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# **Comprehensive Chemical Fingerprinting of High-Quality Cocoa at Early Stages of Processing: Effectiveness of Combined Untargeted and Targeted Approaches for Classification and Discrimination**

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1 **Abstract**

2 This study investigates chemical information in the volatile fractions of high-quality cocoa  
3 (*Theobroma Cacao* L. Malvaceae) from different origins (Mexico, Ecuador, Venezuela, Colombia,  
4 Java, Trinidad, and Sao Tomè) produced for fine chocolate. The study explores the evolution of  
5 the entire pattern of volatiles in relation to cocoa processing (raw, roasted, steamed, and ground  
6 beans). Advanced chemical fingerprinting (e.g., combined *Untargeted* and *Targeted* (UT)  
7 *fingerprinting*) with comprehensive two-dimensional gas chromatography (GC×GC) coupled with  
8 mass spectrometry (MS) enables advanced pattern recognition for classification, discrimination,  
9 and sensory-quality characterizations. The entire data-set is analysed for 595 reliable 2D peak-  
10 regions, including 130 known analytes and 13 potent odorants. Multivariate analysis (MVA) with  
11 unsupervised exploration (principal component analysis (PCA)) and simple supervised  
12 discrimination methods (Fisher ratios and linear regression trees) reveal informative patterns of  
13 similarities and differences and locate characteristic compounds related to samples origin and  
14 manufacturing step.

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16

17

18 **Key-words**

19 *Theobroma Cacao* L.; combined untargeted and targeted fingerprinting; comprehensive two-  
20 dimensional gas chromatography-mass spectrometry; classification and discrimination models;  
21 key-aroma compounds

22

23

## 24 Introduction

25 Cocoa, produced from cocoa beans (*Theobroma Cacao* L. *Malvaceae* family), is a crop of  
26 great economic relevance as the main raw ingredient for chocolate manufacturing.<sup>1</sup> Cocoa and  
27 chocolate are consumed worldwide and their popularity is primarily related to the pleasant  
28 sensory properties, although, recent evidence of several health benefits open new market  
29 perspectives and potential use in functionalized food(s).<sup>2-7</sup>

30 *Theobroma cacao* L. is a tree crop native to tropical forests of American continent. Recent  
31 studies, focused on cocoa germoplasm<sup>8</sup>, defined 10 major genetic clusters, or groups named:  
32 Marañon, Curaray, Criollo, Iquitos, Nanay, Contamana, Amelonado, Purūs, Nacional and Guiana.  
33 The new classification reflected accurately the genetic diversity available overcoming the  
34 traditional classification as Criollo, Forastero or Trinitario.

35 Cocoa quality and economic value are more strictly related to the unique and complex  
36 flavours. The sensory profile (aroma, taste, mouth feeling, and texture) is a key-factor in  
37 obtaining premium quality products suited to consumer preferences. Flavours develop from  
38 complex biochemical and chemical reactions occurring at post-harvesting and vary with  
39 genotype, geographical origin, farming practices, and technological processing.<sup>9</sup> Above all, post-  
40 harvest treatments and, in particular, fermentation<sup>10,11</sup> and roasting<sup>12</sup> are key steps in the  
41 formation of the characteristic cocoa aromas: in fact, the roasting of unfermented beans results  
42 in a product with a poor and unsatisfactory aroma profile.<sup>13</sup> Over the last few decades, several  
43 hundreds of volatiles have been identified in cocoa volatile fractions, including potent odorants  
44 whose particular distribution provides a diagnostic indicator for aroma qualification and products  
45 discrimination. The *molecular sensory science* approach, for example, was adopted to identify  
46 the *aroma blueprint* of different cocoa and chocolate products<sup>14-16</sup> while HS-SPME coupled with  
47 mono-dimensional (1D) GC-MS analysis is the technique of choice for cocoa volatile organic

48 compounds (VOCs) investigations.<sup>17,18</sup> Hyphenated techniques like in-line roasting in cooled  
49 injectors (ILR-CIS) and GC-MS were proposed for assessing process quality<sup>19</sup> and HS-SPME-GC-  
50 MS and direct MS-fingerprinting were combined to characterize cocoa volatiles.<sup>20</sup>

51 In this context, multidimensional analytical techniques, especially comprehensive two-  
52 dimensional gas chromatography (GC×GC) coupled with mass spectrometry (MS) are promising,  
53 powerful approaches for detailed characterization of the complex mixtures of cocoa volatiles as  
54 it has been proven for other foods<sup>21,22</sup>. GC×GC exploits the separation and detection potential of  
55 two separation dimensions providing increased separation power, meaningful 2D  
56 chromatographic patterns with analytes structurally ordered in the chromatographic plane and  
57 enhanced sensitivity derived from the band focusing during modulation.<sup>23–25</sup> Compared to 1D  
58 platforms, GC×GC-MS improves the effectiveness of sample profiling, fingerprinting and, thereby,  
59 classification and discrimination.<sup>21,24,26–29</sup>

60 In the panorama of existing studies, only a few have exploited the full potential of GC×GC  
61 to explain the complex information in cocoa volatile fractions or proposed effective methods  
62 capable of replacing multiple, less-informative, 1D separation methods based on targeted  
63 analysis. In 2009, Humston and co-workers<sup>30</sup> developed and evaluated an analytical procedure  
64 combining HS-SPME and GC×GC with time-of-flight (TOF) MS, to study volatiles from cocoa beans  
65 of different geographical origin, at two storage conditions, and with low or high moisture  
66 content. Within the entire set of detectable analytes, they identified four compounds (i.e., *acetic*  
67 *acid*, *nonanal*, *tetramethylpyrazine*, and *trimethylpyrazine*) showing consistent quantitative  
68 changes depending on bean storage and not on cocoa origin.

69 More recently, Oliveira et al.<sup>31</sup> investigated the volatile fraction of cocoa nibs from Brazil  
70 and Ivory Coast by HS-SPME-GC×GC-MS and GC×GC-FID to select informative analytes for  
71 samples differentiation. First, they applied PCA on GC×GC-FID data to evaluate samples

72 clustering; then, selected samples were submitted to GC×GC-MS to identify the most informative  
73 compounds. Within 20 identified analytes, 15 were found to be present in different amounts in  
74 samples of the two origins under study.

75  
76 The present study investigates the unique VOCs signatures from commercial grade, high-  
77 quality cocoa with a novel pattern recognition strategy that combines untargeted and targeted  
78 fingerprinting to GC×GC-MS data. Samples of interest for fine chocolate production, and from  
79 different geographical provenience (Mexico, Ecuador, Venezuela, Colombia, Java, Trinidad, and  
80 Sao Tomè) are studied along the early stages of industrial processing (raw, roasted, steamed, and  
81 nibs). The complex fraction of volatiles is extracted by automated HS-SPME sampling and  
82 subsequently analyzed by GC×GC-MS with thermal modulation. Advanced pattern recognition by  
83 UT fingerprinting strategy<sup>32</sup> is tested to validate its effectiveness to exploit chemical information  
84 encrypted in VOCs signatures. 2D data matrices are mined to explore different issues such as  
85 origin/process characteristics and sensory profile(s) differentiation.

