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1 ***Artemisia umbelliformis* Lam. and génépi liqueur: volatile profile as diagnostic marker for**
2 **geographic origin and to predict liqueur safety**

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14

15 **ABSTRACT**

16 *A. umbelliformis*, commonly known as "white g n pi", is characterized by a volatile fraction rich in
17 α - and β - thujone, two monoterpenoids; under EU regulations these are limited to 35 mg/L in
18 *Artemisia*-based beverages , because of their recognized activity on human central nervous system.
19 This study reports the results of an investigation to define the geographical origin and thujone
20 content of individual plants of *A. umbelliformis* from different geographical sites, cultivated
21 experimentally at a single site, and to predict the thujone content in the resulting liqueurs, through
22 their volatile fraction. Headspace Solid Phase Microextraction (HS-SPME) combined with Gas
23 Chromatography-Mass Spectrometry (GC-MS) and non-separative HS-SPME-MS were used as
24 analytical platforms to create a database suitable for chemometric description and prediction
25 through Linear Discriminant Analysis (LDA). HS-SPME-MS was applied to shorten analysis time.
26 With both approaches, a diagnostic prediction of: i) plant geographical origin, and ii) thujone
27 content of plant-related liqueurs could be made.

28

29 Keywords: *Artemisia umbelliformis* Lam., g n pi liqueur, food safety

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33 INTRODUCTION

34 *Artemisia* L., with its 500 specific and sub-specific *taxa*, is the largest genus in the tribe *Anthemidae*
35 of the *Asteraceae* family^{1,2}. Its *taxa* grow worldwide at different latitudes, altitudes, and
36 environments, and have traditionally been used for their biological activity¹, even before recent
37 pharmacological developments³⁻⁵.

38 *Artemisia* species were used historically to prepare infusions, wines, and liqueurs⁶. The best-known
39 species is *A. absinthium*, traditionally known as wormwood and in widespread use since Roman
40 times as the base for aromatic wines and liqueurs^{6,7}. Wormwood has been studied in depth because
41 of its diffusion and ready accessibility⁶, but several other *Artemisia* species are also used in
42 beverages. A group of herbs used for liqueur production, traditionally known as "génépi", consists
43 of five rare species growing at high altitudes in the alpine area: *A. eriantha* Ten., *A. genipi* Weber,
44 *A. glacialis* L., *A. nivalis* Br.-Bl. and *A. umbelliformis* Lam.. Génépi species traditionally have less
45 economic importance than wormwood, being almost exclusively used within their production and
46 harvesting areas; however, over the last 40 years, liqueur production have markedly increased, not
47 least thanks to a number of small and medium-sized producers⁶. Génépi species have historically
48 been used in folk and traditional medicine because of their bioactivity⁶. They were known and used
49 as thermogenic agents against the common cold , in infusions against fever, and in aromatic wines
50 and liqueurs, to stimulate the appetite and the digestion,⁸⁻¹⁰ respectively. The extensive ethno-
51 pharmacological knowledge, documented from the second half of the eighteenth century, contrasts
52 with the paucity of scientific studies of génépi species⁶.

53 Liqueur production on a larger scale is not possible if the raw material is limited to the wild plant
54 harvest. Génépi cultivation, which is restricted by the plants' pedoclimatic needs, facilitates
55 harvesting operations and protects wild génépi species⁶. The five génépi species have a common
56 environmental distribution, growing at around 1500-2000 m a.s.l.¹¹. The altitude requirement is due
57 to specific soils in which these species face neither competition from weeds nor fungal pathogens,
58 which become predominant at lower altitudes^{6,12}.

59 *A. umbelliformis*, commonly known as "white génépi", is the génépi species that has been cultivated
60 more successfully, above 1600 m a.s.l.^{6,11,12}; this species has therefore been selected for wider-scale
61 cultivation. Independently of its domestication and production, the supply chain of *A. umbelliformis*
62 faces two issues linked to the food industry: product adulteration and product safety. The need for
63 an authentication process emerges, since it is one of the few Italian infusion liqueurs whose
64 geographic limits are defined in the EU by Reg. EC 110/2008¹³. This regulation introduced three
65 categories for génépi, distinguishing the most important production areas situated in North-West
66 Italy and France: "Genepi della Valle d'Aosta (Italy)", "Genepi del Piemonte (Italy)", and "Génépi
67 des Alpes/Genepi delle Alpi (France, Italy)".

68 Thujones are natural monoterpenoids widely present in *Artemisia* species. α - and β - thujone are the
69 most abundant volatiles in génépi species¹⁴⁻¹⁶, and are characterized by recognized activity on the
70 human central nervous system^{11,17,18}. For this reason, the maximum content of thujone in *Artemisia*-
71 based beverages is limited to 35 mg/L in the EU (Reg. EC 110/2008^{6,13,18}). *A. umbelliformis* is the
72 génépi species with the highest content of thujones¹⁹. The variability of the plant's chemical
73 composition (i.e. the variability of thujone content) is a crucial point for génépi liqueur producers.
74 To overcome this problem, some agronomical selection has been done to create thujone-free
75 chemotypes^{11,18}. However, chemical comparisons between thujone-free and thujone-containing
76 chemotypes have found a marked difference in the content of both aromatic and bitter compounds,
77 responsible for the typical sensory characteristics of the liqueur¹⁸.

78 Chemical analysis combined with chemometrics can be useful to deal with the problems emerging
79 from large-scale production²⁰⁻²⁵. In the present study, Linear Discriminant Analysis (LDA)²⁶ was
80 adopted for its ability to perform class discrimination, instead of Principal Component Analysis
81 (PCA), which is widely used to describe large datasets. LDA is known to elucidate the underlying
82 data structure, a key point when creating a prediction model^{24,27}. Some PCA elaborations were,
83 however, carried out and a comparison of the results to those of LDA confirm that the latter
84 technique has a better prediction capacity (data not shown).

