82 min, 96/2/2. The eluted compounds were detected at 280 nm. Previously, 20 μL external calibration standard was injected into the HPLC system, to calculate the response factors of syringic acid to tyrosol. Then, a volume of 20 µL each sample was injected. Quantification of the phenolic compounds, expressed as tyrosol as reported in COI/T.20/Doc No 29/2009²⁵, was performed according to the concentration of the internal standard and on the basis of the response factor of syringic acid to tyrosol.

Volatile compounds: HS-GC-IMS analysis

170 The analysis of the volatile compounds was performed using a HS-GC-IMS Flavourspec®
171 instrument (G.A.S. Dortmund, Dortmund Germany) connected to a nitrogen generator for
172 carrier/drift gas production (Microprogel, Pordenone, Italy). 2.0 g of each sample were
173 weighted, from which 100 μL of the respective headspace was withdrawn using a 2.5 mL

Hamilton syringe with a 51 mm needle, through an autosampler unit, HT2000H (HTA s.r.l., Brescia, Italy), and introduced in a splitless heated injector (2 mm ID, 6.5 mm OD × 78.5 mm fused quartz glass). From here, the analytes of the headspace are subjected to a double sequential separation: GC, passed into a low polar column FS-SE-54-CB-0.5, 30 m, 0.32 mm ID, film thickness 0.5 µm (94% methyl-5% phenyl-1% vinylsilicone); IMS (equipped with a tritium ionizing radioactive source at 5000 V), passed into a 9.8 cm long drift tube (Gesellschaft für Analytische Sensorsysteme mbH, G.A.S.; Dortmund, Germany).²⁰

Sensorial profile

The evaluation of olive oils was performed under the conditions described in European Union Commission Regulation EEC/2568/91 and its subsequent modifications (Annex XII) by the CREA-Research Centre for Engineering and Agro-Food Processing of Pescara Panel recognized by the International Olive Oil Council (IOC) and the Ministry of Agricultural, Food and Forestry Policies (MiPAAF). Each taster of the panel has smelled and tasted the oil under consideration, according to the profile sheet of the Annex XII and to COI/T.20/DOC. 22 -2005.²⁶ The attributes evaluated were fruity, pungent, bitter, aromatic herbs, tomato, artichoke, almond and grass/leave. Each attribute was evaluated on a scale from 0.0 to 10.0 and statistically processed by calculation of median; the confidence intervals was used considering the attributes with a robust coefficient of variation of 20.0 % or less.

Statistical analyses

A heat map (3D chromatogram) was obtained from each sample, from which it was possible to extrapolate a data matrix. Subsequently, processing this data matrix using statistical software Unscrambler X, version 10.4, it was possible to perform a PCA (Principal 197 Component Analysis).

RESULTS AND DISCUSSION

The analyses for quality indices (Table 1) allowed for all samples of "Olivastro di Bucchianico" to be classified as EVOOs according to EC Regulation (EEC, 1991 and following amendments). Free acidity of the samples was in the range from 0.08 to 0.24 % of oleic acids, peroxide values between 6.2 to 8.8 meq O_2 /kg. Mean values of the specific extinction parameters K_{232} , K_{270} and ΔK were respectively 1.66, 0.12 and -0.003, so within the EVOO limit according to EC regulation. The FAEE content of 2.6 mg/kg is well below the established EU limit for EVOO (35 mg/kg). These fat-soluble compounds are a valuable indicator of oil quality resulting from improper agronomic and technological practices, in particular from fermentation and degradation processes.^{27,28} The waxes content was low, with a range from 27 to 40 mg/kg, typical of extra virgin olive oils. Total tocopherols content was medium-high, around 264 mg/kg (range 218-337 mg/kg). Composition and content of these substances depend e.g. on several agronomic factors, olive processing conditions and the storage.²⁹⁻³² Extra virgin olive oil is one of the foods richest in vitamin E (tocopherols) and the most representative is α -tocopherol, which accounts for about 90% of the total tocopherols with vitaminic and antioxidant actions. The fatty acid composition presented in Table 2 was within EU Regulation limits for olive oils. Oleic, palmitic and linoleic and stearic acids were the most abundant with mean values respectively of 74.71%, 11.94%, 7.81 % and 3.02 %.³³ These values are typical of olive oils: oleic acid not less than 73% and linoleic acid not more than 10% with an oleic/linoleic ratio > 7. Monounsaturated fatty acids values (Σ MUFAs) in "Olivastro di Bucchianico" cv was high, mean around 75.88% with the maximum value 78.14% and minimum value 73.82%. The oleic acid is extremely important for a healthy and balance diet, leading to an increase in oxidative stability, antihypertensive activity, and low-density lipoprotein (LDL) cholesterollowering effects.³⁴ In defining the nutritional properties and varietal characterization of the EVOOs, it is also important MUFA/SFAs ratio, that in the single cultivar studied is average of 4.82.35 The composition of triglycerides, useful for the characterization of single cultivar, was reported in Table 3. As can be seen from this table the most represented compounds are: PLP + OOO + PoPP (42.48%), SOL + POO (23.61%) and OOL + LnPP (12.07%). Sterols composition of "Olivastro di Bucchianico" was shown in Table 4. The most representative sterols have been β -sitosterol (85.8%), Δ -5-avenasterol (6.5%) and campesterol (3.0%). The value of apparent β -sitosterol, the sum of β -sitosterol and four adjacent phytosterols (clerosterol, sitostanol, Δ -5-avenasterol and Δ -5,24-stigmastadienol), was 94.6% superior at EC legal minimal limits established. High levels of apparent β-sitosterol have a positive effect on reducing total plasma cholesterol and LDL cholesterol. 36,37 Total sterols (1274 mg/kg) were superior to the lower limit established by EC legislation (1000 mg/kg). Sterols and triglycerides can characterize the different EVOOs. 38,39 In the unsaponifiable fraction of olive oil, the presence of both diterpenic alcohols (with 20 carbon atoms) and triterpenic alcohols (with 30 carbon atoms) has been determined (Table 5), which make up to 25-30%. Among the first ones the most represented is the Phytol (318.4 mg/kg), partly coming from the degradation of chlorophyll. The composition of the triterpenic fraction seems to be specific to the botanical family from which the oil is derived.⁴⁰ It is mainly composed of cycloartenol (380.4 mg/kg) and 24-Methylen-cycloartanol (317.9 mg/kg) followed by β -amyrin + butyrrospermol (90.2 mg/kg) and α -amyrin (1.2 mg/kg). The alcoholic composition also includes citrostadienol (218.8 mg/kg), a 4-methylsterol, and total aliphatic alcohols whose content (140 mg/kg) is below the limit laid down in the EU standard for EVOOs. In "Olivastro di Bucchianico" the most abundant

