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Adding extra-dimensions to hazelnuts primary metabolome fingerprinting by comprehensive twodimensional gas chromatography combined with time-of-flight mass spectrometry featuring tandem ionization: Insights on the aroma potential

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19 Abstract

20 The information potential of comprehensive two-dimensional gas chromatography combined 21 with time of flight mass spectrometry (GC×GC-TOFMS) featuring tandem hard (70 eV) and soft (12 eV) 22 electron ionization is here applied to accurately delineate high-quality hazelnuts (Corylus avellana L.) 23 primary metabolome fingerprints. The information provided by tandem signals for untargeted and 24 targeted 2D-peaks is examined and exploited with pattern recognition based on template matching 25 algorithms. EI-MS fragmentation pattern similarity, base-peak m/z values at the two examined energies (i.e., 12 and 70 eV) and response relative sensitivity are adopted to evaluate the complementary nature 26 27 of signals.

As challenging bench test, the hazelnut primary metabolome has a large chemical dimensionality that includes various chemical classes such as mono- and disaccharides, amino acids, low-molecular weight acids, and amines, further complicated by oximation/silylation to obtain volatile derivatives.

31 Tandem ionization provides notable benefits including larger relative ratio of structural informing 32 ions due to limited fragmentation at low energies (12 eV), meaningful spectral dissimilarity between 12 33 and 70 eV (direct match factor values range 222-783) and, for several analytes, enhanced relative 34 sensitivity at lower energies. The complementary information provided by tandem ionization is exploited 35 by untargeted/targeted (UT) fingerprinting on samples from different cultivars and geographical origins. 36 The responses of 138 UT-peak-regions are explored to delineate informative patterns by univariate and 37 multivariate statistics, providing insights on correlations between known precursors and (key)-aroma 38 compounds and potent odorants. Strong positive correlations between non-volatile precursors and 39 odorants are highlighted with some interesting linear trends for: 3-methylbutanal with isoleucine (R^2 40 (0.9284); 2,3-butanedione/2,3-pentanedione with monosaccharides (fructose/glucose derivatives) (R^2 0.8543 and 0.8860); 2,5-dimethylpyrazine with alanine (R^2 0.8822); and pyrroles (1H-pyrrole, 3-methyl-41 42 1H-pyrrole, and 1H-pyrrole-2-carboxaldehyde) with ornithine and alanine derivatives (R^2 0.8604). The 43 analytical work-flow provides a solid foundation for a new strategy for hazelnuts quality assessment 44 because aroma potential could be derived from precursors' chemical fingerprints.

45

46 Key-words

47 comprehensive two-dimensional gas chromatography, UT fingerprinting, template matching, time-of-

48 flight mass spectrometry featuring tandem ionization, high quality hazelnuts, aroma potential

50 **1. Introduction**

51 Since its introduction, comprehensive two-dimensional gas chromatography (GC×GC) coupled 52 with mass spectrometry (MS) has shown great potential in profiling and fingerprinting investigations [1] 53 in food [2–6]. 2D patterns of separated compounds well describe sample chemical dimensionality [7].

Although GC×GC is the technique of election to investigate food volatiles, it also enables effective studies on semi-volatiles and on non-volatiles suitably, after derivatization to improve thermal stability and volatility. Examples of research in this direction are the characterization of lipid fractions by GC×GC-MS/FID [8,9], fatty acids methyl esters (FAMEs) profiling [10–12], and primary metabolites in function of harvesting and storage practices [13,14].

59 The current study aims at developing a methodology based on GC×GC-TOF MS that combines 60 profiling and fingerprinting strategies [1] based on 2D-data patterns, to comprehensively map hazelnuts 61 key-aroma compounds precursors and relevant primary metabolites (amino acids, reducing sugars and 62 polyalcohols, organic acids etc..) from raw fruits. To deal with the great complexity of primary 63 metabolome, variable electron ionization (EI) MS is examined. The patented technology, termed tandem ionization[™] [Select eV[™] - US patent number 9,786,480], operates with variable-energy EI across single 64 analytical runs. The acquisition is therefore done by time-switching between two ionization energies so 65 66 that tandem data streams are generated and acquired iteratively. In the ion source, a high potential 67 difference accelerates electrons away from the filament while reducing their energy before their entrance 68 in the ion chamber [15]. This results in a more efficient ionization and a reduced loss of sensitivity at low-69 energy EI with enhanced intensity for structure-diagnostic ions [16].

70 Recent studies, have demonstrated that tandem ionization combining 14 and 70 eV provided 71 successful discrimination of isomeric species in complex mixtures of motor oil samples [17] and in blood 72 volatiles [18]. Additional benefits are derived from dedicated tandem data processing procedures. Freye 73 et al. [19] applied tile-based Fisher ratio analysis on fused tandem signals to detect analytes spiked at 50 74 mg/kg level in diesel samples. Cordero et al. [16] extended the combined untargeted/targeted 75 fingerprinting (UT fingerprinting) approach [20,21] to cocoa volatilome analysis. They confirmed that 76 tandem signals, when summed together, gave better performances, in terms of samples discrimination 77 and classification. In addition, analytes with a reduced fragmentation at 12 and 14 eV gave higher signal-78 to-noise ratio at lower ionization voltages also extending the dynamic range of measurements.

79

80 Challenges posed by hazelnuts primary metabolome fingerprinting are related to the large 81 chemical dimensionality [22], that includes different chemical classes as mono- and disaccharides, amino acids, low-molecular weight acids, and amines, and are furtherly complicated by oximation/silylation to
 obtain volatile derivatives. However, the information embedded in primary metabolites chemical
 signatures could be of high interest if correlated with hazelnuts hedonic qualities. Several aroma active
 compounds [23–26] have known mechanisms of formation from non-volatile precursors [25–27].