85 This research studied *A. umbelliformis* of different geographical origins, cultivated experimentally
86 at a single site. Headspace sampling with Solid Phase Microextraction, combined with gas
87 chromatography - mass spectrometry (HS-SPME-GC-MS) or directly with mass spectrometry (HS-
88 SPME-MS), was used as an analytical platform to create the first database suitable for chemometric
89 description and prediction. Linear Discriminant Analysis (LDA) was here adopted to define and
90 predict geographic origin and thujone content of plants and liqueurs
91 Multiple Headspace- Solid Phase Microextraction (MHS-SPME)^{26,28,29} was applied for thujone
92 content determination.

93

94 **MATERIAL AND METHODS**

95 **Chemicals.** Thujones standard mixture (mixture of α -thujone and β -thujone, purity = 99.9%) and all
96 other reference compounds were from Sigma-Aldrich (St. Louis, MO). Sabinol was kindly supplied
97 by Robertet SA (Grasse, France), and sabinyl ester homologous series were synthesized in the
98 authors' laboratory¹⁸. HPLC and analytical grade solvents were from Carlo Erba Reagenti, Rodano,
99 Italy.

100 **Plant material and experimental site preparation.** Growing fields of *A. umbelliformis* were
101 monitored in spring and summer at various alpine sites in Piedmont in North-West Italy (**Table 1**).
102 The sites are located in the South-West Alps within the area indicated for the liqueur "Genepi del
103 Piemonte"¹³. At the end of the blossoming stage, four *A. umbelliformis* geographic origins were
104 selected (**Table 1**). Plant growing on the verge of the raised beds were excluded, to avoid the edge
105 effect. Mother plants were chosen based on the phenological profile (data not shown) and then
106 carefully harvested to preserve inflorescences and seeds.

107 Plants were dried at 30°C in a dry and aerated room at the Experimental Centre of DISAFA
108 (44°53'11.67"N; 7°41'7.00"E - 231 m a.s.l.) in Tetti Frati, Carmagnola (TO), Italy. The seeds were
109 cleaned to remove extraneous parts, under fan ventilation using a series of sieves of different
110 meshes, by Semina S.r.l. (La Loggia (TO), Italy), and then stored at 6°C in a seed room. The seeds

111 of the four different geographic groups were sown the following summer in trays of a specific peat-
112 based horticultural medium (Neuhaus Huminsubstrat N17; Klasmann-Deilmann® GmbH, Geeste,
113 Niedersachsen, Germany). The trays were placed in an air-conditioned greenhouse providing the
114 necessary supply of water until the seeds germinated. After cotyledon expansion, *A. umbelliformis*
115 was transplanted into 104-cell plastic trays (0.522m × 0.322m × 0.042m; 0.034m upper diameter
116 cells), filled with the above peat medium. Trays were placed in an air-conditioned plastic
117 greenhouse and seedlings were irrigated overhead as required. After secondary leaf development,
118 trays were placed under a plastic film tunnel covered with black shading systems (50%-shade
119 cloth). Before transplanting the *A. umbelliformis* seedlings into the soil, the shade cloth was
120 removed to harden the plants and acclimatize them to outdoor environmental conditions. The count
121 of dead plants was performed weekly (data not shown). Plants were transplanted in the autumn at
122 the experimental site of Pragelato (TO), Italy (45°00'36.41"N; 6°56'14.56"E - 1,500 m a.s.l.). The
123 site is characterized by a soil with a moderate presence of skeleton, absence of slope, and optimal
124 sun exposition. After minimum tillage, a white/black (upper/lower side) plastic mulching film
125 provided with holes arranged in a quincunx formation (0.05 m hole diameter, 0.20 m distance
126 between holes) was used to set 6 mulched raised beds subdivided into 12 plots (4 geographic groups
127 (Gran Paradiso, GP; Val Chisone; Elva; Valle Gesso, VG) ×3 blocks) for a total of ca. 1,350 *A.*
128 *umbelliformis* plants.

129 **Production, phenological and morphological parameters.** The site was inspected to check *A.*
130 *umbelliformis* stand establishment with a ≈ 30 days frequency. The number of dead plants was
131 monitored and recorded 30 days after transplanting, the following summer and autumn (1st year),
132 and again the subsequent summer (2nd year). The percentage of dead plants was calculated and used
133 as an indicator of stand establishment and the plants' adaptation to the site.

134 Phenological and morphological parameters were measured in the 1st summer and autumn and in
135 the 2nd summer after transplanting, on 12 plants per geographic origin and per block, randomly
136 chosen at the beginning of the experiment. Inflorescence parameters were only measured during the

137 2nd summer of experimentation. The phenological parameters measured on the canopy were: canopy
138 diameter, canopy height (**Figure 1A**); on the inflorescences: inflorescence height, inflorescence
139 number (**Figure 1B**). The morphological parameters measured on the canopy leaves were: number
140 of segments, leaf length, leaf segment length and width (**Figure 1C**); on the inflorescence leaves:
141 cauline leaf length, cauline leaf petiole length (**Figure 1D**); on the inflorescences: footstalk length,
142 number of glomerules (at first, second and third internode), number of plant glomerules, number of
143 head glomerules, total number of glomerules (**Figure 1E**).

144 Harvesting took place in the 2nd summer after transplanting, during the blossoming stage. Fresh
145 weight production per plant was measured, and weight after drying at 30°C until constant weight,
146 dry matter percentage was calculated as in the European Pharmacopoeia³⁰.

147 **Headspace Solid Phase Microextraction (HS-SPME) sampling and analysis conditions:**
148 **qualitative profiling of plants.**

149 **Separative analysis.** The 104 selected plants surviving at the end of the 2nd summer were used to
150 analyze the volatile fraction. Portions of 25 mg were sampled, as the minimum plant weight
151 required to have both an adequate informative profile and good repeatability with headspace solid-
152 phase microextraction (HS-SPME).

153 Each plant was sampled in a 20 mL headspace vial (2 replicates of 25 mg) using a 2 cm
154 DVB/CAR/PDMS fiber (Supelco, Bellefonte, PA, USA). Sampling was carried out with a MPS2
155 automatic sampling system (Gerstel, Mülheim a/d Ruhr, Germany) at T=50 °C (15 min of pre-
156 equilibrium, 15 min of sampling). The fiber was automatically transferred to the GC injector of a
157 7890 GC coupled to a 5975C MS (Agilent, Little Falls, DE, USA) from where the sampled analytes
158 were recovered directly by thermal desorption and transferred to the GC column for analysis.