phenols (Table 6) are 3,4 DHPEA-EDA (76.3 mg/kg), p-HPEA-EDA (65.3 mg/kg), pinoresinol and 1-acetoxypinoresinol (58.9 mg/kg), 3,4-DHPEA-EA,H (37.8 mg/kg). Many studies report how derivatives of oleuropein and ligstroside, 3,4-DHPEA-EA and p-HPEA-EDA show antioxidant and health properties. 41,42 Moreover, secoiridoids and lignans have effect on the bitter and pungent attributes in EVOOs. 43-45 The total phenols content was medium (306 mg/kg).46,47 In Figure 2 sensory profile of "Olivastro di Bucchianico" was reported. It was characterized by medium perceptions of fruity, with well-balanced notes of bitter and pungent. Medium notes of grass/leave and almond were present with mean value respectively 2.2 and 3.0 and low notes of artichoke, aromatic herbs and tomato with mean value respectively 2.2, 0.6 and 0.6. In order to verify the discriminatory potential of the HS-GC-IMS method on EVOOs oils on the basis of the cultivars they belong to, Olivastro di Bucchianico samples were analyzed together with other single cultivar oils from Central and Southern Italy. In particular, 10 EVOOs of the Canino, Ogliarola, Coratina, Moraiolo and Peranzana cultivars (2 samples of each cultivar) were subjected to HS-GC-IMS analysis at the same time as Olivastro. From the heat maps obtained, only 15 volatile compounds, selected within the European H2020 project OLEUM and commented on Valli et al., 2020, were considered. Their respective signals present in the form of a monomer and/or dimer in the chromatogram were highlighted using VOCal software (Gesellschaft für Analytische Sensorsysteme mbH, G.A.S.; Dortmund, Germany). Using a specific function of this software, it was possible to export the results as data matrix (Table 7 and Table 8) which was used for the construction of a PCA. Figure 3 shows the scores plot obtained from the PCA (explained variance PC1 and PC2: 70%). The 7 Olivastro di Bucchianico EVOOs resulted separated from the other 10 analyzed

samples, showing a promising discrimination of this cultivar based on the volatile profile with respect to the others.

CONCLUSIONS

The herein performed analytical study conducted on cv. Olivastro di Bucchianico EVOOs, obtained from olives cultivated in restricted area of the Abruzzo region (Italy) highlighted interesting compositional characteristics. Samples showed an average medium-high content of total tocopherols (264 mg/kg), high monounsaturated fatty acids values, around 75.88% and a good oleic/linoleic ratio (9.57). The average total phenols content was medium, 306 mg/kg. Sensory profile of "Olivastro di Bucchianico" was characterized by medium perceptions of fruity, with well-balanced notes of bitter and pungent, accentuated hints of grass/leave and almond with mean value respectively, 2.2 and 3.0 and low intensity notes of artichoke, aromatic herbs and tomato. The research was extended to the analysis of volatile compounds by HS-GC-IMS for verifying the discriminatory potential of the method according to the cultivar. In particular, Olivastro di Bucchianico cy EVOOs samples were analyzed as well as other 10 single cultivars of the Canino, Ogliarola, Coratina, Moraiolo and Peranzana cv. The chemometric analysis of the data allowed to distinguish the Olivastro di Bucchianico cv EVOOs from the others. This preliminary study put in evidence the compositional characteristics of the studied single cultivar EVOOs, in order to propose a characterization study to the competent institutions. This contribution can also be evaluated for the purpose of defining the specifications PDOs, PGIs, PATs.