Intriguing empirical evidence on the instability, and/or evolution, of the volatile metabolome 86 87 during shelf-life and storage of hazelnuts was discussed by Cialiè Rosso et al. [28]. In that study, GC×GC-88 MS, combined with advanced fingerprinting based on template matching, highlighted informative 89 patterns of odorants strongly correlated, in raw hazelnuts, to post-harvest drying practices and storage 90 conditions within a 12-month shelf-life [28]. At the same time, the aroma potential of nuts was studied by mapping volatiles fingerprints after lab-scale roasting across the shelf-life (e.g., 12 months after 91 92 harvest). Results obtained with two origins (Turkish Ordu and Italian Nocciola Romana or Tonda Gentile 93 Romana PDO - IT/PDO/0005/0573), showed that volatile patterns were greatly impacted by storage conditions. Samples stored under modified atmosphere (99% $N_2 - 1\% O_2$) at 5°C were characterized by 94 95 higher abundance of alkyl pyrazines, Streker aldehydes, and ketones compared to samples kept in a 96 storage atmosphere richer in oxygen (78% $N_2 - 21\% O_2$) and/or at ambient temperature (18°C). Most 97 components are Maillard reaction products and/or formed by thermal degradation of reducing sugars, so 98 it was hypothesized that storage would have impacted the distribution/availability of some non-volatile 99 precursors in the fruits.

100 In this study, for the first time, GC×GC-TOF MS featuring tandem ionization is tested for its 101 effectiveness on hazelnuts primary metabolome fingerprinting. In particular, spectral 102 similarity/dissimilarity between high (70 eV) and low (12 eV) ionization energy is examined, including 103 complementary information potential, sensitivity, and dynamic range of response. The analytical strategy 104 is then validated for its informative potential and reliability in providing robust data for correlation studies 105 with potent odorants distribution.

106

107 **2. Experimental**

108 **2.1 Chemicals and reference solutions**

109 Pure standards of *n*-alkanes (from n-C9 to n-C25) for system evaluation and linear retention 110 indexes (I^{T}) determination and α -tujone for volatiles internal standardization (IS) were from Merck (Milan, 111 Italy).

112 The mixture of *n*-alkanes for the I^{T} solution was prepared in cyclohexane at a concentration of 100 113 mg/L; internal standard (IS) α -thujone solution was prepared in diethyl phthalate (Sigma Aldrich 99% of 114 purity) at a concentration of 100 mg/L.

Pure standards for identity confirmation of pyruvic acid, lactic acid, malonic acid, acetoacetic acid, phosphoric acid, succinic acid, glyceric acid, fumaric acid, malic acid, citric acid, Alanine-Ala, Asparagine-Asn, Aspartic acid-Asp, Cysteine-Cys, Glutamic acid-Glu, Glycine-Gln, Isoleucine-IIe, Leucine-Leu, Lysine-Lys, Methionine-Met, Ornithine-Orn, Phenylalanine-Phe, Proline-Pro, Serine-Ser, Threonine-Thr, Tryptophan-Trp, Tyrosine-Tyr, Valine-Val, glycerol, xylitol, mannitol, myo-inositol, fructose, glucose, saccharose, and the internal standards (IS), 4- chlorophenylalanine (quality control - QC for derivatization) and 1,4-dibromobenzene (QC for GC normalization), were from Merck (Milan, Italy).

Derivatization reagents and HPLC grade solvents: O-methylhydroxylamine hydro- chloride (MOX), (N,O-bis(trimethylsilyl)trifluoroacetamide (BSTFA) methanol, pyridine, *n*-hexane, di- chloromethane, and toluene were from Merck (Milan, Italy).

125 2.2 Hazelnut samples

Commercial grade samples of raw hazelnuts (*Corylus avellana* L.) with uniform caliber of 13 mm and harvested in 2016 were supplied by Soremartec Italia Srl (Alba-CN, Italy). They were from different geographical areas: (*a*) mono-cultivar Nocciola Romana (TR), also known as Tonda Gentile Romana, a Protected Denomination Origin (PDO) product (EU Quality registration code IT/PDO/0005/0573) here defined as *Roman*; (*b*) mono-cultivar Tonda Gentile delle Langhe (TGL) or Nocciola del Piemonte, a Protected Geographical Indication (IGP) product here defined as *Piedmont* and, (*c*) Turkish blend harvested in the Ordu (OR) region with *Tombul, Palaz* and *Çakildak* cultivars here defined as *Ordu*.

Raw hazelnuts were grinded after freezing with liquid nitrogen and stored at -18 °C before analysis. Lab-scale roasting was by a ventilated oven for 15 minutes at 160 °C on 100 g aliquots; these conditions were set in a previous study and optimized to develop major key-odorants [26] and roasting marker compounds [29].

137 **2.3 Primary metabolites extraction and derivatization**

138 *2.3.1 Defatting*

139 Micro-scale defatting was conducted in sealed glass vials (20 mL volume) on 1.00 g of ground 140 hazelnuts alignots with 5.0 mL of *n*-hexane as extraction solvent at ambient temperature and with the aid 141 of ultrasound (US - 40 KHz ± 5%). Steps of 15' were set and exhaustiveness was verified after 6-7 successive 142 extractions with fresh *n*-hexane aliquots. To avoid any contamination of the hydrophilic phase and to 143 ensure efficient derivatization in the next sample preparation step, 10 extractions were set in the final 144 protocol for samples defatting. A schematic diagram of the extraction/derivatization procedure is 145 provided as **Supplementary Figure 1 – SF1** together with experimental results on fat fraction extraction 146 yields on a representative sample (Supplementary Figure 2 – SF2). The defatted hazelnut powder was 147 then kept under a gentle stream of nitrogen to remove residual solvent and stored at -18°C before 148 extraction and derivatization.