86

## 87 **Materials and methods**

### 88 **Reference compounds and cocoa samples**

89 Pure reference standards for identity confirmation (key-aroma compounds and  
90 informative volatiles) of *acetic acid*, *3-methylbutanoic acid*, *3-methylbutanal*, *2-phenylethanol*, *2-*  
91 *heptanol*, *butanoic acid*, *2-methylbutanal*, *linalool*, *phenylacetaldehyde*, *2-ethyl-3,5-*  
92 *dimethylpyrazine*, *4-hydroxy-2,5-dimethyl-3(2H)-furanone*, *2-ethyl-3,6-dimethylpyrazine*, *(E,E)-*  
93 *2,4-nonadienal*, *dimethyl trisulfide*, *2-methylpropanoic acid*, *ethyl-2-methylbutanoate*, and *n-*  
94 *alkanes (n-C9 to n-C25)* for Linear Retention Index ( $I^T_s$ ) determination were from Sigma-Aldrich  
95 (Milan, Italy).



96 Internal standards (ISTDs) for analyte response normalization were  $\alpha$ - and  $\beta$ -*thujone* from  
97 Sigma Aldrich (Milan, Italy). A standard stock solution of ISTDs at 100 mg/L was prepared in  
98 *dibutylphthalate* (Sigma-Aldrich, Milan, Italy) and stored in a sealed vial at -18°C.

99 High-quality cocoa samples (*Theobroma cacao* L.) of commercial grade were selected by  
100 confectionery experts on the basis of their peculiar sensory characteristics. Descriptive sensory  
101 analysis (data not shown) was performed by company internal panel to drive processing  
102 parameters toward a desirable sensory quality. Origins were: Ecuador, Venezuela, Colombia,  
103 Trinidad, Mexico from Chontalpa region of Tabasco, Java and Sao Tomè. Samples information is  
104 provided as supplementary material in Supplementary Table 1 - ST1. Chontalpa is a top-quality  
105 area for cocoa production that was recognized by the Slow Food Presidium in 2007 after severe  
106 floods destroyed most of the cocoa plantations.

107 All samples were harvested in 2014; they were analyzed at four different technological  
108 stages: raw, roasted, steamed nibs obtained after the removal of bean shells (4 processing steps).

109 Processing was by Guido Gobino srl (Turin, Italy) in three replicated batches using time  
110 and temperature protocols between 100 and 130°C for a timing from 20 up to 40 minutes.  
111 Processing was optimized for each origin and driven by a desirable flavour development. Hot-air  
112 roasting was conducted in a vertical roaster designed by Bühler AG (Uzwil, Switzerland).

113 Cocoa samples were freeze in liquid nitrogen immediately after each step of processing  
114 and then stored at -80°C. Before headspace analysis, samples were ground in a laboratory mill  
115 up to about 300  $\mu$ m (Grindomix GM200, Retsch, Haan, Germany); particle size homogeneity was  
116 verified by visual inspection. The resulting cocoa powder was then precisely weighted (1.500 g)  
117 in headspace glass vials (20 mL) and submitted to automated HS-SPME sampling.

118

119 **Automated Head Space Solid Phase Micro Extraction: sampling devices and conditions**

120 Automated HS-SPME was performed using a MPS-2 multipurpose sampler (Gerstel,  
121 Mülheim a/d Ruhr, Germany) installed on the GC×GC-MS system. SPME fibers,  
122 Divinylbenzene/Carboxen/Polydimethyl siloxane (DVB/CAR/PDMS)  $d_f$  50/30  $\mu$ m - 2 cm were from  
123 Supelco (Bellefonte, PA, USA). Fibers were conditioned before use as recommended by the  
124 manufacturer. The standard-in-fiber procedure was adopted to pre-load the ISTDs ( $\alpha$ - and  $\beta$ -  
125 thujone) onto the fiber before sampling. 5.0  $\mu$ L of ISTDs solution were placed into a 20 mL glass  
126 vial and submitted to HS-SPME at 50°C for 10 min.

127 After ISTDs loading, the SPME device was exposed to the headspace of cocoa samples (1.500 g)  
128 for 40 min at 50°C. Extracted analytes were recovered by thermal desorption of the fiber into the  
129 split/splitless (S/SL) injection port of the GC×GC system at 250°C for 5 min. Each sample was  
130 analyzed in duplicate.

131

### 132 **GC×GC-MS instrument set-up and analytical conditions**

133 GC×GC analyses were performed on an Agilent 6890 GC unit coupled with an Agilent  
134 5975C MS inert detector operating in the EI mode at 70 eV (Agilent, Little Falls, DE, USA). The  
135 transfer line was set at 270°C. An *Auto Tune* option was used and the scan range was set at  $m/z$   
136 40-240 with a scan rate of 12,500 amu/s to obtain a sampling frequency of 28 Hz.

137 The system was equipped with a two-stage KT 2004 loop thermal modulator (Zoex  
138 Corporation, Houston, TX) cooled with liquid nitrogen and controlled by Optimode™ V.2 (SRA  
139 Instruments, Cernusco sul Naviglio, MI, Italy). Hot jet pulse time was set at 250 ms, modulation  
140 time was 3s, and cold-jet total flow was progressively reduced with a linear function from 40% of  
141 Mass Flow Controller (MFC) at initial conditions to 8% at the end of the run. A deactivated fused  
142 silica capillary loop (1 m × 0.1 mm  $d_c$ ) was used.

143 The column set was configured as follows: <sup>1</sup>D SolGel-Wax column (100% polyethylene  
144 glycol) (30 m × 0.25 mm d<sub>c</sub>, 0.25 μm d<sub>f</sub>) from SGE Analytical Science (Ringwood, Australia) coupled  
145 with a <sup>2</sup>D OV1701 column (86% polydimethylsiloxane, 7% phenyl, 7% cyanopropyl) (1 m × 0.1 mm  
146 d<sub>c</sub>, 0.10 μm d<sub>f</sub>), from J&W (Agilent, Little Falls, DE, USA).

147 SPME thermal desorption into the GC injector port was under the following conditions:  
148 split/splitless injector in split mode, split ratio 1:5. Carrier gas was helium at a constant flow of  
149 1.2 mL/min. The oven temperature program was: from 40°C (1 min) to 200°C at 3°C/min and to  
150 250°C at 10°C/min (5 min).

151 The *n*-alkanes liquid sample solution for  $I_s^T$  determination was analyzed under the  
152 following conditions: split/splitless injector in split mode, split ratio 1:50, injector temperature  
153 250°C, and injection volume 2 μL.

154

#### 155 **Data acquisition and data elaboration**

156 Data were acquired by Agilent MSD ChemStation ver D.02.00.275 and processed by GC  
157 Image® GC×GC Edition Software, Release 2.6 (GC Image, LLC Lincoln NE, USA). Statistical analysis  
158 was performed with XLstat (Addinsoft, New York, NY USA).