291 ACKNOWLEDGMENTS

The authors thank Cesare Rossini (LabService Analytica S.R.L., Bologna, Italy) for the concession of the HS-GC-IMSinstrument. The set-up of the HS-GC-IMS analytical method as well as the selection of the 15 volatile compounds was performed within the EU Horizon 2020 project "Advanced solutions for assuring the overall authenticity and quality of olive oil (OLEUM) – Grant Agreement number: 635690".

REFERENCES

- 299 1. Miazzi MM, Di Rienzo V, Mascio I, Montemurro C, Sion S, Sabetta W, Vivaldi GA,
 300 Camposeo S, Caponio F, Squeo, G, Difonzo G, Loconsole G, Bottalico G, Venerito P, Montilon
 301 V, Saponari A, Altamura G, Mita G, Petrontino A, Fucilli V and Bozzo F, Re. Ger. O.P.: an
 302 integrated project for the recovery of ancient and rare olive germplasm. *Frontiers in Plant*
- *Science* **11**:128. (2020) https://doi.org/10.3389/fpls.2020.00073.
- Muzzalupo I, Olive Germplasm–Italian Catalogue of Olive Varieties, ed. By Muzzalupo Intech, Italy, pp. 249-250. (2012). http://dx.doi.org/10.5772/51719.
- 306 3. Di Serio MG, Di Giacinto L, Di Loreto G, Giansante L, Pellegrino M, Vito R and Perri E, 307 Chemical and sensory characteristics of Italian virgin oils from Grossa di Gerace *cv. Eur J Lipid* 308 *Sci Technol* 118:288-298 (2016).
- Cecchi T, Passamonti P, Alfei B and Cecchi P, Monovarietal Extra Virgin Olive Oils from
 the Marche region, Italy: analytical and sensory characterization. *Int J Food Prop* 14:483–495
 (2011).
- 5. Di Loreto G, Di Giacinto L, Zema V, Pollastri L, Serraiocco A and Giorgetti M, ed. ARSSA. *Primo anno di studio sulla caratterizzazione di oli vergini di oliva delle varietà* "Rustica" e "Gentile dell'Aquila" in Valle Peligna (Abruzzo). "La caratterizzazione degli oli vergini di oliva in Valle Peligna". Raiano, L'Aquila, Italy (2011).

- 316 6. Rotondi A, Magli M, Ricciolini C and Baldoni L, Morphological and molecular analyses
- for the characterization of a group of Italian olive cultivars. *Euphytica* 132:129–137 (2003).
- 318 7. Stefanoudaki E, Kotsifaki F and Koutsaftakis A, Sensory and chemical profiles of three
- 319 European olive varieties (Olea europea L.): an approach for the characterisation and
- authentication of the extracted oils. *J Sci Food Agric* 80:381-389 (2000).
- 321 8. Marcelino G, Aiko Hiane P, De Cássia Freitas K, Figueiredo Santana L, Pott A,
- Rodrigues Donadon J and de Cássia Avellaneda Guimarães R, Effects of Olive Oil and Its
- 323 Minor Components on Cardiovascular Diseases, Inflammation, and Gut Microbiota. *Nutrients*
- **11(8):1826 (2019)**.
- 325 9. Buckland G and Gonzalez CA, The role of olive oil in disease prevention: a focus on
- 326 the recent epidemiological evidence from cohort studies and dietary intervention trials. Br J
- 327 Nutr 94:101 (2015). doi: 10.1017/S0007114514003936
- 328 10. Brunelleschi S, Amoruso A, Bardelli C, Romani A, Ieri F and Franconi F, Chapter 117 -
- 329 Minor Polar Compounds in Olive Oil and NF- kB Translocation. In Olives and Olive Oil in
- *Health and Disease Prevention.* **1079-1086 (2010)**.
- 331 11. Saldeen K and Saldeen T, Importance of tocopherols beyond α -tocopherol: evidence
- 332 from animal and human studies. *Nutr Res* 25 (10):877-889 (2005).
- 333 12. Bruno L, Picardi E, Pacenza M, Chiappetta A, Muto A, Gagliardi O, Muzzalupo I, Pesole
- 334 G and Bitonti MB, Changes in gene expression and metabolic profile of drupes of Olea
- 335 europaea L. cv Carolea in relation to maturation stage and cultivation area. BMC Plant Biol
- **19:428-445 (2019)**.