149

2.3.2 Extraction of primary metabolites

150 An aliquot of 0.100 g of defatted hazelnuts was placed in a centrifuge glass tube with 5.0 mL of 151 H_2O/CH_3OH (98:2 v/v) mixture. Extraction was conducted at ambient temperature and with the aid of 152 ultrasound (US - 40 KHz ± 5%) for 15'. Exhaustiveness was verified after 3-10 successive extractions, 153 depending on the targeted metabolite, with fresh solvent aliquots. A schematic diagram of the extraction/derivatization procedure is provided as Supplementary Figure 1 - SF1 together with 154 155 experimental results on targeted primary metabolites extraction yields Supplementary Figure 2- SF2. 156 After extraction, centrifugation was carried out at 5,500 rpm for 10 min, the supernatant was then 157 carefully collected and filtered with Nylon HPLC filters with 20 μ m pores.

158 2.3.3 Derivatization

159 1 mL of the water-methanol extract from the first four extractions were collected together, spiked 160 with 20 μ L of 4-chlorophenylalanine solution (4 mg/mL in CH₃OH) and dried under a gentle stream of 161 nitrogen into 1.5 mL glass vials. 45 µL of MOX solution (20 mg/mL in pyridine) were added and metoximation reaction was carried out at 60 °C for 2 h. Lastly, 60 µL of (N,O-162 163 bis(trimethylsilyl)trifluoroacetamide - BSTFA were added to the reaction mixture. The sylilation reaction 164 was carried out at 60 °C for 1 h. At the end of the derivatization step, the reaction mixture was spiked with 20 μ L of 1,4-dibromobenzene (IS 1 g/L in CH₂Cl₂); an additional 75 μ L of CH₂Cl₂ were added up to a final 165 166 volume of 200 μ L.

For primary metabolites identity confirmation, 1.00 mL of primary metabolites standards mixture (listed in *section 2.1*) was submitted to the derivatization procedure and analyzed under conditions described in *section 2.5*.

170 **2.4 Headspace Solid Phase Microextraction devices and sampling conditions**

171 Automated HS-SPME sampling was performed using a MPS-2 multipurpose sampler (Gerstel, 172 Mülheim a/d Ruhr, Germany) installed on the GC×GC–TOF-MS system. SPME fibers, 173 divinylbenzene/carboxen/polydimethyl siloxane (DVB/CAR/PDMS) df50/30m - 2 cm, were from Supelco 174 (Bellefonte, PA, USA). Fibers were conditioned before their use as recommended by the manufacturer. 175 The IS (α -thujone) used for peak response normalization was pre-loaded into SPME fiber before sampling 176 by exposing the extraction device (e.g., the SPME fiber) to 5 µL of ISTD standard stock solution for 20 min 177 at 50°CC. Hazelnut samples were frozen before milling, using liquid nitrogen, to ensure homogeneous 178 particle size distribution thus stored at -80°C until analyzed. Samples were exactly weighed (1.50 g) in 179 headspace glass vials (20 mL) and submitted to headspace sampling for 40 min at 50°C.

180 **2.5 GC×GC-TOF MS featuring Tandem Ionization: instrument set-up and experimental conditions**

GC×GC analyses were performed on an Agilent 7890B GC unit coupled with a Bench TOF-Select[™] system (Markes International, Llantrisant, UK) featuring Tandem EI. Hard ionization at 70 eV was set for identity confirmation while 12 eV was applied to explore spectral complementarity. The ion source and transfer line were set at 290°C. The MS optimization option was set to operate in Tandem Ionization with a mass range between 35 and 550 *m/z*; data acquisition frequency was 50 Hz per channel; filament voltage was set at 1.60 V.

The system was equipped with a two-stage KT 2004 loop thermal modulator (Zoex Corporation, Houston, TX) cooled with liquid nitrogen controlled by Optimode[™] V.2 (SRA Instruments, Cernusco sul Naviglio, MI, Italy). The hot jet pulse time was set at 280 ms, modulation period was 5 s, and cold-jet total flow was progressively reduced with a linear function from 30% of Mass Flow Controller (MFC) at initial conditions to 5% at the end of the run.

192 2.6 GC×GC columns and settings for primary metabolites profiling

193The column set was configured as follows: ¹D DB-5 column (95% polydimethylsiloxane, 5% phenyl;19430 m × 0.25 mm d_c, 0.25 μ m d_f) coupled with a ²D OV1701 column (86% polydimethylsiloxane, 7% phenyl,1957% cyanopropyl; 2 m × 0.1 mm d_c, 0.10 μ m d_f), from J&W (Agilent, Little Falls, DE, USA). The first 0.80 m196of the ²D column, connected in series to the 1D column by a silTite μ -union (Trajan Scientific and Medical,197Ringwood, Victoria, Australia), were wrapped in the modulator slit and used as loop-capillary for cryogenic

modulation. The carrier gas was helium at a constant flow of 1.6 mL/min. The oven temperature program
was from 75°C (1 min) to 290°C (15 min) at 4°C/min.

For primary metabolites profiling, 2.0 μ L of the derivatized solution (section *2.3.3*) was analyzed under the following conditions: split/splitless injector in split mode, split ratio 1:20, injector temperature 290°C. The *n*-alkanes liquid sample solution for *I*^T determination was analyzed under the following conditions: split/splitless injector in split mode, split ratio 1:50, injector temperature 290°C, and injection volume 1 μ L.