159

#### 160 **UT fingerprinting work-flow**

161 *Untargeted and Targeted (UT) fingerprinting* was carried out by the template matching  
162 approach, introduced by Reichenbach and co-workers in 2009<sup>33</sup> and following a work-flow  
163 previously validated for olive oil volatiles investigation.<sup>32</sup> The approach uses metadata collected  
164 from 2D peak patterns (retention times, MS fragmentation patterns, and single ions and/or total  
165 ions response) and establishes reliable correspondences between the same chemical entities

166 across multiple chromatograms. The output is a data matrix of aligned 2D peaks and peak-regions  
167 and their related metadata available for comparative purposes and further processing.

168 Targeted analysis focused on 130 compounds tentatively identified by matching their EI-  
169 MS fragmentation pattern (NIST MS Search algorithm, ver 2.0, National Institute of Standards  
170 and Technology, Gaithersburg, MD, USA, with Direct Matching threshold 900 and Reverse  
171 Matching threshold 950) with those collected in commercial (NIST2014 and Wiley 7n) and in-  
172 house databases. As a further check for identification, experimental Linear Retention Indices ( $I^T_s$ )  
173 were computed and compared to the tabulated indices.<sup>34</sup>

174 Untargeted analysis was based on peak-regions features<sup>35,36</sup> and was performed  
175 automatically by GC Image Investigator™ R 2.6 (GC-Image LLC, Lincoln NE, USA). The untargeted  
176 analysis included *all peak-regions* above the fixed peak response threshold of 5,000 counts  
177 together with all targeted peaks and related metadata. This process<sup>32,35–39</sup> aligned the feature  
178 template to each of the 168 chromatograms (7 cocoa origins × 4 technological steps × 3 technical  
179 batches × 2 analytical replicates) using a set of *registration peaks* that were reliably matched  
180 across all chromatograms. The resulting data matrix for untargeted and targeted reliable peak-  
181 regions was 168 × 595; column bleeding and SPME fiber interferent peaks were removed before  
182 chemometric analysis. Response data from all cross-aligned 2D peak-regions were used for  
183 multivariate analysis (MVA) and supervised discrimination approaches (Fisher ratio and  
184 regression trees).

185 Fisher ratios were used to measure class separation for individual features relative to the  
186 variance within classes. For the same number of observations in two classes, the square-root of  
187 the Fisher ratio is the t-value. For more than 20 samples (e.g., 21 samples at each of the four  
188 processing stages), a Fisher ratio of 1 has a p-value of 16%, a Fisher ratio of 1.77 exceeds 90%  
189 confidence, and a Fisher ratio of 6.45 exceeds 99% confidence. In this study, Fisher ratios (F value)

190 were calculated during the UT fingerprinting elaboration by the Image Investigator™ (GC Image  
191 v2.6) on normalized 2D peak-region volumes considering each class against the superset of all  
192 other classes (one vs. all).

193 Repeatability and intermediate precision results on retention times ( $^1t_R$  and  $^2t_R$ ) and on  
194 Normalized 2D volumes is reported as supplementary material. Repeatability was evaluated on  
195 single batch Chontalpa nibs replicate analyses over a three days time interval (three replicated  
196 samples) while intermediate precision was calculated on ISTDs ( $\alpha$ - and  $\beta$ -thujone) 2D peaks from  
197 all nibs samples analyzed over the one-month period (42 runs). Data refers of good method  
198 precision<sup>40</sup> on both: (a) retention times, where RSD % ranges from 0.06 to 3.43 (average value  
199 0.59) for  $^1D$  and from 0.83 and 6.12 (average value 2.68) for the  $^2D$ . Normalized 2D Volumes were  
200 always below 20% with an average RSD of 6.85%. 2D Normalized Volumes of the two ISTDs ( $\alpha$ -  
201 and  $\beta$ -thujone), monitored over a wider period, never exceeded the 14% of RSD.

202

## 203 **Results and discussion**

204 This study exploits the power of GC×GC-MS for the detailed chemical profiling of complex  
205 samples, harnessing its intrinsic potential as a highly informative fingerprinting tool. Thanks to  
206 dedicated pattern recognition approaches, the large amount of (chemical) information encrypted  
207 in cocoa volatiles distributions, can be rationalized and mined to find compositional  
208 similarities/differences (fingerprinting) and to explain the informative role of single chemicals,  
209 whose distribution provides indications for origin traceability, effects of manufacturing  
210 processes, and aroma quality.

211 The following sections illustrate: (a) the chemical complexity of the volatile fraction of  
212 high-quality *Theobroma cacao* samples, as revealed by combining targeted and untargeted  
213 investigations; (b) the particular distributions of informative analytes (key-aroma compounds

214 and technological sensitive analytes) within samples and their evolution along processing steps,  
215 (c) how simple supervised approaches could support the selection of informative chemicals to  
216 discriminate samples.

217

## 218 **Information encrypted on cocoa volatiles distribution**

219 The high chemical complexity of cocoa volatile fractions results from many chemical  
220 reactions, most of them catalyzed by specific enzymes (endogenous or exogenous from moulds,  
221 yeasts and bacteria) and occurring at the different stages of its processing. Influential factors  
222 have been extensively reviewed by Afoakwa et al<sup>9,41</sup> and include some of the variables considered  
223 in our sampling design: roasting (time/temperature) and other physical and mechanical  
224 treatments such as debacterization by steaming, and grinding.

225 Within the 595 detected VOCs by GC×GC-MS (peak-regions corresponding to detectable  
226 analytes in at least two samples of the set), 130 analytes were tentatively identified and reported  
227 in **Table 1**. Each analyte is characterized by absolute retention times ( $^1t_R$  - min and  $^2t_R$  - sec),  
228 experimental  $I^T_s$ , and odor descriptors as reported in reference literature.

229 Figure 1 visualizes a heat-map of the relative distributions (Normalized 2D Peak Volumes)  
230 of 595 untargeted peak-regions, including the 130 known analytes. Columns follow processing  
231 stages from raw to grinded beans after steaming. Analytes are ordered according to their inter-  
232 class variance. Normalized peak volumes values were mean and centered before colorization.  
233 Color scale varies between red (low abundance) to green (high abundance).

234 The evolution of the volatiles profile along the different steps of processing is illustrated  
235 by changes in the heat-map colour spots (Fig. 1). In particular, after roasting and steaming, when  
236 volatiles are developed from their non-volatile aroma precursors, dark spots predominate while

237 several analytes, already present in raw beans, increase their relative abundances (quantitative  
238 changes).