- 13. El Qarnifa S, El Antari A and Hafidi A, Effect of Maturity and Environmental Conditions
 338 on Chemical Composition of Olive Oils of Introduced Cultivars in Morocco. *J Food Qual* 1-14
- 339 (2019).
- 340 14. Marra FP, Buffa R, Campisi G, Costa F, Di Vaio C, La Farina M, La Mantia M, Mafrica R,
- 341 Motisi A, Zappia R and Caruso T, Morphological and SSR molecular markers based genetic
- 342 variability in 39 olive cultivars (Olea europea L.) originated in Southern Italy. Second
- 343 International Seminar Olivebioteq. Marsala-Mazara del Vallo, vol. I:213-216 [5-10 november
- **2006**].
- 345 15. Kotti F, Chiavaro E, Cerretani L, Barnaba C, Gargouri M and Bendini A, Chemical and
- thermal characterization of Tunisian extra virgin olive oil from Chetoui and Chemlali cultivars
- and different geographical origin. Eur Food Res Technol 228:735-742 (2009).
- 348 16. Blazakis KN, Kosma M, Kostelenos G, Baldoni L, Bufacchi M and Kalaitzis P,
- 349 Description of olive morphological parameters by using open access software. Plant Meth
- 350 13(1):111 (2017). doi: 10.1186/s13007-017-0261-8.
- 351 17. Romero N, Saavedra J, Tapia F, Sepúlveda B and Aparicio R, Influence of agroclimatic
- 352 parameters on phenolic and volatile compounds of Chilean virgin olive oils and
- 353 characterization based on geographical origin, cultivar and ripening stage. J Sci Food Agric
- 354 96:583–592 (2016).
- 355 18. Rotondi A, Alfei B, Magli M and Pannelli G, Influence of genetic matrix and crop year
- 356 on chemical and sensory profiles of Italian monovarietal extra-virgin olive oils. J Sci Food
- *Agric* 90(15):2641-2648 (2010).

- 358 19. Contreras MDM, Arroyo-Manzanares N, Arce C and Arce L, HS-GC-IMS and
- 359 chemometric data treatment for food authenticity assessment: Olive oil mapping and
- classification through two different devices as an example. *Food Control* 98:82-93 (2019).
- 361 20. Valli E, Panni F, Casadei E, Barbieri S, Cevoli C, Bendini A, García-González DL and
- 362 Gallina Toschi T, An HS-GC-IMS method for the quality classification of virgin olive oils as
- 363 screening support for the panel test. *Foods* 9:657 (2020). doi: <u>10.3390/foods9050657</u>.
- 364 21. Muzzalupo I, Salimonti A, Caravita MA, Pellegrino M and Perri E, SSR markers for
- 365 characterization and identification of cultivars of Olea Europaea L. in the Abruzzo and Molise
- 366 regions in south-central Italy. *Adv Hort Sci* 22(2):129–135 (2008).
- 367 22. Pietrangeli F and Russo A, ed. by Regione Abruzzo. Olivi d'Abruzzo Contributo alla
- 368 conoscenza del germoplasma olivicolo autoctono, 2nd edn. Guardiagrele, Chieti, Italy (2004).
- 369 23. REG. EEC/2568 Consolidated version of the Commission Regulation EEC No 2568/91
- of 11 July 1991 on the characteristics of olive oil and olive-residue oil and on the relevant
- 371 methods of analysis (1991).
- 372 24. Rovellini P, Azzolini M and Cortesi N, Tocoferoli e tocotrienoli in oli e grassi vegetali
- 373 mediante HPLC. Riv Ital Sost Grasse 74:1-5 (1997).
- 374 25. International Olive Oil Council, Determination of Biophenols in Olive Oils by HPLC,
- 375 COI/T20/Doc 29, (2009).
- 376 26. International Olive Oil Council, Method for the Organoleptic Assessment of Extra
- 377 Virgin Olive Oil Applying to Use a Designation of Origin, COI/T20/Doc 22, (2005).
- 378 27. Di Giacinto L, Di Loreto G, Di Serio MG, Giansante L, Faberi A, Marianella RM,
- 379 Ricchetti L, Perri E, Serraiocco A and Vito R, ed. by CREA. Monitoraggio degli alchil esteri (AE)