205 2.7 GC-MS configuration and settings for volatiles profiling

GC-MS profiling of volatiles from lab-scale roasted samples was by a system configured as follows: Agilent 7890 GC unit coupled to an Agilent 5977B MS detector (Agilent, Little Falls, DE. USA) with High Efficiency Source (HES) operating in EI mode at 70 eV. The transfer line was set to 270°C, HES Tune was applied and the scan range was set to m/z 35-350 with a scanning rate of 2,500 *amu*/s.

A Solgel-Wax[™] capillary column was used (100% polyethylene glycol, 30 m × 0.25 mm dc, 0.25 μm
 df) Trajan Scientific and Medical (Ringwood, Victoria, Australia).

The analytes sampled by HS-SPME were thermally desorbed from the fiber for 5 min, directly into the GC injector, under the following conditions: split mode, split ratio 1/20, injector temperature 250°C. The carrier gas was helium, at a constant flow rate of 1.0 mL/min. The oven temperature program was 40°C (1 min) to 180°C at 3°C/min and to 240°C at 15°C/min (5 min). MS source was set at 250°C while MS quadrupole was set at 180°C.

217 **2.8** Method performance parameters: retention times and response repeatability

218 Method validation was run on a three-weeks basis and aimed at the evaluation of repeatability of 219 retention times and UT peaks response precision [30]. Retention times in both chromatographic 220 dimensions (${}^{1}t_{R}$ and ${}^{2}t_{R}$) were collected from the UT peak-regions at 70 eV of analyzed samples (3 221 cultivar/origins × 5 extraction/derivatization batches × 2 analytical replicates) for a total of 30 analytical 222 runs processed. Results are reported as relative standard deviation (RSD) in Supplementary Table 1 – ST1 with quite good retention-times stability with an average RSD of 4.78E-03 for ${}^{1}t_{R}$ and 1.80E-02 for ${}^{2}t_{R}$. 223 224 Response repeatability was calculated on all UT peak-regions at 70 eV and was based on normalized 225 responses from Piedmont samples (5 extraction/derivatization batches × 2 analytical replicates). As 226 reported in in **Supplementary Table 1 – ST1**, repeatability RSD averaged of 0.10, with a minimal of 0.03 227 for feature (35) and a maximum of 0.20 for (238), both near the end of the run.

228 **2.9 Data acquisition and 2D data processing**

GC×GC data were acquired by TOF-DS software (Markes International, Llantrisant, UK), 1D-GC-MS data were acquired and processed by Enhanced MassHunter (Agilent, Little Falls, DE, USA). GC×GC-TOF MS data were processed using GC Image GC×GC Edition, ver 2.8 (GC Image, LLC, Lincoln NE, USA). Data elaboration and results visualization were by XL-Stat (Addinsoft Inc, New York, USA) and by open source Gene-E (Broadinstitute.org).

234 3. Results and Discussion

235 In this study, hazelnut primary metabolome fingerprints are, for the first time, explored by 236 GC×GC-TOF MS featuring tandem ionization and the information potential embedded in 2D-patterns 237 evaluated in terms of both discrimination power between cultivars and origins and aroma precursors 238 distribution. In the panorama of existing studies aimed at delineating hazelnuts nutritional quality, there 239 is a lack of systematic investigations on non-volatile aroma precursors distribution studied for their 240 correlation to potent odorants formed after technological roasting. Alasalvar et al. [31] studied the 241 proximate composition, minerals, vitamins, dietary fiber, amino acids, and taste active components (free 242 amino acids, sugars, and organic acids) distribution in Tombul hazelnuts. They applied a multi-platform 243 analytical approach to profile major and essential nutrients, writing that the Tombul hazelnut would 244 "serve as a good source of vital nutrients and taste active components" [31]. The study then was extended 245 to other cultivars/origins [32,33] and also to other tree nuts [34] by also integrating the analytical profiling 246 with bio-active phytochemicals [35].

The first attempt to correlate hazelnut metabolome and its volatile composition was by Kiefl [26], who, based on available literature, reviewed the most relevant non-volatile precursors responsible of the formation of key-aroma compounds and the reaction pathways in hazelnuts after dry-roasting. However, the study mostly focused on the volatile metabolome, lacking experimental evidence of the expected strong correlations between precursors and reaction products.

252 In this study, we designed an analytical approach based on a single analytical platform, to map 253 the characteristic distribution (quali-quantitative) of informative analytes selected from those playing a 254 major role in defining hazelnuts aroma potential and sensory quality. Reactions, promoted by dry-255 roasting, include carbohydrates caramelization, consisting of dehydration and isomerization to form α -256 dicarbonyl compounds (2,3-butanedione and 2,3-pentanedione), and furanones (5-hydroxymethyfurfural 257 and 4-hydroxy-2,5-dimethyl-3(2H)-furanone). 2-/3-Methylbutanal and phenylacetaldehyde are formed by 258 Strecker degradation of amino acids in presence of α -dicarbonyls from Ile, Leu and Phe respectively [36]. 259 Moreover, α -amino carbonyl compounds, resulting from Strecker reaction, are pyrazine precursors; their 260 dimerization leads to the formation of dihydropyrazines that can either oxidize into pyrazines or react

with aldehydes to generate substituted pyrazines [37]. Additionally, reactions between Ala, Arg, Lys, Pro and Orn, and sugar degradation products (i.e., deoxyosones) forms 2-acetyl and 2-propyonilpyrroline that can be further oxidized to pyrrole derivatives [38]. Finally, thermal degradation of primary metabolites also can affect the lipid fraction, promoting autoxidation and formation of secondary products such as saturated and unsaturated aldehydes (hexanal, octanal, (*Z*)-2-octenal, (*E*)-2-octenal, (*Z*)-2-nonenal, (*Z*)-2decenal and (*E*, *E*)-2,4-nonadienal, etc.).