239

#### 240 **Potent odorants distribution within samples and their evolution along processing**

241 Within the volatile fraction, the most significant changes occur for key-aroma and some  
242 technologically sensitive analytes (technological markers), in close accordance with reference  
243 studies.<sup>1,42,41</sup>

244 Cocoa key-aroma compounds, identified by Schieberle and co-workers,<sup>14–16</sup> deserve a  
245 detailed discussion, in that their distribution is fundamental for aroma properties. They include  
246 several chemical classes, especially alkyl pyrazines (*2,3,5-trimethylpyrazine*, *2-ethyl-3,5-*  
247 *dimethylpyrazine*, and *3,5-diethyl-2-methylpyrazine*) which impart characteristic earthy notes.  
248 Another important set of key-volatiles are short-chain and branched fatty acids: *acetic acid*,  
249 *butanoic acid*, *2-methylpropanoic acid*, and *3-methylbutanoic acid*, whose presence, at high  
250 concentrations, can impart off-flavours due to their rancid, sour, and sweaty notes. Strecker  
251 aldehydes (*2-* and *3-methylbutanal*), formed during fermentation and roasting, impress malty  
252 and buttery notes, and *phenylacetaldehyde*, derived from L-phenylalanine (L-Phe), is responsible  
253 for a pleasant honey-like note. Other key analytes are esters (*ethyl-2-methylbutanoate* – fruity,  
254 *2-phenylethyl acetate* – flowery), linear alcohols (*2-heptanol* – citrusy), phenyl propanoids  
255 derivatives (*2-phenylethanol* – flowery), and sulphurous derived compounds (*dimethyl trisulfide*).

256 Raw cocoa beans (just fermented) have specific distributions of potent odorants related  
257 to origin. Profiling data, in agreement with reference studies,<sup>1,41</sup> show that the volatile fraction  
258 of raw beans is dominated by short-chain fatty acids, especially *3-methylbutanoic* and *acetic acid*,  
259 that result from the enzymatic degradation of the pulp during fermentation. In particular, *acetic*  
260 *acid* is the most abundant volatile and is present at high levels in unroasted beans (high Odour

261 Activity Value <sup>16</sup>), giving an intense vinegar-like perception which can affect cocoa aroma quality.  
262 However, during cocoa processing (roasting, above all) and later, during chocolate manufacturing  
263 (conching and refining), undesired volatiles with low boiling points, such as *acetic acid*, are  
264 removed resulting in a drastic decrease of its concentration (up to 70%).<sup>16</sup>

265 During the fermentation of raw beans, non-volatile aroma precursors obtained through  
266 the degradation of seeds storage proteins and carbohydrates react, mainly under enzymatic  
267 control, and generate odor-active volatiles (alcohols, esters, aldehydes, and organic acids).  
268 Bacteria and moulds are fundamental at this stage<sup>9,41</sup>.

269 Roasting has a larger impact on aroma: alkyl pyrazines and Strecker aldehydes (*3-*  
270 *methylbutanal* and, in some cases, *phenylacetaldehyde*) show a large increase after this stage.  
271 Roasting has only a minor impact on *3-methylbutanoic acid* and esters (rancid smelling), which  
272 were detected in similar amounts before and after this process. As general consideration, the  
273 differences in the volatile profiles between unroasted and roasted beans are quantitative rather  
274 than qualitative. Supplementary Figure 1 (SF1) illustrates GC×GC patterns and their evolution  
275 across stages for cocoa harvested in the Chontalpa region (Tabasco, Mexico). Relative  
276 distribution differences of some potent odorants, between raw and roasted beans, are visually  
277 shown, in logarithmic scale, on spider diagrams in Figure 2. Sample origins illustrated are  
278 Chontalpa, Mexico (2A); Venezuela (2B); and Sao Tomè (2C) and quantitative changes refer to  
279 raw (green lines) and roasted (brown lines) stages.

280 The Chontalpa sample from Mexico (2A) average profile shows a remarkable increase for  
281 the Strecker aldehyde *3-methylbutanal*; alkyl pyrazines, with earthy and roasty notes; *2-*  
282 *heptanol*, with a citrusy smell; and *dimethyl trisulfide*, whereas the amounts of other  
283 characteristic odorants (*butanoic acid*, *3-methylbutanoic acid*, *ethyl-2-methylbutanoate*,  
284 *phenylethylalcohol*, etc.) remain rather similar, even after roasting.



285 The distribution profile of the Venezuela sample (2B) is characterised by a more significant  
286 increase (compared to Chontalpa) of *3-methylbutanal* (malty odour), a significant increase for  
287 *phenylacetaldehyde* (opposite the trend for Chotalpa), and no change for *2-heptanol*.

288 The Sao Tomè sample (2C) shows a different behaviour: even though some aroma and  
289 technological markers increase after roasting (*3-methylbutanal* and pyrazines), the raw and  
290 roasted cocoa present very similar patterns, as is the case for the Java sample (data not shown).

291

292

### 293 **Untargeted and Targeted (UT) Fingerprinting results**

294 The distribution of all detected VOCs (known and unknown analytes) is a potentially  
295 informative fingerprint for geographical origin and manufacturing stage differentiation.  
296 Unsupervised multivariate analysis, i.e., PCA, was applied to map the natural conformation  
297 (groups) of samples and to localize informative chemicals responsible for variations.

298 In the first step, PCA was performed on the matrix combining information from 130  
299 targeted analytes in samples with different origins (CH-Chontalpa, VE-Venezuela, CO-Colombia,  
300 EC-Ecuador, JA-Java, TR-Trinidad, ST-Sao Tomè), manufacturing stages, and processing batches  
301 (3). Analytical replicates (2) were averaged. Auto-scaling was applied as pre-processing step and  
302 baseline correction was performed on the 2D data by GC Image software.

303 Figures 3A-B show the scores plot for the first two principal components (F1-F2 plane) for  
304 raw (Fig.3A) and roasted (Fig.3B) cocoa, based on the 84 × 130 matrix (samples × targets). The  
305 variance explained by the first two components was similar in all elaborations (including those  
306 based on steamed and nibs, not shown), ranging from a minimum of 48.83% for roasted samples  
307 (30.22 % for F1, 18.61% for F2) to the 53.74 % of raw cocoa (30.00 % for F1 and 23.74 % for F2).  
308 Origin dominates group conformation and grouping is maintained through manufacturing steps.

309 In particular, PCA clusters cocoa samples in three main sub-groups. In the first sub-group  
310 (highlighted with blue circles), cocoa from Ecuador, Venezuela and Colombia are close together  
311 at all stages. This outcome is consistent with their aroma profiles, considered relatively similar by  
312 confectionery experts. In the second sub-group (green circles), cocoa from Trinidad has a  
313 distinctive chemical fingerprint that yields independent clustering at all stages. In the third sub-  
314 group (red circles), Chontalpa,Java (and Sao Tomè show more similar chemical fingerprints,  
315 despite their different geographical provenience.