- 380 nell'olio extra vergine di olive. "MONITORALCHIL" Risultati dell'attività sperimentale 2012-
- *2014*, Roma, Italy (2014).
- 382 28. Mariani C and Bellan G, Sul possibile aumento degli alchil esteri negli oli extra vergini
- 383 di oliva. *Riv Ital Sost Grasse* 1:3-10 (2011).
- 384 29. Jukić Špika M, Kraljić K and Škevin D, Tocopherols: Chemical Structure, Bioactivity,
- and Variability in Croatian Virgin Olive Oils, in Products from Olive Tree. Chapter 17, pp. 317-
- 386 329 (2016).
- 387 30. Bengana M, Bakhouche A, Lozano-Sánchez J, Youcef, AY, Youyou A, Segura-Carretero
- 388 A and Fernández-Gutiérrez A, Influence of olive ripeness on chemical properties and
- 389 phenolic composition of Chemlal extra-virgin olive oil. Food Research Int 2013, 54:1868-
- 390 1875 (2013). doi:10.1016/j.foodres.2013.08.037
- 391 31. Beltrán G, Jiménez A, Del Rio C, Sánchez S, Martínez L, Uceda M and Aguilera MP,
- 392 Variability of vitamin E in virgin olive oil by agronomical and genetic factors. J Food Compos
- *Anal* 23(6):633–639 (2010). doi:10.1016/j.jfca.2010.03.003
- 394 32. Deiana M, Rosa A, Falqui Cao C, Pirisi FM, Bandino G and Dessì MA, Novel approch to
- 395 study oxidative stability of extra virgin olive oils: importance of alpha-tocopherol
- 396 concentration. *J Agric Food Chem* 50:4342-4346 (2002).
- 397 33. Marongui B, Özcan MM, Rosa A, Dessi MA, Piras A and Al Juhaimi F, Monitoring of
- 398 the fatty acid compositions of some olive oils. Riv Ital Sost Grasse. Vol.XCII:39-42 (2015).
- 399 34. Psaltopoulou T, Naska A, Orfanos P, Trichopoulos D, Mountokalakis T and
- 400 Trichopoulou A, Olive oil, the Mediterranean diet, and arterial blood pressure: the Greek
- 401 European prospective investigation into cancer and nutrition (EPIC) study. Amer J Clin Nutr
- 402 80(4):1012–1018 (2004).

- 403 35. Montaño A, Hernández M, Garrido I, Llerena JL and Espinosa F, Fatty acid and
- 404 phenolic compound concentrations in eight different monovarietal virgin olive oils from
- 405 Extremadura and the relationship with oxidative stability. Int J Mol Sci 17 (11):1960 (2016).
- 406 doi:10.3390/ijms17111960
- 407 36. Mena C, González AZ, Olivero-David R and Pérez-Jiménez MA, Characterization of
- 408 'Castellana' virgin olive oils with regard to olive ripening. HortTechnology 28(1):48-57 (2018).
- 409 37. St-Onge MP, Lamarche B, Mauger JF and Jones PJH, Consumption of a functional oil
- 410 rich in phytosterols and medium-chain triglyceride oil improves plasma lipid profiles in men.
- 411 J Nutr 133(6):1815–1820 (2003).
- 412 38. Yorulmaz A, Yavuz H and Tekin A, Characterization of Turkish Olive Oils by
- Triacylglycerol Structures and Sterol Profiles. J Am Oil Chem Soc 91:2077–2090 (2014).
- 414 39. Galeano Diaz T, Durán Merás I, Sánchez Casas J and Alexandre MF, Characterization
- 415 of virgin olive oils according to its triglycerides and sterols composition by chemometric
- 416 methods. Food Control 16:339-347 (2005).
- 417 40. Ben Temime S, Manai H, Abaza L, Baccouri B, Daoud D and Zarrouk M, Sterol and
- 418 triterpene alcohols profile of Chètoui virgin olive oils. Second International Seminar
- 419 Olivebioteq 2006 pp 481-483 (2006).
- 420 41. El Riachy M, Priego-Capote F, León L, Rallo L and Luque de Castro MD, Hydrophilic
- 421 antioxidants of virgin olive oil. Part 2: Biosynthesis and biotransformation of phenolic
- 422 compounds in virgin olive oil as affected by agronomic and processing factors. Eur J Lipid Sci
- *Technol* 113:692-707 (2011).
- 42. Servili M, Selvaggini R, Esposto S, Taticchi A, Montedoro GF and Morozzi G, Health
- 425 and sensory properties of virgin olive oil hydrophilic phenols: agronomic and technological

- 426 aspects of production that affect their occurrence in the oil. J Chromatogr A. 1054 (1-2):113-
- 427 127 (2004).
- 428 43. Lanza B, Di Serio MG, Giansante L, Di Loreto G and Di Giacinto L, Effect of shelf
- 429 conditions on the phenolic fraction and oxidation indices of monovarietal extra virgin olive
- 430 oil from cv. Taggiasca. *Acta Aliment* 44 (4):585-592 (2015).
- 431 44. Esti M, Contini M, Moneta E and Sinesio F, Phenolics compounds and temporal
- 432 perception of bitterness and pungency in extra-virgin olive oils: Changes occurring
- 433 throughout storage. Food Chem 113:1095-1100 (2009).
- 434 45. Preziuso SM, Di Loreto G and Biasone A, Studio delle correlazioni tra le intensità degli
- 435 attributi organolettici di amaro e piccante e le concentrazioni dei composti che ne sono
- 436 responsabili. Abstracts Book Primo Convegno Nazionale dell'Olivo e dell'Olio di Portici (NA).
- 437 pp.70. (2009).
- 438 46. Montedoro G, Servili M, Baldioli M and Miniati E, Simple and hydrolyzable phenolic
- 439 compounds in virgin olive oil. 1. Their extraction, separation, and quantitative and
- 440 semiquantitative evaluation by HPLC. J Agric Food Chem 40:1571-1576 (1992).
- 441 https://doi.org/10.1021/jf00021a019.
- 442 47. Ghanbari Shendi E, Sivri Ozay D and Taha Ozkaya M, Effects of filtration process on
- 443 the minor constituents and oxidative stability of virgin olive oil during 24 months storage
- 444 time. OCL. 27:37 (2020). http://doi.org/10.1051/ocl/2020030.