159 *GC-MS conditions.* Inlet T=250°C, split injection (5 min, 1/10 split ratio); Helium was used as
160 carrier gas at a flow rate of 1 mL/min. Column: MEGA5MS (30 m × 0.25 mm i.d. × 0.25 µm;
161 MEGA, Legnano, MI, Italy). Temperature program: 50 °C (1 min)-3 °C/min-250 °C (5 min). MS
162 operated in EI mode at 70 eV with a mass range from 35 to 350 amu in full scan mode.

163 Data were processed with Agilent MSD ChemStation D.03.00.611 (Agilent Technologies).
164 Components were identified by comparing their linear retention indices (I^T s) (calculated *versus* a
165 C₉-C₂₅ hydrocarbon mixture) and their mass spectra to those of authentic standards.

166 The HS-SPME-GC-MS method repeatability was tested on ten samples in five replicates for the 30
167 selected compounds, reaching an overall value of relative standard deviation (RSD%) of 9.3%. As
168 reported in **Supplementary Table 1**, the minimum RSD% was detected with sabinyl valerianate
169 (2.5%) and the maximum RSD% with sabinene (29.3%).

170 **Non-separative analysis.** The same plant samples analyzed by HS-SPME-GC-MS were also
171 analyzed without performing GC separation. Sampling was carried out on 2 replicates of 150 mg of
172 each plant weighted in a 20 mL headspace vial using a 2 cm DVB/CAR/PDMS fiber (Supelco,
173 Bellefonte, PA, USA) at 100 °C for 5 min. Sampling and injection were managed by a Gerstel
174 MPS2 automatic system (Gerstel, Mülheim a/d Ruhr, Germany). The fiber was automatically
175 transferred to the GC injector of a 7890 GC coupled to a 5975C MS (Agilent, Little Falls, DE,
176 USA) and the sampled analytes thermally desorbed directly into the empty tubing for MS analysis.

177 *MS analysis conditions.* Inlet T:250°C, split injection (5 min, 1/20 split ratio); carrier gas: helium,
178 flow rate: 1 mL/min. MS transfer column: deactivated fused silica empty tubing (8 m × 0.25 mm
179 i.d.; MEGA, Legnano, MI, Italy). Oven temperature: 250 °C (5 min). MS operated in EI mode at 70
180 eV with a mass range from 35 to 250 amu in full scan mode.

181 MS data were processed with Agilent MSD ChemStation D.03.00.611 (Agilent Technologies) and
182 Pirouette 4.0.2 (Infometrix Inc., Bothell, WA, USA).

183 The HS-SPME-MS method repeatability was tested on ten samples in five replicates for the 30 most
184 abundant fragment ions, reaching an overall relative standard deviation (RSD%) of 1.8%. As
185 reported in **Supplementary Table 2**, the minimum RSD% was found for $m/z=43$ (0.6%) and the
186 maximum for $m/z=107$ (2.5%).

187 **Multiple headspace solid phase microextraction (MHS-SPME-GC-MS): quantitation of**
188 **thujones in plants and liqueurs.** Thirty six samples of *A. umbelliformis* among the 104 samples
189 employed in the previous steps were selected to produce experimental liqueurs. For each plant, one
190 individual liqueur was prepared, following the guidelines for liqueur preparation given in Annex II
191 (32) of EC Regulation No 110/2008¹³. MHS-SPME was then applied to the liqueurs (three
192 replicates of 10 µL for each sample introduced in a 20 mL headspace vial, three HS-SPME
193 samplings for each replicate).

194 Beside liqueur analysis, each plant sample was also analyzed by MHS-SPME-GC-MS (three
195 replicates of 3 mg for each sample weighted in a 20 mL headspace vial, three HS-SPME samplings
196 for each replicate).

197 Quantitative calibration was carried out by MHS-SPME using seven concentration levels of α - and
198 β -thujone standard mixture (92.6% α -thujone, 7.4% β -thujone) between 0.1 g/L and 10 g/L in
199 cyclohexane (two replicates of 10 µL for each level, three HS-SPME samplings for each
200 replicate)³¹.

201 *Sampling conditions.* MHS-SPME analyses were carried out using a 2 cm
202 divinylbenzene/carboxen/polydimethylsiloxane (DVB/CAR/PDMS) fiber (Supelco, Bellefonte, PA,
203 USA) at 60 °C for 30 min. Sampling and injection were managed by a Gerstel Multipurpose
204 Sampler 2 (MPS2) (Gerstel, Mülheim a/d Ruhr, Germany). The fiber was automatically transferred
205 to a 7890 GC unit coupled to a 5975C MS to analyze the sampled analytes (Agilent, Little Falls,
206 DE, USA). During MHS-SPME sampling, the sample was not heated between sampling, but placed
207 back into the sample tray after each sampling.

208 *GC-MS conditions.* Inlet T:250°C, split injection (5 min, 1/20 split ratio); carrier gas: helium, flow
209 rate: 1 mL/min. Column: MEGA-SE52 (30 m × 0.25 mm i.d. × 0.25 µm; Mega, Legnano, MI,
210 Italy). Temperature program: 50°C-3°C/min-105°C-20°C/min-250°C (1 min). MS operated in EI
211 mode at 70 eV with a mass range from 35 to 350 amu in full scan mode. Data were processed with
212 Agilent MSD ChemStation D.03.00.611 (Agilent Technologies).

213 **Statistical data treatment.** Preliminary descriptive, significance analyses of variance (ANOVA)
214 and linear regression attempts were run on MHS-SPME-GC-MS data. Linear Discriminant Analysis
215 (LDA)^{27,32} was applied to correlate MHS-SPME-GC-MS data with HS-SPME-GC-MS data and
216 HS-SPME-MS data. Statistical analyses were performed by using SPSS 15.0 (IBM Corporation) or
217 R³³; RStudio³⁴ was used to write a LDA cycling script. Other R packages used were: caret³⁵,
218 MASS³⁶ and e1071³⁷.