The analytical strategy includes multi-analyte profiling and fingerprinting by GC×GC-TOF MS after oximation/silylation of non-volatile precursors. Tandem ionization is evaluated by combining 70 and 12 eV ionization energies to improve the confidence on analytes identification and to validate the comparative capacity of 2D-fingerprints. The high energy (70eV) enables confident identification based on the similarity of characteristic fragmentation patterns; the lower energy (12eV) provides complementary information and maximizes the relative intensity of structural informative fragments [16– 18].

Experimental data is based on a selection of three relevant hazelnuts cultivar/origins (*Roman*, *Piedmont*, and *Ordu*) from harvest year 2015 analyzed at time zero (i.e. after in-field post-harvest drying). Five different subsets of samples which were independently defatted/extracted/derivatized and two analytical replicates were run for each.

The next sections include: (*a*) a discussion of the informative power of the approach and on the coverage of the non-volatile precursors, e.g., primary metabolites, of interest; (*b*) an evaluation of the complementary nature of tandem signals and their synergy in revealing compositional differences between samples; (*c*) an evaluation of existing correlations between primary metabolites and volatiles generated by lab-scale roasting with some insights on specific compounds generating key-aromas.

283 **3.1** Fingerprinting information power

284 In the development of the analytical protocol, reference studies based on GC-MS and GC×GC-MS 285 [13,14,39–41] were considered for their potential to cover the chemical dimensions [22] of interest. 286 Extraction and derivatization were therefore validated for their exhaustiveness and efficacy toward 287 aminoacids, sugars and polyalcohols and some low-molecular weight acids. Exhaustiveness was evaluated 288 by analysing each aliquot of the ten successive extractions until none of the targeted primary metabolites 289 (including all silvlated forms) were detected. Results are illustrated visually in Supplementary Figure 2 -290 SF2 as normalized responses (Normalized 2D-Peak Volume) while some considerations about 291 derivatization issues are commented and supported by dedicated references.

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В

Experimental results indicate that AA are almost 100% recovered after four extractions while for most of the acids, the exhaustive recovery is after 5 extractions. Exceptions are phosphoric acid, malic acid, lactic acid, and citric acid. According to literature, sugars and polyalcohols are major metabolites (excluding the fat components) and, for most of them, a dedicated procedure would be necessary to achieve their complete extraction. However, the first four aliquots were collected and unified before oximation/silylation to obtain more representative extracts.

300 The application of the combined untargeted/targeted (UT) fingerprinting procedure [20,42] on 301 the set of the 30 2D-patterns acquired at 70 eV (3 cultivars/origins × 5 extraction/derivatization batches 302 × 2 analytical replicates) revealed 138 UT peak-regions included in a UT feature template. They contained 303 108 targeted peaks putatively identified through their 70 eV EI-MS fragmentation pattern, by comparing 304 them to those collected in commercial and in-house databases and for which a Direct Match Factor (DMF) 305 ≥ 900 was obtained by applying the NIST Similarity match algorithm [43] on 2D-peak spectrum. The latter 306 is the average spectrum obtained from the highest modulation within the 2D-peak region. Positive 307 identifications were accepted for candidates with an experimental ${}^{1}D I^{T} \pm 15$ units tolerance. Within this 308 list of 110 known analytes, 25 were confirmed by reference standards (see section 2.1). The identities of 309 30 analytes could not be confirmed and are tagged with unique numerical identifiers [(#)] Their re-310 alignment across the 30 2D-patterns was by template matching and by applying retention times and mass spectral similarity constraints (${}^{1}t_{R}$ and ${}^{2}t_{R} \pm 15\%$ and DMF ≥ 800) with affine transformation [44,45]. The 311 312 complete list of UT peak-regions together with their average retention times (${}^{1}t_{R}$ min, ${}^{2}t_{R}$ sec), 313 experimentally determined ${}^{1}D I^{T}$ values and tabulated ones (NIST database [43]), is reported in the 314 Supplementary Table 1 – ST1.

The next section discusses the complementary nature of tandem signals acquired at 70 and 12 eV and their information power in the chemical fingerprinting of hazelnut primary metabolome.

317 **3.2** Tandem ionization: the complementary nature of tandem signals

The impact of different ionization voltages on spectral profiles and, thereby, on total ion response can be highlighted by applying datapoint features fingerprinting [46]. The approach compares the response, or the relative response, between image pairs based on a pixel-basis and promptly evidences those pattern regions in which detector responses greatly varies between the two data streams. **Figure 1** reports the comparative visualization, rendered as the *colorized fuzzy ratio*, between the 12 eV data stream (*analyzed* image) and the 70 eV signal (*reference* image) from a *Piedmont* hazelnut extract. Analyte relative response in the two detector signals is highlighted by color coding (green, red and light-grey). Regions colored in green indicate analytes for which the relative intensity (Signal-to-Noise) was higher in the *analyzed* image (12 eV) and those colored in red indicate analytes with a higher relative intensity in the *reference* image (70 eV).

328 Insert Figure 1 here

UT peak-regions corresponding to some monosaccharides (glucose and fructose derivatives), sucrose and polyalcohols (xylitol and myo-inositol), and AA, showed a higher absolute and relative response on the 12 eV signal. This suggests that the lower ionization energy would have a great impact on analyte fragmentations. This evidence has been confirmed by calculating DMF and Reverse Match Factors (RMF) values on a selection of analytes at 12 eV and 70 eV. They are listed in **Table 1** together with base peak (BP) information at the two ionization energies and SNR values.