446 FIGURE LEGENDS

- **Figure 1** Olivastro di Bucchianico cultivation area in the Abruzzo region (Italy)
- 448 Figure 2 Sensory profile of the EVOOs from Olivastro di Bucchianico cv

449 Figure 3 Scores plot obtained from the PCA

- 451 APPENDIX: ABBREVIATIONS USED
- 452 EVOOs, Extra Virgin Olive Oils, FAEE, Fatty Acid Ethyl Esters, SFAs, Saturated Fatty Acids,
- **MUFAs,** Monounsaturated Fatty Acids, **PUFAs,** Polyunsaturated Fatty Acids, **PCA,** Principal
- 454 Component Analysis.



Table 1. Qualitative characteristics of the EVOOs from Olivastro di Bucchianico cv.

Determinations:	Mean	Max	Min	S. D.
Free acidity (% oleic acid)	0.15	0.24	0.08	0.06
Peroxide value (mEqO ₂ /kg)	7.5	8.8	6.2	1.0
K_{232}	1.66	1.98	1.44	0.17
K_{270}	0.12	0.20	0.07	0.043
ΔK	-0.003	-0.002	-0.004	0.001
FAEE (mg/kg)	2.6	4.3	0.1	1.7
Waxes (mg/kg)	33	40	27	5
Tocopherols (mg/kg)	264	337	218	47

S.D.: standard deviation

Table 2. Fatty acid composition of the EVOOs from Olivastro di Bucchianico cv.

Fatty acid composition (%):	Mean	Max	Min	S.D.
Myristic acid (C14:0)	0.01	0.01	0.00	0.004
Palmitic acid (C16:0)	11.94	14.14	10.89	1.30
Palmitoleic acid (C16:1)	0.71	1.00	0.52	0.19
Heptadecanoic acid (C17:0)	0.12	0.15	0.08	0.03
Heptadecenoic acid (C17:1)	0.18	0.26	0.12	0.04
Stearic acid (C18:0)	3.02	3.17	2.82	0.14
Oleic acid (C18:1)	74.71	77.12	72.46	1.66
Linoleic acid (C18:2)	7.81	8.32	6.63	0.60
Arachic acid (C20:0)	0.46	0.54	0.42	0.04
Linolenic acid (C18:3)	0.58	0.64	0.53	0.04
Eicosenoic acid (C20:1)	0.27	0.31	0.24	0.03
Behenic acid (C22:0)	0.11	0.19	0.06	0.04
Lignoceric acid (C24:0)	0.08	0.15	0.04	0.04
Oleic/linoleic	9.57	10.93	9.27	0.99
\sum SFAs	15.74	17.69	14.69	1.11
\sum MUFAs	75.88	78.14	73.82	1.51
\sum PUFAs	8.39	8.93	7.17	0.62
MUFAs/SFAs	4.82	5.03	4.42	0.42

SFAs: saturated fatty acids; MUFAs: monounsaturated fatty acids; PUFAs: polyunsaturated fatty acids; S.D.: standard deviation

Table 3. Triglycerides composition of the EVOOs from Olivastro di Bucchianico cv.

Triglycerides composition (%):	Mean	Max	Min	S.D.
LLL	0.08	0.17	0.03	0.06
OLLn+PoLL	0.14	0.27	0.08	0.07
PLLn	0.06	0.07	0.02	0.02
OLL	1.50	1.65	1.12	0.19
OOLn+PoOL	1.18	1.33	1.08	0.09
PLL+PoPoO	0.49	0.63	0.28	0.12
POLn+PPoPo+PPoL	0.61	0.75	0.52	0.07
OOL+LnPP	12.07	13.00	11.14	0.64
PoOO	1.10	1.65	0.69	0.39
SLL+PLO	5.61	6.45	4.49	0.71
PoOP+SPoL+SOLn+SPoPo	0.70	0.98	0.53	0.18
PLP+OOO+PoPP	42.48	46.38	39.35	2.77
SOL+POO	23.61	25.33	22.31	1.23
POP	3.88	5.01	3.29	0.67
SOO	5.21	5.72	4.76	0.43
POS+SLS	1.28	1.52	1.18	0.11

P: palmitic; Po: palmitoleic; S: stearic; O: oleic; L: linoleic acids; S.D.: standard deviation

Table 4. Sterol composition and erythrodiol and uvaol content of the EVOOs from Olivastro di Bucchianico cv.