219 LDA describes and predicts multivariate data matrices in which samples are described by one
220 categorical variable and n continuous variables. The LDA model calculates Discriminant Functions
221 (DFs) which are used first to describe the group distribution in the analyzed dataset and then to
222 predict the groups of samples. DFs are calculated on the basis of the relationships between the
223 continuous variables describing the dataset and the categorical variables defining the group of each
224 sample. DFs can be considered similar to the Principal Components in PCA, but they differ for
225 some characteristics, such as their meaning (they maximize distances between the groups of the
226 dataset instead of maximizing the dataset explained variance), and for their number (PC number =
227 variable number; DF number = group number *minus* one)²⁷.

228 LDA was carried out on four different matrices based on different data: i) plant geographic origin as
229 categorical variable and plant morphological and phenological data as continuous variables; ii) plant
230 geographic origin as categorical variable and plant HS-SPME-GC-MS compound abundances as
231 continuous variables; iii) liqueur MHS-SPME-GC-MS thujone level as categorical variable and
232 plant HS-SPME-GC-MS compound abundances as continuous variables; iv) liqueur MHS-SPME-
233 GC-MS thujones level as categorical variable and plant HS-SPME-MS fragment ions abundances as
234 continuous variables.

235

236 **RESULTS AND DISCUSSION**

237 The study comprised three main steps: i) to describe statistically-significant differences relating to
238 the geographical distribution of cultivated *A. umbelliformis* plants; ii) to develop and apply

239 analytical methods to rapidly quantify thujone content of *A. umbelliformis*-based liqueur (génépi);
240 iii) to create multivariate statistical models to predict the above two points.

241 The morphological/phenological dataset and the chemical dataset were processed with LDA .

242 **Origin discrimination through phenological and morphological data.** At the end of the 2nd year
243 of field tests, 61.8% of the *A. umbelliformis* plants survived without showing significant differences
244 between plants from different geographical origins (data not shown). Morphological data collected
245 in the 2nd year included both vegetative and reproductive parts of the 104 plant samples (**Figure 1,**
246 **Supplementary Table 3**).

247 ANOVA was carried out, when possible ($p < 0.05$), together with the Tukey Honestly Significant
248 Difference (HSD) test. The measurements related to the length of footstalks and to the number of
249 glomerules were found to be the most significant parameters in differentiating plant individuals
250 originating from different geographical areas (**Supplementary Table 4**).

251 LDA was used to describe the morphological differentiation of the plant groups and was carried out
252 in two steps: cross validation (CV) and prediction. The dataset was divided into two subsets.
253 Approximately 60% of the samples were used to create the prediction model based on three
254 Discriminant Functions (DF). Subsequently, the residual 40% of the samples were predicted by the
255 three Discriminant Functions, in order to calculate the predictive power of the LDA model.

256 LDA was carried out considering phenological and morphological parameters, and the three DFs
257 explain the total variance (DF1 = 51.0%; DF2 = 38.6%; DF3 = 10.4%). The scatter plot reported in
258 **Figure 2** does not show a clear distribution of the samples in function of their geographical origin. .

259 However Elva and Valle Gesso are separated along DF2 DF1 and DF2 showed higher correlations
260 with the variables related to the length of the footstalk and the number of glomerules. The variables
261 related to the canopy and the leaves showed a higher correlation with DF3, which, however, did not
262 provide significant results (**Supplementary Table 5**).

263 Discriminant Functions can be used not only to describe the phenological and morphological
264 differences, but also to predict the origin of the plant samples.

265 LDA usually reports the success of a prediction model with a confusion matrix that describes the
266 recognition success of the group prediction performed on the basis of DFs. Cross validation and
267 prediction results report the percentage of the positive prediction divided by group.

268 **Table 2** shows that the LDA model recognizes samples' geographical origin in the range of 66.7-
269 77.8% in cross validation and 29.6%-51.9% in prediction. This result shows that the use of LDA
270 with morphological and phenological data does not give satisfactory results.

271 **Differentiation of plant origin by HS-SPME-GC-MS. Figure 3** reports the HS-SPME-GC-MS
272 profile of an *A. umbelliformis* sample with thirty identified compounds used to create the
273 multivariate data matrix. Similarly to the morphological data, LDA was applied to find differences
274 between plants with different geographic origins.

275 Each of the three Discriminant Functions calculated was significant, unlike the morphological
276 dataset. The scatter plot of the plant samples presents a distribution similar to the one obtained with
277 the morphological data along DF1 (**Figure 4**). Groups are better separated, especially Elva (EL) and
278 Valle Gesso (VG), while Gran Paradiso (GP) and Val Chisone still significantly overlap. DF3 helps
279 to differentiate GP from VC (**Supplementary Table 5**).

280 The improvement with HS-SPME-GC-MS data was considerable, especially in terms of prediction
281 rates obtained with the LDA model. As reported in the related confusion matrix (**Table 3**),
282 classification rates were calculated to be above 90% (mean value) both in validation (min=88.9%;
283 max=100.0%; mean=93.7%) and in prediction (min=82.6%; max=96.3%; mean=90.3%).

284 **Quantitation of thujones in plants and liqueurs by MHS-SPME-GC-MS.** MHS-SPME-GC-MS
285 analysis was used to quantify α - and β -thujone in plants and liqueurs and, then, to find a correlation
286 between the total thujone amount in plants and liqueurs. Thirty-six plant and liqueur samples were
287 analyzed in three replicates, by performing three subsequent HS-SPME-GC-MS analyses on each
288 replicate. Total α - and β -thujone GC peak area estimation of each replicate was quantified by MHS-
289 SPME-GC-MS via external calibration, analyzing α - and β -thujone standard mixture at seven

290 concentration levels. (α -thujone: $y = 2205.2x - 88987.3$; $R^2 = 0.9996$; β -thujone: $y = 1977.6x -$
291 250263 ; $R^2 = 0.9995$).

292 The thujone amount in liqueur and plant material was used to calculate a linear regression, which
293 could then be used to predict the thujone content in liqueurs on the basis of the related plant
294 material (i.e. before the liqueur preparation, in agreement with the EC protocol¹³).

295 The linear regression model between thujone quantities in plants and liqueurs was not satisfactory
296 for prediction (**Supplementary Figure 1**). As a consequence, the possibility of predicting the
297 thujone level in liqueurs had to be investigated by a different approach. LDA was then adopted to
298 predict the probability of a liqueur to enter (or not) within the thujone EC limit on the basis of the
299 thujone content in the plant used to produce it.