316 Cocoa clustering results from different variables (loadings plots not reported): for  
317 example, raw beans from Chontalpa and Sao Tomè had higher levels of some potent odorants  
318 such as *acetic acid* (sour), *phenylacetaldehyde* (honey-like), *2-phenylethanol*, and other volatiles  
319 such as esters (*ethyl hexanoate*, *ethyl octanoate*, and *ethyl decanoate*) and organic acids. The  
320 volatiles signatures of South America cocoa (Ecuador, Colombia, Venezuela) is connoted by the  
321 presence of short chain primary alcohols (*1-butanol*, *1-pentanol*, *1-hexanol*, *2-ethyl-1-hexanol*, *2-*  
322 *hexanol*, and *2-heptanol* (citrusy)) and *3-methylbutanoic acid* (rancid). These analytes (esters,  
323 alcohols, and acids) and some detectable linear aldehydes (*hexanal*, *octanal* and *nonanal*) are  
324 formed mostly during fermentation. The cluster of roasted samples from Chontalpa, Java, and  
325 Sao Tomè have a distinctive fingerprint of *alkyl pyrazines* (*2,3,5-trimethyl*, *2-ethyl-3,5-dimethyl*,  
326 *2-ethyl-5(6)-methyl* and *3,5-diethyl-2-methyl pyrazine*), important processing markers. Roasted  
327 beans of South American cocoa are connoted by higher amounts of aromatic ketones (*1-hydroxy-*  
328 *2-propanone*, *2,3-pentanedione* and *2,3-butanedione*) and other volatiles such as *1H-pyrrole-2-*  
329 *carboxaldehyde* and *2-furanmethanol*.

330 Fisher ratio values were therefore used for supervised ranking and selection of highly  
331 informative features characterising the chemical fingerprints of different sample sets. Fisher  
332 ratios (F value) were calculated automatically during the *UT fingerprinting* elaboration on

333 normalized 2D peak-region volumes considering each class against the superset of all other  
334 classes (one vs. all).

335 Figure 4 shows bar plots of F values for classes of three origins (Chontalpa-Mexico, Java,  
336 and Trinidad) and two processing steps (roasting and steaming), with an arbitrarily fixed cut-off  
337 of 30. As seen in this plot, several analytes are distinctive for origin independent of processing  
338 (those with paired cyan and orange bars). In most cases, cocoa origins are described by the same  
339 variables at roasted and/or steamed stage.

340 Chontalpa and Java, which clustered together with Sao Tomè in the PCA elaboration, have  
341 distinctive signatures: Java has a characteristic distribution of alkyl pyrazines (*tetramethyl-, 2-*  
342 *ethyl-3,5-dimethyl-, 2,3,5-trimethyl-* and *3,5-diethyl-2-methylpyrazine*) that is preserved after  
343 steaming, with most of those F values increasing (e.g., from 67 to 413 for *tetramethylpyrazine*,  
344 from 320 to 820 for *2-ethyl-3,5-dimethylpyrazine*, etc.) indicating a stronger diagnostic role.

345 Chontalpa from Mexico is characterized by esters, responsible for fruity notes, which  
346 probably derive from fermentation processes. The most significant ones are *hexyl acetate* (F  
347 value 1139 for roasted, but only 73 for steamed) and *1-butanol-3-methyl acetate* (F value 440 for  
348 roasted and 409 for steamed). Moreover, *3-hydroxy-2-butanone*, a technological marker  
349 influencing buttery perception, plays a less significant role for both roasted and steamed  
350 samples.

351 Cocoa from Trinidad, independently clustered at all stages of processing, is connoted by  
352 a distinctive signature of phenyl-propanoid derivatives (*benzaldehyde* and *2-phenylalcohol*),  
353 some process markers (*2,6* and *2,3-dimethylpyrazine*), and *trans-linalool oxide*. The highest F  
354 value (495) is observed in the roasted sample for *ethyl butanoate* (sweet, fruity), an analyte that  
355 does not keep its information potential after steaming.

356 To confirm the results obtained with the targeted fingerprinting and to evaluate if new  
357 informative markers could be revealed within the entire volatile fraction, the study was extended  
358 to all detected analytes, including unknowns. The set of 595 peak-regions, included the 130  
359 target analytes (tentatively identified), was thereby used to validate targeted analysis results.

360 Figures 3C-D visualise PCA results with the 595 reliable peak-regions for raw (Fig. 3C) and  
361 roasted (Fig.3D) cocoa. Results are highly consistent with those from the targeted peaks  
362 distributions. Samples are clustered into three groups: Ecuador-Venezuela-Colombia (blue  
363 circles), Chontalpa-Sao Tomè-Java (red circles), and Trinidad (green circles). The total explained  
364 variability here ranges from 40.66% for roasted beans to 48.50% for steamed cocoa (data not  
365 shown).

366 Targeted peak-regions, included in the untargeted approach, cross-validate the  
367 classification based on previous PCAs: samples are described by almost the same variables and  
368 no additional informative roles of unknown features were hypothesized. This approach clearly  
369 highlights the strong accordance between targeted and untargeted fingerprinting for sample  
370 classification purposes suggesting that for some applications, untargeted fingerprinting is  
371 effective, efficient, and less time-consuming than targeted analysis.

372

### 373 **Samples classification and discrimination: variables selection strategy**

374 The classification and prediction potential of the proposed approach has a high risk of  
375 over-fittings due to the large number of analyte variables and the limited number of the samples  
376 under study. However, to demonstrate the flexibility of such comprehensive fingerprinting for  
377 pattern recognition, simple classification approaches have been adopted to define key-variables  
378 (explanatory quantitative variables) suitable to discriminate one sample, or a group of them,  
379 from others. This is illustrated by two following examples: (a) the identification of a univocal set

380 of processing variables capable of distinguishing raw from processed cocoa independently of  
381 origin and (b) the definition of origin-specific variables sensitive to thermal treatments (roasting  
382 and steaming).

383 The explanatory approach adopted was a regression tree analysis based on the CHAID  
384 algorithm.<sup>43,44</sup> The entire set of samples × target analyte variables was explored to find univocal  
385 variables indicating the effect of processing on the raw cocoa, independent of origin. The sample  
386 set was divided into estimation samples and validation samples. The validation set included the  
387 second of the three replicated batches of analyses (28 samples, then not included in the  
388 estimation set). The resulting regression tree correctly classified all samples from the  
389 estimation/training set (i.e., the confusion matrix for all processing steps had 100% true  
390 positives). In the validation test, the predictive model failed in classifying five steamed samples  
391 belonging to the nibs (3) and roasted (2) classes, but it was successful for all others (i.e., better  
392 than 82% correct). The most informative classification variable for discriminating raw from  
393 processed cocoas was *2,3-dihydro-3,5-dihydroxy-6-methyl-4H-pyran-4-one*. Its formation,  
394 promoted by heating, is related to the presence of fructose and β-alanine in raw cocoa.<sup>45</sup>  
395 Variables with a secondary role for discriminating processing stages were: *2,6-dimethylpyrazine*,  
396 *2,3,5-trimethylpyrazine*, *2-ethyl-5-methylpyrazine*, and the potent odorant (*E*)-*2-phenyl-2-*  
397 *butenal* (intense chocolate note).