Determinations:	Mean	Max	Min	S.D.
Sterol composition (%):				
Colesterol	0.2	0.4	0.1	0.1
Campesterol	3.0	3.2	2.9	0.1
Campestanol	0.0	0.1	0.0	0.0
Stigmasterol	0.7	0.8	0.6	0.1
Δ -7-Campesterol	0.1	0.1	0.0	0.0
Δ -5.23-Stigmastadienol	0.0	0.1	0.0	0.0
Clerosterol	0.6	0.8	0.4	0.2
β-Sitosterol	85.8	88.5	80.3	3.0
Sitostanol	1.0	1.1	0.7	0.1
Δ-5-Avenasterol	6.5	11.5	3.7	2.9
Δ-5.24-Stigmastadienol	0.8	1.0	0.6	0.2
Δ-7-Stigmastenol	0.5	0.8	0.3	0.2
Δ-7-Avenasterol	0.8	1.1	0.7	0.1
Apparent β-Sitosterol	94.6	95.0	94.3	0.2
Total sterols (mg/kg)	1274	1409	1099	128
Erythrodiol +Uvaol (%)	1.48	1.98	1.01	0.35

S.D.: standard deviation

Table 5. Alcoholic content of EVOOs from Olivastro di Bucchianico cv.

Mean	Max	Min	S.D.
318.4	418.8	205.3	85.5
14.5	19.6	10.4	2.9
5.2	6.9	3.9	1.2
22.1	27.8	16.1	5.1
66.7	84.3	50.1	16.0
28.1	41.4	20.1	8.4
90.2	127.3	55.2	29.9
1.2	3.5	0.1	1.2
380.4	516.2	200.5	128.3
317.9	420.9	168.8	114.6
218.8	289.9	145.4	55.8
140	187	94	39
	318.4 14.5 5.2 22.1 66.7 28.1 90.2 1.2 380.4 317.9 218.8	318.4 418.8 14.5 19.6 5.2 6.9 22.1 27.8 66.7 84.3 28.1 41.4 90.2 127.3 1.2 3.5 380.4 516.2 317.9 420.9 218.8 289.9 140 187 andard deviation	318.4 418.8 205.3 14.5 19.6 10.4 5.2 6.9 3.9 22.1 27.8 16.1 66.7 84.3 50.1 28.1 41.4 20.1 90.2 127.3 55.2 1.2 3.5 0.1 380.4 516.2 200.5 317.9 420.9 168.8 218.8 289.9 145.4 140 187 94

Table 6. Phenols content of EVOOs from Olivastro di Bucchianico cv.

Phenols content (mg/kg of tyrosol):	Mean	Max	Min	S.D.
3,4-DHPEA	1.0	1.9	0.3	0.6
p-HPEA	2.4	5.5	0.0	2.4
Vanillic acid	1.2	3.2	0.4	1.2
Vanillin	2.0	2.7	1.0	0.6
p-Coumaric acid	1.7	1.9	1.5	0.2
Hydroxytyrosyl acetate	0.4	0.7	0.0	0.3
Ferulic acid	3.0	8.4	0.6	3.2
o-Coumaric acid	0.1	0.2	0.0	0.1
3,4-DHPEA-EDA ox	0.0	0.0	0.0	0.0
3,4-DHPEA-EDA	76.3	203.2	24.8	74.2
3,4-DHPEA-EA	23.8	46.9	6.8	17.2
Tyrosyl acetate	0.6	1.0	0.3	0.3
p-HPEA-EDA ox	15.2	25.2	6.5	7.6
p-HPEA-EDA	65.3	130.1	37.8	37.6
Pinoresinol, 1-acetoxypinoresinol	58.9	73.3	52.3	9.5
Cinnamic acid	3.5	5.1	1.8	1.4
p-HPEA EA	0.8	1.5	0.0	0.7
3,4-DHPEA-EDA, -EA, -H ox	0.8	1.2	0.5	0.3
Luteolin	3.1	4.9	2.4	1.0
3,4-DHPEA-EA, -H	37.8	73.7	19.2	21.1
p-HPEA-EA, -H ox	5.8	9.4	2.8	2.4
Apigenin	1.0	1.4	0.2	0.5
Methyl-luteolin	8.2	14.5	5.1	3.7
p-HPEA-EA, -H	4.1	7.6	1.9	2.6
Total phenols	306	535	182	135

S.D.: standard deviation

Table 7. Data matrix used to build the PCA. The values (mV) represents the intensity of the dimer signals of each volatile compounds present in the heat maps for each samples.