300 **Thujone level prediction in liqueurs.** A new matrix was created using the total thujone amount as
301 categorical variable and HS-SPME-GC-MS compound abundances as continuous describing
302 variables. The samples were divided into two groups based on the total thujone concentration, using
303 the 35 mg/L EC limit to discriminate the high-thujone group from the low-thujone group.

304 With LDA, only one DF was calculated to discriminate the two groups, because the categorical
305 response variable was represented by only two possibilities (above and below the EC limit).

306 As reported in **Figure 5**, a clear discrimination between the two groups was possible using HS-
307 SPME-GC-MS data. This result shows that the samples above the limit have the lowest values in
308 DF scores, while the samples below the limit have the highest DF scores. The result is clear
309 considering that the two group distributions did not overlap, giving the model a strong possibility of
310 recognizing the response group. The model efficiency is supported by the confusion matrix (**Table**
311 **4**) that reports high validation (<35 ppm=97.6%; >35 ppm=100.0%) and prediction (<35
312 ppm=95.2%; >35 ppm=92.9%) rates.

313 The combination of HS-SPME-GC-MS data with LDA modeling achieved satisfactory prediction
314 rates. However, the analytical process (HS-SPME-GC-MS) is still too time-consuming (70
315 min/sample). The method proposed is reliable in qualitative and quantitative analyses when it is

316 necessary to consider each compound, but it should be made easier and quicker to fit better with the
317 aim of this study. The analysis can significantly be speeded up by eliminating the GC separation
318 step and reducing the sampling time to five minutes, thus reducing the total analysis time to eight
319 minutes.

320 The scan range of the non-separative HS-SPME-MS method has been fixed between 35 and 250
321 *m/z*. As in the previous LDA model, a matrix was created based on the total thujone amount as
322 categorical variable, but using the 215 fragment ions as continuous describing variables of each
323 sample.

324 As reported in **Figure 6** and in **Table 5**, the LDA results on non-separative MS data were slightly
325 less predictive than those obtained with GC-MS data, but they were still in a significant range of
326 significance and prediction. The model efficiency for non-separative approach is supported by the
327 confusion matrix (**Table 5**) that reports good validation (<35 ppm=97.7%; >35 ppm=96.4%;
328 mean=97.1%) and prediction (<35 ppm=88.6%; >35 ppm=89.3%; mean=89.0%) rates.

329 In conclusion, the combination of chromatographic techniques (HS-SPME-GC-MS, HS-SPME-MS
330 and MHS-SPME-GC-MS) with a chemometric tool (LDA) has successfully been used to create
331 prediction models that can help to solve two important issues concerning *A. umbelliformis* and its
332 related g n pi liqueur. HS-SPME-GC-MS data recovered from a plant sample were found to be
333 good predictive markers for: i) plant geographical origin, and ii) thujone level in the plant-related
334 liqueur. Furthermore, a non-separative methods (HS-SPME-MS) was also developed to markedly
335 shorten the analysis time, while at the same time maintaining its capacity as prediction method for
336 thujone level in the related liqueur.

337 **ACKNOWLEDGMENT**

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339 metabolismo secondario di matrici di origine vegetale” financially supported by the Ricerca Locale
340 (Ex 60% 2015) of the University of Turin, Turin (Italy).

341

342 **ASSOCIATED CONTENT**

343 Supporting Information

344 **Supplementary Figure 1.**

345 Linear regression model between total thujone content of plant and liqueur. A, Correlation between
346 plant and liqueur MHS-SPME-GC-MS data. B, Correlation between MHS-SPME-GC-MS liqueur
347 data and plant-based predicted liqueur values

348 **Supplementary Table 1.**

349 Repeatability of HS-SPME-GC-MS analysis of all compounds detected.

350 **Supplementary Table 2**

351 Repeatability of HS-SPME -MS analysis of the 30 most abundant MS fragments.

352 **Supplementary Table 3**

353 ANOVA and Tukey Honestly Significant Difference (HSD) test on morphological parameters.

354 **Supplementary Table 4**

355 Results of LDA performed with morphological parameters. Discriminant Functions are analyzed
356 testing their significance in discriminating the different geographical origin in the Wilks' Lambda
357 Table. The second table reports the correlation between the three Discriminant Functions and the
358 morphological parameters.

359 **Supplementary Table 5**

360 Results of LDA performed with compounds detected by HS-SPME-GC-MS. Discriminant
361 Functions are analyzed testing their significance in discriminating the different geographical origin
362 in the Wilks' Lambda Table. The second table reports the DF value for each geographical origin.
363 The third table reports the correlation between the three Discriminant Functions and the HS-SPME-
364 GC-MS compounds.

365

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465

466

467 **Figure captions**

468 **Figure 1**

469 Génépi illustrated to highlight phenological (A, B) and morphological (C, D, E) parameters
470 measured (Drawing: Karla Alejandra Palacios Antenucci).

471 **Figure 2**

472 Scatter plot of the first two Discriminant Functions of morphological data LDA. The geographic
473 origin is here reported by numbers: 1, Gran Paradiso; 2, Val Chisone; 3 Elva; 4, Valle Gesso.

474 **Figure 3**

475 *A. umbelliformis* Lam. HS-SPME-GC-MS chromatographic profile. Legend: 1) α -thujene, 2) α -
476 pinene, 3) camphene, 4) sabinene, 5) β -pinene, 6) α -phellandrene, 7) α -terpinene, 8) *p*-cymene, 9)
477 1.8-cineole, 10) γ -terpinene, 11) *cis*-sabinene hydrate, 12) α -terpinolene, 13) α -thujone, 14) β -
478 thujone, 15) *trans*-pinocarveol, 16) sabinol, 17) pinocarvone, 18) borneol, 19) terpinen-4-ol, 20) α -
479 terpineol, 21) cuminaldehyde, 22) α -copaene, 23) sabinyl propionate, 24) sabinyl isobutyrate, 25)
480 *trans*- β -farnesene, 26) sabinyl isovalerianate, 27) sabinyl valerianate, 28) δ -cadinene, 29)
481 caryophyllene oxide, 30) neryl isovalerianate

482 **Figure 4**

483 Scatter plot of the first two Discriminant Functions of HS-SPME-GC-MS data LDA. The
484 geographic origin is here reported by numbers: 1, Gran Paradiso; 2, Val Chisone; 3 Elva; 4, Valle
485 Gesso.