335 The DMF values averaged 608 with minimal values for Valine 2TMS derivative – 222, Serine 3TMS 336 – 253, Valine TMS derivative – 254, and Alanine 2TMS – 317. The RMF averaged 657, slightly higher 337 because the similarity computation does not include the analyzed spectra (12 eV) fragments not present 338 in the reference at 70 eV. Smaller RMF values were for Valine 2TMS derivative – 228, Serine 3TMS – 374, 339 Alanine 2TMS – 376, and Glutaric acid 2TMS – 389. Compared to average values registered in a previous 340 study focused on the cocoa volatilome [16], for which DMF averaged 779 and RMF 787, spectral 341 dissimilarity is higher emphasizing the complementary nature of high and low ionization energies. This 342 evidence suggests that a differential impact on sensitivity has to be expected (see below).

343 An additional parameter considered in this comparative process was the base peak (BP) m/z value. 344 UT-peaks spectra were manually inspected, background subtraction and deconvolution were applied, 345 when necessary, to isolate a clean reference spectrum from which BP could be defined. Results are 346 reported in **Table 1**. For 53 of 72 2D-peaks (74%), the BP at 70 eV had lower m/z value when compared to 347 that recorded at 12 eV (shown as negative values in the BP Diff column). This result is quite interesting; it 348 indicates that structure-informing fragments are more prevalent at lower ionization, thus providing an 349 additional basis to identify analytes and, furthermore, additional specificity could be explored when the 350 two data streams are combined. To note, just 3 analytes (Glucopyranose 5TMS, Glycine 2TMS, and Serine 351 3TMS) presented a larger intensity BP at 70 eV (shown as a difference positive).

To further evaluate the complementary nature of tandem signals, relative sensitivity was considered through the ratio of the experimental SNR values registered at 12 and 70 eV data streams (data is the average value between data streams and replicates). Results indicate that for 38 of 138 *UT*peaks (27%) listed in **Supplementary Table 1 – ST1**, the lower ionization energy signal has a larger relative intensity including several targeted analytes of interest for their role as aroma precursors. They include sucrose 8TMS (SNR ratio 13), fructose 5TMS syn (SNR ratio 5), glucose 5TMS (SNR ratio 4), xylitol, 5TMS
derivative (SNR ratio 4), malic acid 3TMS (SNR ratio 4), myo-inositol 6TMS derivative (SNR ratio 3), glycerol
3TMS (SNR ratio 3), glycine 3TMS (SNR ratio 2), ornithine 4TMS derivative (SNR ratio 2), and citric acid
4TMS derivative (SNR ratio 2).

361 Figure 2 reports the spectra of four analytes showing dissimilar fragmentation patterns: glucose 362 in its cyclic form (i.e., glucopyranose 5TMS), Gly 3TMS, citric acid 4TMS, and succinic acid 2TMS. In all cases, the molecular ion was detected at both 70 and 12 eV. In the glucopyranose spectrum at lower 363 ionization energy, the molecular ion (i.e, 540 m/z) was slightly larger (1.0E-02 vs. 9.8E-04 percent 364 365 intensity). In general, spectra at 12 eV are dominated by structurally informing fragments whereas those 366 at 70 eV are dominated by the fragments of silvlating agent (i.e., TMS - 73 m/z) with low informing power. 367 In the case of glucose and citric acid derivatives, the base peak (BP) between 12 (red) and 70 (blue) eV 368 spectra is different. These results confirm the complementary nature of tandem ionization signals, a 369 characteristic that is very useful for the accurate fingerprinting of complex samples. For analytes showing 370 larger SNR values at 12 eV, the relative sensitivity of the method increases at lower energies and additional 371 benefits could be also expected on the linearity range. At lower energies the amount of ionized molecules 372 is reduced, compared to 70 eV, and the risk of detector saturation is minimized. Experimental evidence 373 on the extended linearity range of responses for volatile compounds is discussed in an application of 374 interest for the flavor and fragrance field [47].

375 Insert Figure 2 here

376 The UT feature template was applied to the 30 2D-patterns obtained at 70 eV and re-aligned to 377 the 12 eV data streams, then response information from all UT peak-regions was extracted to exploit the 378 complementary information from tandem signals in terms of fingerprinting information potential. The 379 final data matrix was 60×144 (60 2D-patterns [3 cultivar/origins × 5 extraction/derivatization batches × 2 380 ionization energies × 2 analytical replicates] × 144 UT peak-regions). Results are visualized as heat-map in 381 Figure 3. Figure 3A shows the 70 eV data stream and represents, in a color scale from blue to red, the 382 normalized UT peak-regions volume distribution; Figure 3B shows, in a color scale from green to red, the 383 12 eV UT peak-regions. Hierarchical clustering (HC) is based on Euclidean distances and was applied after 384 Z-score normalization of the data (i.e., subtract mean and divide by standard deviation).

385 Insert Figure 3 here

Primary metabolites fingerprints can independently cluster samples based on their cultivar/origin.
 The two data streams show a high consistency in their discrimination potential: both data streams closely
 cluster *Piedmont* and *Roman* samples although informative UT peaks contributing to this classification are

389 different (see rows/UT peaks HC clusters). In particular, Ordu samples are connoted by a lower relative 390 concentration of primary metabolites (confirmed by both data streams). Only a few analytes show a 391 higher relative abundance: Gly and tartaric acid within the known compounds set and (1199) and (1153) 392 within the unknown compounds set. On the other hand, the discrimination between Piedmont and Roman 393 samples is mainly driven by: (a) sugars that are more abundant in the Piedmont cultivar (e.g. glucose, 394 galactose, maltose, and fructose (considering all derivatives); (b) AA like Trp, Orn, and Tyr more abundant 395 in *Piedmont* and Leu, Ile, Met, Val, Phe, Pro, and pyroglutamic acid with a more intense response in the 396 Roman samples; (c) acids such as lactic acid (monomer 2TMS and dimer 2TMS), glutaric acid, galacturonic 397 acid, fumaric acid, tartaric acid, and oxalic acid which are more abundant in *Roman* samples. Univariate 398 statistics, in box-plot diagrams, for a selection of UT peaks are reported in the Supplementary Figure 3 – 399 SF3.