398 Figure 5A shows the samples distribution as a function of two discriminating variables:  
399 *2,3-dihydro-3,5-dihydroxy-6-methyl-4H-pyran-4-one* and *2-ethyl-5-methylpyrazine*. Raw cocoa  
400 (green markers in Fig. 5A) is clearly differentiated by processed derivatives independent of the  
401 origin; as those samples are closely clustered in the bottom-left of plot. Roasted cocoa is  
402 relatively well distinguished, but more dispersed along the x-axis, with *2,3-dihydro-3,5-dihydroxy-*  
403 *6-methyl-4H-pyran-4-one* increasing from left to right. Steamed and ground samples are less

404 differentiated along the *y*-axis, representing the relative abundance of the earthy pyrazine (2-  
405 *ethyl-5-methylpyrazine*).

406 The second model was developed to discriminate cocoa nibs (i.e., the last stage of  
407 processing considered here) based on their origin. In this case, the model was effective with just  
408 three variables: *2-pentylfuran*, *2,3,5-trimethylpyrazine*, and *linalool*. Figure 5B shows the  
409 distribution of samples in three variables: *x*-axis *linalool*; *y*-axis *2,3,5-trimethylpyrazine* ; and  
410 bubble-size *2-pentylfuran*). This model for nibs discrimination confirms what it was shown by  
411 unsupervised approaches (PCA on targeted and on UT data, shown in Fig. 3). Samples from  
412 Ecuador and Colombia are aligned along *x* axis (higher abundance of *linalool*) together with the  
413 Venezuela samples. Java samples are connoted by a strong pyrazines signature (*2,3,5-*  
414 *trimethylpyrazine* is one of the most origin sensitive), whereas Chontalpa, Sao Tomè, and  
415 Venezuela samples are coherently positioned in the Cartesian space with lower amounts of both  
416 chemicals.

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## Figure Captions

**Figure 1:** heat map showing *UT fingerprinting* results (untargeted and targeted) on 595 reliable peak-regions detected in the headspace of cocoa samples of seven origins and analyzed at each step of technological processing (raw, roasted, steamed and nibs). The heat-map quantitative descriptors (Normalized 2D Peak Volumes) are colorized according to a linear scale. Colour intensity goes from red (minimum) to light green (maximum).

**Figure 2:** Distribution of key-aroma compounds in raw/fermented (green line) and roasted (brown line) cocoa from Chontalpa-Mexico, Venezuela and Sao Tomè. Relative abundances reported in logarithmic scale refer to normalized 2D Peak Volumes.

**Figure 3:** Scores plots on the first two principal components (F1-F2 plane), based on the targeted fingerprinting of (3A) raw/fermented cocoa beans and (3B) roasted cocoa beans of all origins (Chontalpa, Mexico (CH), Ecuador (EC), Venezuela (VE), Colombia (CO), Java (JA), Sao Tomè (ST), Trinidad (TR). The complete set of untargeted + targeted peak-regions (i.e., 595 peak-regions above the fixed threshold of 5,000 counts) resulted in the distribution shown in 3C for raw and in 3D for roasted cocoa samples.

Each origin is represented by three processing batches while the two analytical replicates have been averaged before statistical analysis.

**Figure 4:** histograms with most significant Fisher Ratio values obtained with *one-vs-all* comparison; (a) roasted and (b) steamed Chontalpa, (c) roasted and (d) steamed Java, (e) roasted and (f) steamed Trinidad, (g) roasted and (h) steamed Ecuador. F values were selected by above the fixed threshold of 30.

**Figure 5:** dispersion graphs illustrating the discrimination potential of: (5A) *2,3-dihydro-3,5-dihydroxy-6-methyl-4H-pyran-4-one* and *2-ethyl-5-methylpyrazine* on processed cocoa; and (5B) *2,3,5-trimethylpyrazine*, *linalool* and *2-pentylfuran* on cocoa nibs from different origin.

**Table 1:** list of targeted volatiles together with their absolute retention times ( $^1t_R$  min and  $^2t_R$  sec), experimental  $I^1_s$ , informative role and odour descriptors as reported in the reference literature <sup>14-</sup>

16,46,47

ID	Compound Name	$^1t_R$ (min)	$^2t_R$ (sec)	Exp $I^1_s$	Compound Confirmation	Informative Role	Odour descriptor
1	2-Methylpropanal	4.19	0.35	833	a	-	Green, pungent
2	Methyl acetate	4.59	0.52	853	a	-	-
3	2-Methyl tetrahydrofuran	4.94	0.69	870	b	-	-
4	Ethyl Acetate	5.09	0.59	878	a	-	Fruity, aromatic
5	2-Methylbutanal	5.44	0.64	895	a	-	Malty
6	3-Methylbutanal	5.50	0.65	898	a	Key-aroma marker	Malty
7	Ethanol	5.79	0.41	913	a	-	Ethanol-like
8	Ethyl propanoate	6.24	0.79	935	b	-	-
9	Ethyl-2-methylpropanoate	6.59	1.03	953	b	-	Fruity
10	2,3-Butanedione	6.64	0.55	955	a	Technological marker	Buttery
11	2-Pentanone	6.69	0.79	958	a	-	Fruity
12	Pentanal	6.84	0.79	965	a	-	Almond-like, pungent, malt
13	1-Methylpropyl acetate	6.84	1.03	966	b	-	-
14	2-Methylpropyl acetate	7.49	1.00	998	a	-	-
15	2-Butanol	7.60	0.55	1001	a	-	Winey
16	$\alpha$ -Pinene	7.69	1.93	1005	a	-	Harsh, terpene-like, minty
17	2-Ethyl-5-methyl-furan	7.89	1.00	1012	b	-	-
18	Ethyl butanoate	7.99	1.21	1016	b	-	Sweet, fruity
19	2-Methyl-3-Buten-2-ol	8.19	1.24	1022	b	-	-
20	Ethyl-2-methylbutanoate	8.54	1.41	1035	b	Key-aroma marker	Fruity
21	2,3-Pentandione	8.74	0.76	1041	a	Technological marker	Caramel
22	Ethyl-3-methylbutanoate	8.99	1.38	1050	b	-	Fruity
23	Dimethyl disulfide	9.13	0.83	1055	a	Technological marker	Sulfurous
24	2-Pentyl acetate	9.15	1.38	1056	b	-	-
25	Butyl acetate	9.19	1.14	1057	a	-	Fruity, herbaceous
26	Hexanal	9.54	1.14	1069	a	-	Tallowy, leaf-like
27	2-Methyl-1-propanol	9.69	0.59	1074	a	-	-
28	2-Methyl-2-butenal	10.04	0.86	1086	a	-	-
29	2-Pentanol	10.74	0.69	1108	a	-	Light, seedy, sharp
30	3-Methylbut-1-yl acetate	10.89	1.38	1112	a	-	-
31	Ethyl pentanoate	11.01	1.39	1115	b	-	Fruity, sweet
32	Butyl-2-methylpropanoate	11.19	1.83	1120	b	-	Fruity, sweet
33	4-Methyl-3-penten-2-one	11.29	1.00	1123	b	-	-
34	1-Butanol	11.64	0.59	1132	a	-	Winey
35	$\beta$ -Myrcene	12.24	1.83	1148	a	-	-
36	1-Pentylacetate	12.79	1.41	1163	a	-	Fruity, metallic, green
37	2-Heptanone	13.19	1.38	1173	a	-	Sweet, fruity
38	2-Ethylhexanal	13.39	1.76	1179	a	-	-
39	Limonene	13.74	1.93	1188	a	-	Citrus, mint
40	2-Methyl-1-butanol	14.19	0.66	1200	a	-	Fermented, fatty
41	Pyrazine	14.39	0.75	1205	a	-	Earthy
42	Butyl butanoate	14.64	1.86	1211	b	-	Fruity, flowery, sweet