486 **Figure 5**

487 Histograms representing the frequencies of discriminant function scores of the analyzed samples by
488 HS-SPME-GC-MS. The upper panel shows samples below the limit, while the lower panel shows
489 samples above the limit

490 **Figure 6**

491 Histograms representing the frequencies of discriminant function scores of the analyzed samples by
492 HS-SPME-MS. The upper panel shows samples below the limit, while the lower panel shows
493 samples above the limit.

Tables

Table 1 - Geographic origin of *A. umbelliformis* used in the experiment as mother plants.

Geographic origin	Code	Locations	Altitude	Geographic coordinates
Gran Paradiso	GP	Campiglia (Torino)	1,400 m a.s.l.	45°32'27.00"N, 7°32'17.78"E
Val Chisone	VC	Pragelato (Torino)	1,500 m a.s.l.	45°00'36.41"N, 6°56'14.56"E
Elva	EL	Elva (Cuneo)	1,600 m a.s.l.	44°32'16.31"N, 7°05'33.87"E
Valle Gesso	VG	Castelmagno (Cuneo)	1,900 m a.s.l.	44°24'13.10"N, 7°09'53.98"E

Table 2 - Validation and prediction confusion matrices in Gran Paradiso (GP); Val Chisone (VC); Elva (EL); Valle Gesso (VG). The recognition rates of the model are not successful.

		Predicted Group Membership (%)			
		GP	VC	EL	VG
Validation	GP	66.7	25.0	0.0	8.3
	VC	18.5	66.7	3.7	11.1
	EL	3.8	19.2	76.9	0.0
	VG	7.4	7.4	7.4	77.8
Prediction	GP	50.0	33.3	4.2	12.5
	VC	51.9	29.6	3.7	14.8
	EL	7.7	42.3	42.3	7.7
	VG	11.1	14.8	22.2	51.9

Table 3 - Confusion matrices in validation and prediction in Gran Paradiso (GP); Val Chisone (VC); Elva (EL); Valle Gesso (VG) using LDA model created with HS-SPME-GC-MS data.

		Predicted Group Membership (%)			
		GP	VC	EL	VG
Validation	GP	89.1	10.9	0.0	0.0
	VC	11.1	88.9	0.0	0.0
	EL	1.9	1.9	96.2	0.0
	VG	0.0	0.0	0.0	100.0
Prediction	GP	82.6	17.4	0.0	0.0
	VC	13.0	87.0	0.0	0.0
	EL	1.9	1.9	94.3	1.9
	VG	0.0	0.0	3.7	96.3

Table 4 - Confusion matrix for liqueur thujone level prediction with HS-SPME-GC-MS data.

		Predicted Group (%)	
		< 35 ppm	> 35 ppm
Validation	< 35 ppm	97.6	2.4
	> 35 ppm	0.0	100.0
Prediction	< 35 ppm	95.2	4.8
	> 35 ppm	7.1	92.9

Table 5 - Confusion matrix for liqueur thujone level prediction with HS-SPME-MS data.

		Predicted Group (%)	
		< 35 ppm	> 35 ppm
Validation	< 35 ppm	97.7	2.3
	> 35 ppm	3.6	96.4
Prediction	< 35 ppm	88.6	11.4
	> 35 ppm	10.7	89.3