Tandem signals, when independently processed, enable cross validation of fingerprinting results. For some analytes, linear regression analysis between absolute responses recorded at 12 eV (dependent variable) and at 70 eV (independent variable) give good results ($R2 \ge 0.90$) as shown in **Figure 4** for some aroma precursors: glucose, Orn, Leu, and Ile. The coherent trend of analytes response validate the differential distribution of markers within sample set, while the different response factors between ionization channels, described by regression functions with slopes \neq 1, confirms their complementary nature.

407 The next section deals with the correlation between primary metabolites signatures and hazelnuts 408 volatile metabolome, in particular with key-aroma compounds and marker analytes distribution after lab-409 scale roasting.

- 410 Insert Figure 4 here
- 411

412 **3.3** Hazelnuts aroma potential and correlation with primary metabolites signatures

Samples were submitted to lab-scale roasting, designed in a previous study, to evaluate the aroma quality of roasted hazelnuts. The protocol includes a dry-roasting in a ventilated oven at 160°C for 15 minutes [28]. Roasted samples then were submitted to HS-SPME sampling to extract volatiles and GC-MS profiling to monitor quali-quantitative changes in volatile markers including potent odorants [26,29,48– 50].

The list of targeted volatiles is provided in the **Supplementary Table 2 – ST2**, together with the experimental I^{T} , odor quality and odor threshold for markers. They include key-aroma compounds validated by sensomics [23]: 2- and 3-methylbutanal, with *malty* notes; 2,3-butanedione and 2,3421 pentanedione, with buttery odor quality; 3-methyl-4-heptanone, 5-methyl-(Z)-2-hepten-4-one and 5-422 methyl-(E)-2-hepten-4-one (i.e., filbertone), with characteristic nutty and fruity odors; the earthy 423 pyrazines 2,3,5-trimethylpyrazine and 3,5-dimethyl-2-ethylpyrazine; acetic acid, responsible of the sour 424 note; phenylacetaldehyde, with honey and flowery odors; 2-methylbutyric acid, with a sweaty note; and 425 4-hydroxy-2,5-dimethyl-3(2H)-furanone (i.e., furaneol), with the characteristic sweet and caramel-like 426 note. Within the volatile fraction of roasted samples, several other compounds were targeted. They 427 belong to the group of roasting indicators, i.e., compounds formed during thermal processing and for 428 which a meaningful increase was registered during lab-scale roasting [29]. They include: carbonyl 429 derivatives (2-methylpropanal, 2-butenal and 3-hydroxy-2-butanone), alkyl pyrazines (pyrazine, 2-430 methylpyrazine, 2,5-dimethylpyrazine, 2,6-dimethylpyrazine, ethylpyrazine, 2,3-dimethylpyrazine, 2-431 ethyl-6-methylpyrazine, 2-ethyl-5-methylpyrazine, 2,3,5-trimethylpyrazine and 3-ethyl-2,5-432 dimethylpyrazine), furanones (furfural, 5-methylfurfural, dihydro-2(3H)-furanone and furfuryl alcohol), 433 and pyrroles (1(H)-pyrrole, 1-methyl-1(H)pyrrole and 1(H)-pyrrole-2-carboxaldehyde). The present 434 discussion does not include secondary products of lipid oxidation that contribute to some odor qualities 435 and inform about shelf-life stability, although a dedicated procedure to study the fat fraction has been 436 developed.

The squared Pearson correlation coefficient (*r*) was calculated to evaluate the existence of positive correlations between primary metabolites and volatiles. The *p*-values computed for each coefficient afford testing the null hypothesis that *r* values are not significantly different from 0 and informs about the relevance of the correlation.

441 The data matrix included the normalized responses (i.e. normalized 2D peak-regions from GC×GC 442 data and normalized chromatographic areas for 1D-GC data) for targeted primary metabolites and all 443 volatiles respectively (see ST1 and ST2). The resulting matrix was 30×162 features (30 2D-patterns [3 444 cultivar/origins × 5 extractions/derivatization batches × 1 ionization energy 70 eV × 2 analytical replicates] 445 × 108 targeted peak-regions + 54 volatiles). Before processing, data were scaled by Pareto scaling [51] in 446 order to reduce the relevance of large values while keeping data structure almost intact. The resulting 447 Pearson correlation matrix, showing r values, is illustrated as heat-map in Figure 5. The color scale, from 448 blue to red, has been adapted to emphasize (red color) r values ≥ 0.8 , arbitrarily considered meaningful 449 in this specific data-set for notable correlations. HC is based on "one minus Pearson correlation" value as 450 metrics.

451 Insert Figure 5 here

452 Results show: (a) high r values (accompanied by meaningful p-values – α =0.05) within primary 453 metabolites (Figure 5 – box 1) confirming fingerprinting results where Piedmont and Roman samples 454 showed a higher relative abundance for most of the non-volatile precursors and metabolites compared 455 to Ordu origin (see Fig. 3A and SF3); (b) high correlation values ($r \ge 0.8$ - Figure 5 - box 2) within volatiles 456 including 3-methyl-4-heptanone, 2,3-pentanedione, (E)-3-penten-2-one, 3-hydroxy-2-butanone, 457 benzenemethanol, furaneol, furfuryl alcohol, 3-methylbutanal, 1H-pyrrole, 2,3-dimethylpyrazine, ethylpyrazine, methylpyrazine, and trimethyl pyrazine; and (c) between primary metabolites (and/or non-458 459 volatile precursors) and volatiles with a high informing power in terms of aroma quality. Further details 460 on these last correlations are discussed below.