43	2-Hexanol	14.69	0.83	1212	a	-	Mushroom, green
44	Ethyl hexanoate	14.94	1.74	1219	a	-	Fruity
45	2-Pentylfuran	15.09	1.52	1222	b	-	Buttery, green bean-like
46	(E)-2-methyl-2-butenolate	15.39	1.31	1229	b	-	-
47	1-Pentanol	15.89	0.69	1241	a	-	Sweet, pungent
48	2,4-Dimethyl-3-pentanol	16.34	0.66	1252	b	-	-
49	Methylpyrazine	16.69	0.79	1260	a	Technological marker	Earthy
50	Hexyl acetate	16.89	1.62	1265	a	-	Fruity
51	3-Hydroxy-2-butanone	17.24	0.62	1274	a	Technological marker	Buttery
52	2-Octanone	17.49	1.55	1280	a	-	Mould, green
53	Octanal	17.64	1.59	1283	a	-	Fatty, sharp
54	1-Hydroxy-2-propanone	17.94	0.52	1290	a	Technological marker	Buttery
55	2-Methyl-1-pentanol	17.99	0.76	1292	a	-	-
56	2-Ethyl-(E)-2-hexenal	18.04	1.59	1293	a	-	-
57	3-Hepten-2-one	18.09	1.56	1294	b	-	-
58	2-Heptanol	18.94	0.90	1314	a	Key-aroma marker	Citrusy
59	2,3-Octanedione	19.14	1.28	1319	b	Technological marker	-
60	2,5-Dimethylpyrazine	19.24	0.88	1321	a	Technological marker	Earthy
61	2,6-Dimethylpyrazine	19.34	0.90	1324	b	Technological marker	Earthy
62	Ethylpyrazine	19.54	0.89	1328	b	Technological marker	Earthy
63	6-Methyl-5-hepten-2-one	19.64	1.28	1331	a	-	Pungent, green
64	2,3-Dimethylpyrazine	20.09	0.93	1341	b	Technological marker	Earthy
65	1-Hexanol	20.29	0.79	1346	a	-	-
66	4-Hydroxy-4-methyl-2-pentanone	20.54	0.83	1352	b	-	Fruity, banana, soft
67	Dimethyl trisulfide	21.24	1.03	1368	a	Key-aroma marker	sulfury, cabbage
68	2-Ethyl-6-methylpyrazine	21.74	1.07	1380	a	Technological marker	Earthy
69	2-Nonanone	21.94	1.72	1385	a	-	-
70	2-Ethyl-5-methylpyrazine	22.04	1.07	1387	a	Technological marker	Earthy
71	Nonanal	22.14	1.72	1389	a	-	Fatty, waxy, pungent
72	2,3,5-Trimethylpyrazine	22.64	1.03	1401	a	Key-aroma marker	Earthy
<b>73</b>	<b><math>\alpha</math>-Thujone</b>	<b>23.19</b>	<b>1.79</b>	<b>1414</b>	<b>a</b>	<b>ISTD</b>	<b>-</b>
74	2-Octanol	23.49	1.07	1422	a	-	Mushroom, fatty, creamy
75	Ethyl octanoate	23.89	2.00	1431	a	-	-
76	1-Octen-3-ol	23.94	0.64	1433	a	-	Mould, earthy
77	Acetic acid	23.99	0.57	1434	a	Key-aroma marker	Sour, vinegary
78	2-Ethyl-3,6-dimethylpyrazine	24.29	1.20	1441	a	Technological marker	Earthy
79	Furfural	24.88	0.69	1455	a	Technological marker	Sweet, bread-like
80	1-Acetyloxy-2-propanone	24.89	1.31	1455	b	Technological marker	-
81	2-Ethyl-3,5-dimethylpyrazine	24.94	1.17	1457	a	Key-aroma marker	Earthy
82	<i>Trans</i> -linalool oxide	25.29	1.21	1465	a	-	Sweet floral, citrus, fruity
83	2,6-Dimethyl-4-heptanol	25.44	1.40	1469	b	-	-
84	Tetramethylpyrazine	25.54	1.14	1471	a	Technological marker	Earthy
85	2,3-Butanediol diacetate	25.99	1.14	1482	b	-	-
86	2-Ethyl-1-hexanol	26.04	0.97	1483	a	-	-
87	Decanal	26.54	1.86	1495	a	-	Penetrating, sweet, waxy
88	2-Acetylfuran	26.64	0.72	1498	b	-	-
89	3,5-Diethyl-2-methylpyrazine	27.09	0.55	1509	b	Key-aroma marker	Earthy
90	Benzaldehyde	27.34	0.79	1515	a	-	Almond, burnt sugar
91	2,3-Butanediol diacetate	27.49	1.10	1519	b	-	-
92	Furfuryl acetate	27.79	0.79	1526	a	-	-