Figures

Figure 1

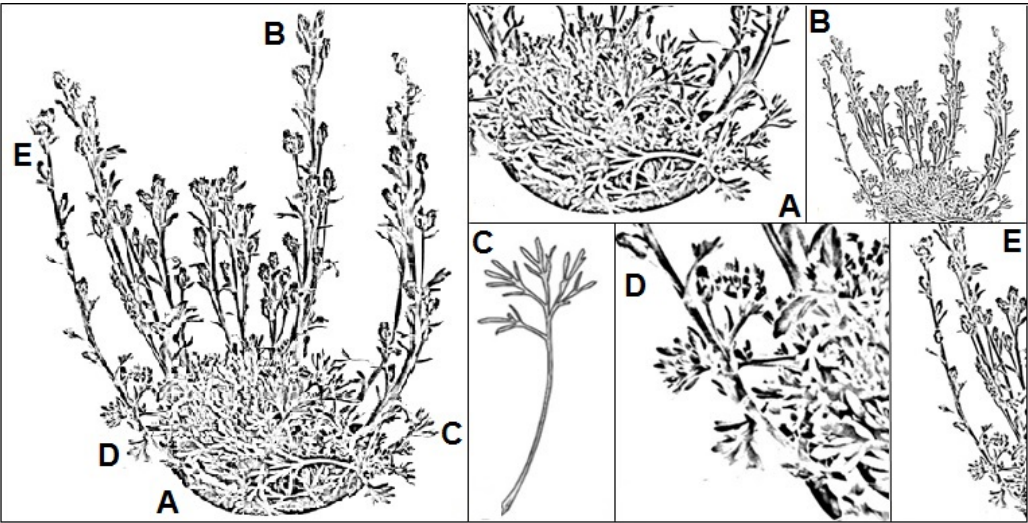


Figure 2

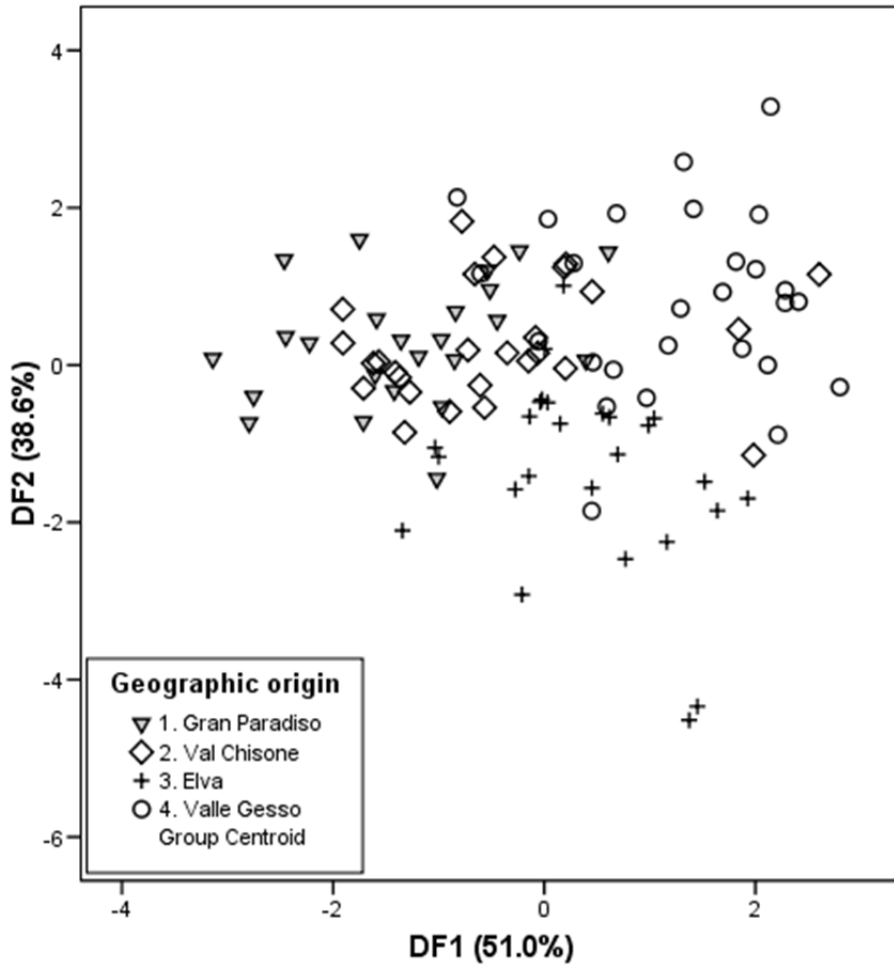


Figure 3

Abundance

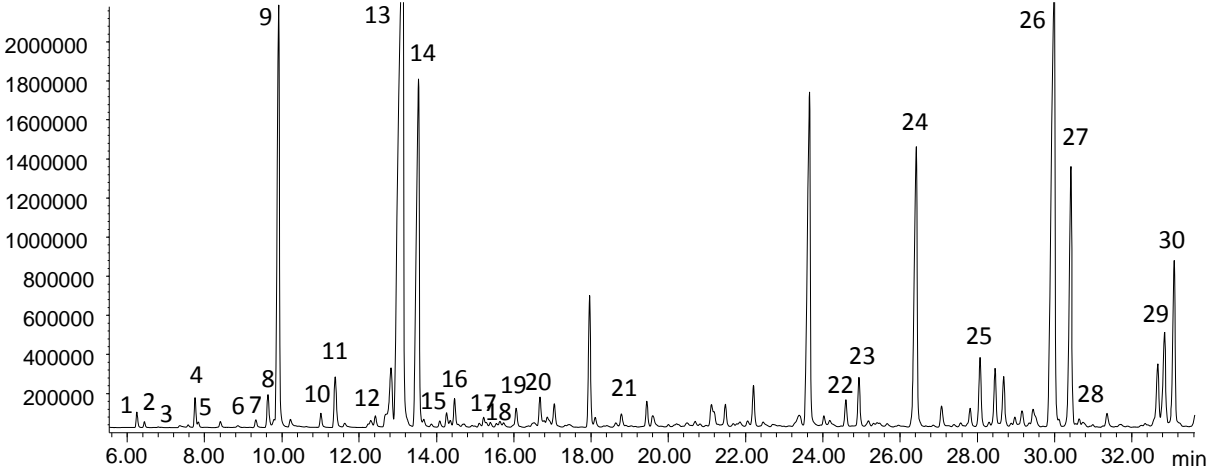


Figure 4