461 These correlations were tested for their significance (p-value) and linearity; the coefficients of 462 determination (R^2) of the regression were calculated, considering the precursor(s) as independent variable 463 (x) and key-volatiles as dependent variable (y). Figure 6A reports the regression function between 3-464 methylbutanal and Leu (R^2 0.9577), Fig. 6B between 3-methylbutanal and Ile (R^2 0.9284), Fig. 6C-D 465 between 2,3-butanedione and 2,3-pentanedione and the sum of fructose (Fructose 5TMS syn- and anti-466 forms) and glucose (Glucopyranose 5TMS and Glucose 5TMS) derivatives (R² 0.8543 and 0.8860), Fig. 6E 467 between 2,5-dimethylpyrazine and Ala (R^2 0.8822) and pyrroles (1H-pyrrole, 3-methyl-1H-pyrrole and 1H-468 pyrrole-2-carboxaldehyde) and the sum of Orn and Ala derivatives (R^2 0.8604).

469

Insert Figure 6 here

These results are interesting, and evidence a solid foundation for adoption of the proposed approach for a comprehensive primary metabolome fingerprinting as an informative tool to characterize aroma potential of hazelnuts. However, the approach needs validation through a more extensive sampling that includes further variables influencing primary metabolites distribution in hazelnuts: cultivar and origins should be accompanied by a selection of multiple harvest years, shelf-life, and storage conditions that also are relevant because of their known impact on the aroma potential [28].

476

477 4. Conclusions

The present study is focused on variable EI energy TOF MS adopted to extend the analytical dimensions of GC×GC for hazelnut primary metabolites profiling. By combining standard 70 eV and 12 eV, tandem data streams are generated and spectra show a complementary nature. This characteristic can be exploited in identification and structural elucidation studies. At low energy, structure informing fragments prevail while overall fragmentation is reduced, achieving, for some analytes, a higher relative sensitivity that results in a wider dynamic range of the method. Tandem data responses from *UT*-peaks in 484 complex chemical signatures are strongly correlated but exhibit different response factors that open485 interesting perspectives for quantitative studies.

Response data recorded from primary metabolites fingerprints in different cultivars/origins show good correlation to volatiles formed during dry-roasting and validate empirical observations from previous experiments [28]. Linear regressions with meaningful *R*² give solid foundations to the causal relationship between metabolome and sensobolome signatures. The proposed approach, once validated over a wider sampling design, would introduce a new concept in hazelnuts quality assessment while opening new perspectives for breeding studies and shelf-life quality evaluation.

492

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495

496 **Compliance with ethical standards Notes**

- 497 Prof. Stephen E. Reichenbach has a financial interest in GC Image, LLC.
- 498 Melanie Charron, Federica Manini, Roberto Menta and Mauro Fontana are employees of Soremartec
- 499 Italia Srl, Alba (CN, Italy).

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676 **Figure Captions** 677 Figure 1: comparative visualization, rendered as the colorized fuzzy ratio, between the 12 eV data stream 678 (analyzed image) and the 70 eV signal (reference image) from a Piedmont hazelnut extract. Green 679 highlighted areas indicate peak-regions where the differential response between channels was higher on 680 12 eV ionization energy. 681 682 Figure 2: fragmentation patterns recorded for a selection of analytes at 12 eV (red trace) and 70 eV (blue 683 trace) ionization energy. 684 685 Figure 3: heat-map visualization of the UT peak-regions normalized response from 70 eV (Fig. 3A) and 12 686 eV (Fig. 3B) data streams. Analytical replicates were averaged resulting in a (15+15) × 138 data matrix. 687 Hierarchical clustering (HC) is based Euclidean distances - Z-score normalization of the data (i.e., subtract 688 median and divide by standard deviation). 689 690 Figure 4: regression analysis between absolute responses recorded at 12 eV (dependent variable) and at 691 70 eV (independent variable) for aroma precursors: glucose, Orn, Leu and Ile. 692 693 Figure 5: heat-map illustrating Pearson correlation matrix results referred as r values. Color scale, from 694 blue-to-red, is adapted to emphasize (red color) r values \geq 0.8; HC is based on "one minus Pearson" 695 correlation and green squares indicates analytes pairs commented in the text. 696 697 Figure 6: regression functions between 3-methylbutanal and Leu (R^2 0.9577), 3-methylbutanal and IIe (R^2 698 0.9284), 2,3-butanedione and 2,3-pentanedione and the sum of fructose (Fructose 5TMS syn- and anti-699 forms) and glucose (Glucopyranose 5TMS and Glucose 5TMS) derivatives (R² 0.8543 and 0.8860), 2,5-700 dimethylpyrazine and Ala (R² 0.8822) and pyrroles (1H-pyrrole, 3-methyl-1H-pyrrole and 1H-pyrrole-2-701 carboxaldehyde) and the sum of Orn and Ala derivatives (R^2 0.8604). Graph legend (—) linear regression 702 function, (---) confidence interval mean 95%, (---) confidence interval observ. 95% 703 704

705 Table Captions:

- **Table 1:** selected targeted *UT* peak-regions corresponding to informative primary metabolites listed
- together with their average retention times (${}^{1}t_{R}$ min, ${}^{2}t_{R}$ sec), experimental and tabulated (NIST database [21]) f,
- base peak (BP) m/z values recorded from the 2D-apex spectrum and their difference (BP diff), direct match factor
- 709 (DMF) and reverse match factors (RMF) values calculated between 12 eV vs. 70 eV and SNR value between
- 710 12 and 70 eV channels (values in bold highlight analytes where SNR at 12 eV was higher compared to 70
- 711 eV).