Samples	Ethyl acetate	Ethyl propanoate	3-methyl- 1-butanol	Propanoic acid	(E,E)-2,4 -hexadienal	(E)-2-heptenal	6-methyl-5 -hepten-2-one	Ethanol	Acetic acid	Hexanal	(E)-2-hexenal	1-hexanol	1-octen-3-ol	(Z)-3-hexenyl acetate	Nonanal
C1	203.7	331.4	229.2	397.7	377.4	396.3	419.4	548.9	333.2	800.5	845.0	395.1	399.8	401.0	376.4
C2	336.1	325.9	229.2	567.7	384.0	396.3	433.4	431.6	345.6	924.1	1000.0	389.1	392.2	406.7	376.4
CR1	493.4	314.8	284.5	549.8	357.5	354.9	440.4	1000.0	388.8	951.1	871.1	353.2	376.8	344.5	339.4
CR2	434.8	602.0	221.3	447.0	350.9	343.1	454.4	852.5	444.3	1000.0	869.2	347.2	369.1	344.5	333.2
M1	527.7	314.8	245.0	676.8	364.1	354.9	447.4	681.8	499.8	595.9	801.4	353.2	384.5	344.5	345.6
M2	450.7	309.3	237.1	570.8	364.1	354.9	433.4	612.8	351.7	577.9	717.3	347.2	376.8	350.2	345.6
O1	455.9	336.9	272.7	784.9	377.4	396.3	419.4	647.5	432.0	888.0	818.5	395.1	376.8	406.7	376.4
O2	462.3	325.9	260.8	568.7	384.0	396.3	433.4	653.0	413.5	898.3	820.9	395.1	384.5	395.4	376.4
OL1	809.5	331.4	229.2	745.0	377.4	408.2	412.4	281.7	407.3	508.4	801.0	389.1	376.8	395.4	388.8
OL2	1000.0	458.4	347.8	1000.0	523.0	573.8	496.3	673.1	549.2	279.3	942.0	568.7	446.0	593.1	555.4
OL3	768.9	314.8	715.3	978.0	364.1	349.0	398.5	592.3	450.5	360.4	355.5	359.2	384.5	355.8	357.9
OL4	779.9	314.8	446.6	929.7	364.1	354.9	405.5	615.2	456.6	818.5	550.2	359.2	392.2	350.2	357.9
OL5	777.3	309.3	656.0	967.5	364.1	360.8	405.5	468.1	382.6	388.7	385.1	353.2	376.8	355.8	345.6
OL6	806.3	314.8	407.0	827.9	364.1	354.9	426.4	719.3	555.4	598.5	734.4	359.2	392.2	350.2	345.6
OL7	816.4	314.8	565.1	995.8	357.5	360.8	426.4	195.0	357.9	455.6	639.0	359.2	384.5	350.2	351.7
P1	683.9	353.5	237.1	369.4	397.2	408.2	433.4	443.4	382.6	741.3	945.1	395.1	384.5	406.7	388.8
P2	333.0	336.9	233.2	244.5	377.4	402.2	405.5	256.1	382.6	854.6	423.3	401.1	399.8	418.0	382.6

Table 8. Data matrix used to build the PCA. The values (mV) represents the intensity of the monomer signals of each volatile compounds present in the heat maps for each samples.

Samples	(E)-2-heptenal_M	(E,E)-2,4-hexadienal_M	Ethyl acetate_M	Ethyl propanoate_M	3-methyl-1-butanol_M	(Z)-3-hexenyl acetate_M	Hexanal_M	(E)-2-hexenal_M	1-hexanol_M	Acetic acid_M
C1	382.2	387.9	374.4	351.4	146.4	368.7	977.9	878.3	280.7	263.9
C2	374.9	395.4	421.7	375.6	191.8	368.7	985.7	879.1	280.7	265.2
CR1	360.2	402.8	615.4	393.8	501.0	354.5	975.7	864.2	285.6	353.6
CR2	367.5	387.9	343.8	351.4	278.4	354.5	1000.0	865.0	280.7	399.7
M1	367.5	387.9	494.9	339.3	387.6	368.7	847.7	861.1	270.9	389.2
M2	374.9	380.5	410.6	351.4	346.4	368.7	801.3	862.6	270.9	331.1
O1	382.2	387.9	863.8	454.4	443.3	375.8	986.8	873.6	290.6	827.2
O2	389.6	395.4	684.0	393.8	362.9	368.7	968.0	871.3	315.2	1000.0
OL1	382.2	402.8	532.9	351.4	177.3	375.8	835.5	876.8	334.9	476.3
OL2	470.4	462.5	918.4	563.5	459.8	489.2	681.0	1000.0	645.2	773.1
OL3	374.9	395.4	528.3	393.8	983.5	382.8	716.3	821.0	295.5	497.4
OL4	374.9	402.8	832.3	490.8	732.0	368.7	933.8	859.5	310.3	547.5
OL5	374.9	395.4	628.4	399.9	1000.0	375.8	732.9	830.5	354.6	558.0
OL6	374.9	402.8	790.5	339.3	688.7	368.7	802.4	859.5	330.0	753.3
OL7	374.9	380.5	1000.0	357.5	913.4	368.7	742.8	863.4	270.9	409.0
P1	389.6	402.8	532.0	363.5	146.4	382.8	854.3	873.6	275.8	325.9
P2	374.9	402.8	420.8	357.5	127.8	368.7	875.3	861.1	265.9	298.2

M: monomer