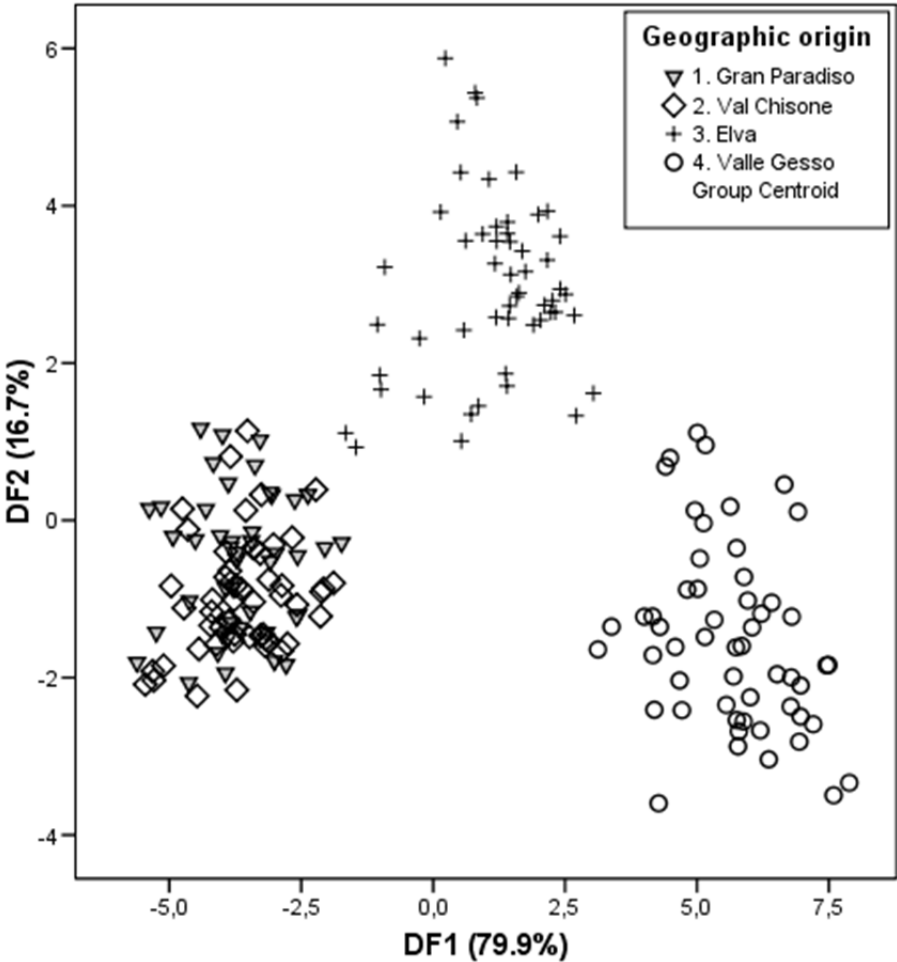


Figure 5

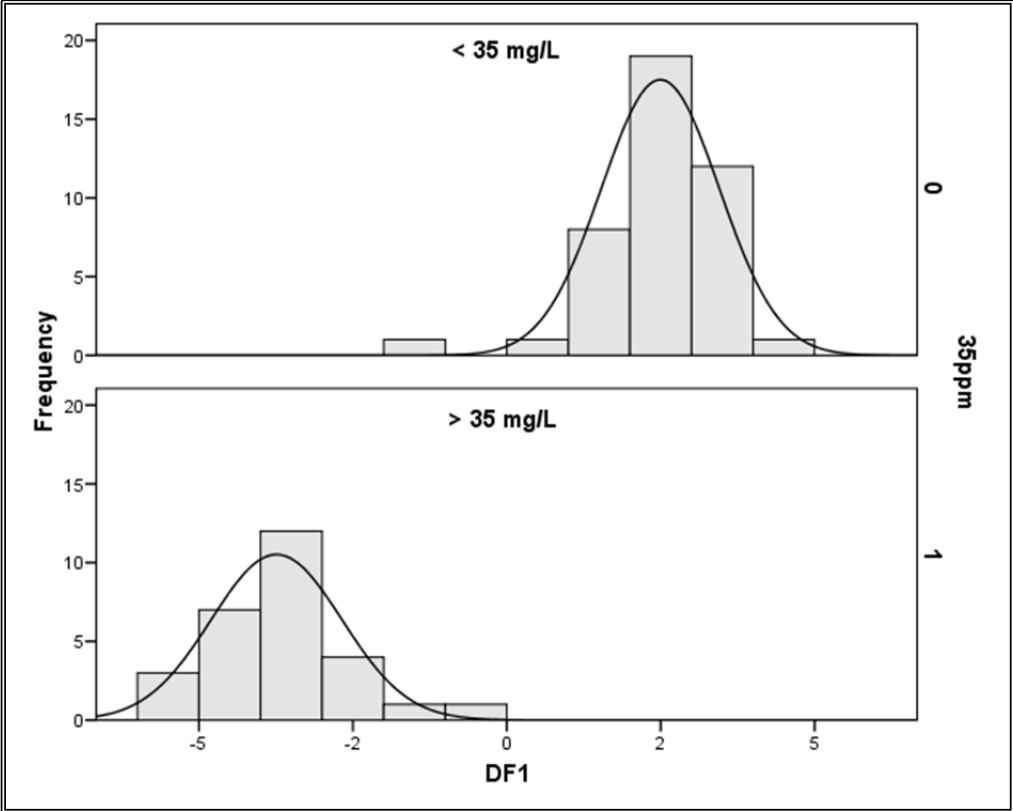
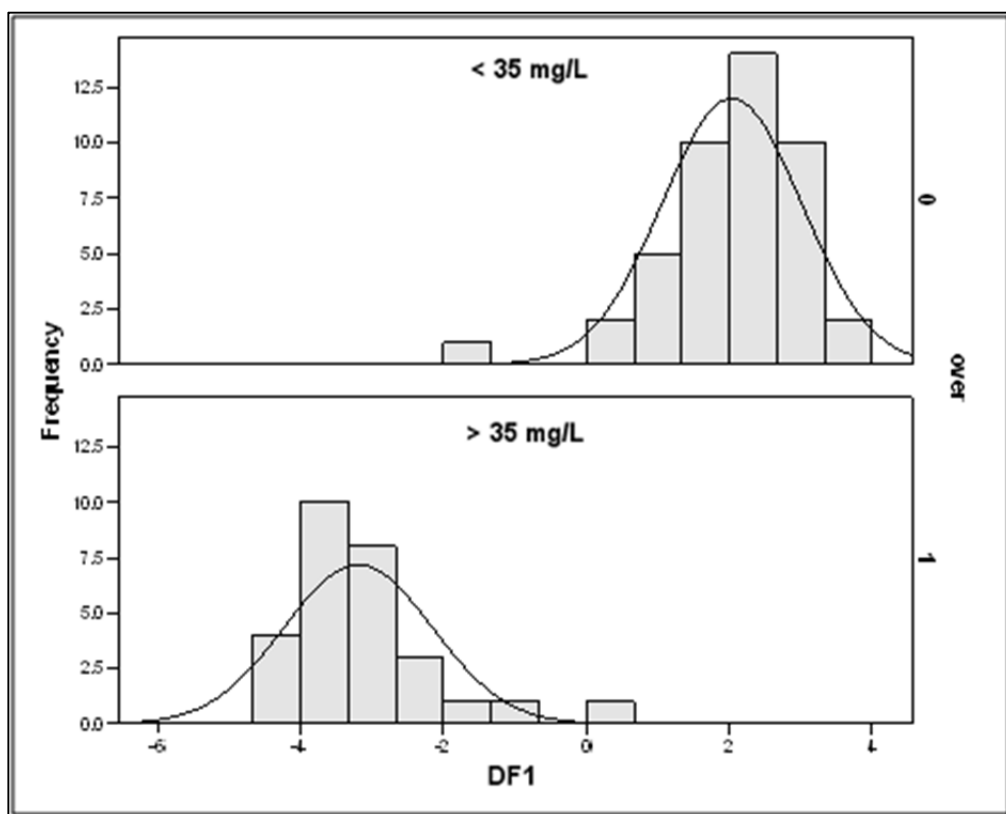


Figure 6



GRAPHIC FOR TABLE OF CONTENTS