93	2-Nonanol	27.83	0.94	1527	a	-	-
94	2,3-Butanediol	27.99	0.55	1531	a	-	-
95	Propanoic acid	27.99	0.40	1531	a	-	Fruity, pungent
96	Linalool	28.44	1.00	1542	a	-	Citrus
97	1-Octanol	28.84	0.93	1552	a	-	Moss, nut, mushroom
98	2-Methylpropanoic acid	29.09	0.48	1558	b	Key-aroma marker	Rancid
99	2,3-Butanediol	29.44	0.52	1567	a	-	-
100	Dihydro-2(3H)-furanone	31.34	0.76	1614	b	-	-
101	Butanoic acid	31.54	0.48	1619	a	Key-aroma marker	Sweaty, rancid
102	Phenylacetaldehyde	32.04	0.83	1632	a	Key-aroma marker	Honey-like
103	Ethyl decanoate	32.14	2.31	1635	a	-	Fruity
104	Acetophenone	32.34	0.86	1640	a	-	-
105	2-Furanmethanol	32.59	0.52	1646	a	Technological marker	Burned
106	Ethyl benzoate	33.04	1.49	1658	a	-	-
107	3-Methylbutanoic acid	33.09	0.52	1659	b	Key-aroma marker	Rancid
108	Dodecanal	34.94	2.01	1707	a	-	Fatty, citrus-like
109	Pentanoic acid	35.94	0.46	1734	a	-	Sweaty
110	4-Ethylphenyl acetate	37.39	1.03	1773	b	-	-
111	4-Methylpentanoic acid	38.19	0.52	1795	b	-	-
112	1-Phenylethanol	38.24	0.63	1796	b	-	-
113	2-Phenylethyl acetate	38.39	0.62	1800	b	Key-aroma marker	Flowery
114	Hexanoic acid	39.69	0.52	1837	a	-	Rancid
115	Ethyl dodecanoate	39.74	2.48	1838	a	-	-
116	Guaiacol	39.84	0.75	1841	a	-	Spicy
117	2-Methyl propyl benzoate	40.19	1.21	1851	b	-	-
118	Benzyl alcohol	40.44	0.59	1858	a	-	Sweet, fruity
119	Phenylethylalcohol	41.64	0.66	1892	a	Key-aroma marker	Honey-like
120	(E)-2-Phenyl-2-butenal	42.44	0.97	1915	b	Technological marker	-
121	Acetyl pyrrole	43.64	0.59	1950	a	-	Popcorn-like
122	Phenol	44.74	0.52	1982	a	-	-
123	1H-Pyrrole-2-carboxaldehyde	45.34	0.52	2000	a	-	-
124	4-Hydroxy-2,5-dimethyl-3(2H)-furanone	45.64	0.59	2009	a	Technological marker	Caramel-like
125	Octanoic acid	46.94	0.55	2049	a	-	Sweaty
126	5-Methyl-2-phenyl-2-(Z)-hexenal	47.24	1.21	2058	b	-	-
127	Nonanoic acid	50.34	0.57	2156	a	-	Sweaty, waxy
128	2,3-Dihydro-3,5-dihydroxy-6-methyl-4H-pyran-4-one	52.74	1.41	2231	b	-	-
129	Decanoic acid	53.49	0.66	2259	a	-	Soap-like, fatty
130	2-Phenylacetic acid	59.44	0.79	2549	b	-	Honey-like

<sup>a</sup>: targets identified by means of authentic standards

<sup>b</sup>: targets tentatively identified on MS fragmentation patterns and Linear Retention Indices available in commercial libraries

## **Associated content**

**Supplementary Table 1 (ST1):** Samples characteristics

**Supplementary Table 2 (ST2):** Validation data. Repeatability and intermediate precision on retention times and 2D peaks quantitative descriptors (Normalized 2D Volumes).

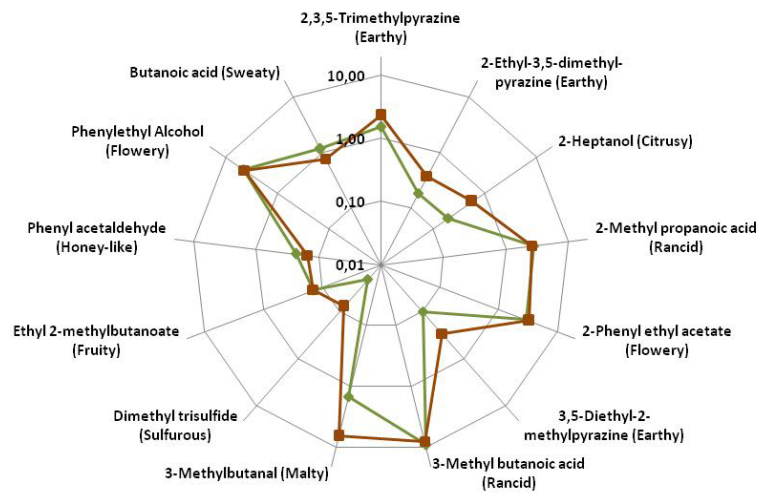
**Supplementary Figure 1 (SF1):** 2D patterns of volatiles from cocoa samples harvested in the Chontalpa region (Tabasco - Mexico) from raw (SF1A), to roasted (SF1B) than steamed (SF1C) and at the end to nibs (SF1D). Light blue circles indicate the positions of targeted peaks, pink the untargeted and yellow circles the ISTDs peaks ( $\alpha$ - and  $\beta$ -thujones).

Figure 1

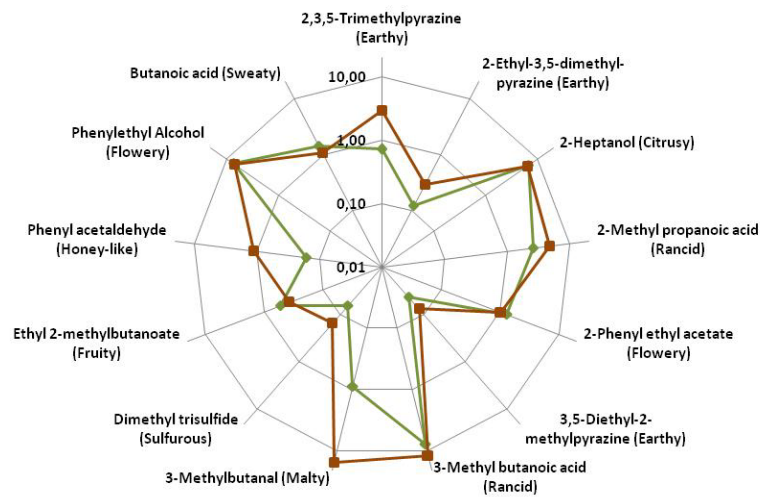


**Figure 2**

**A**  
Chontalpa (Trinitario)  
Raw vs Roasted



**B**  
Venezuela (Trinitario)  
Raw vs Roasted



**C**  
Sao Tomè (Forastero)  
Raw vs Roasted

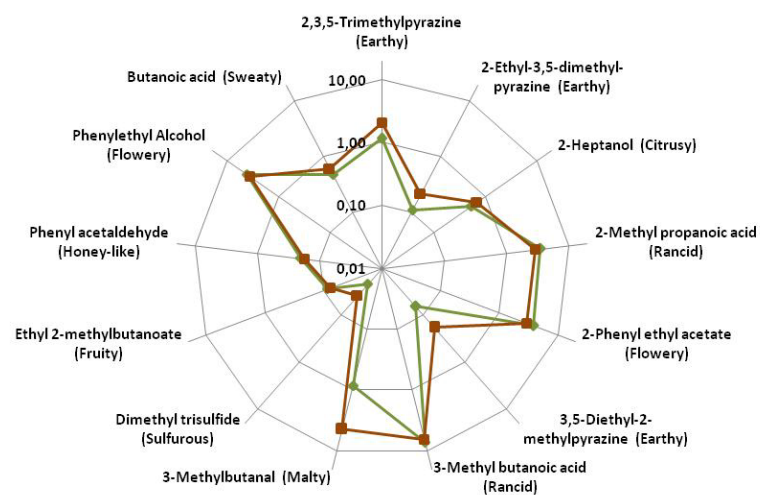
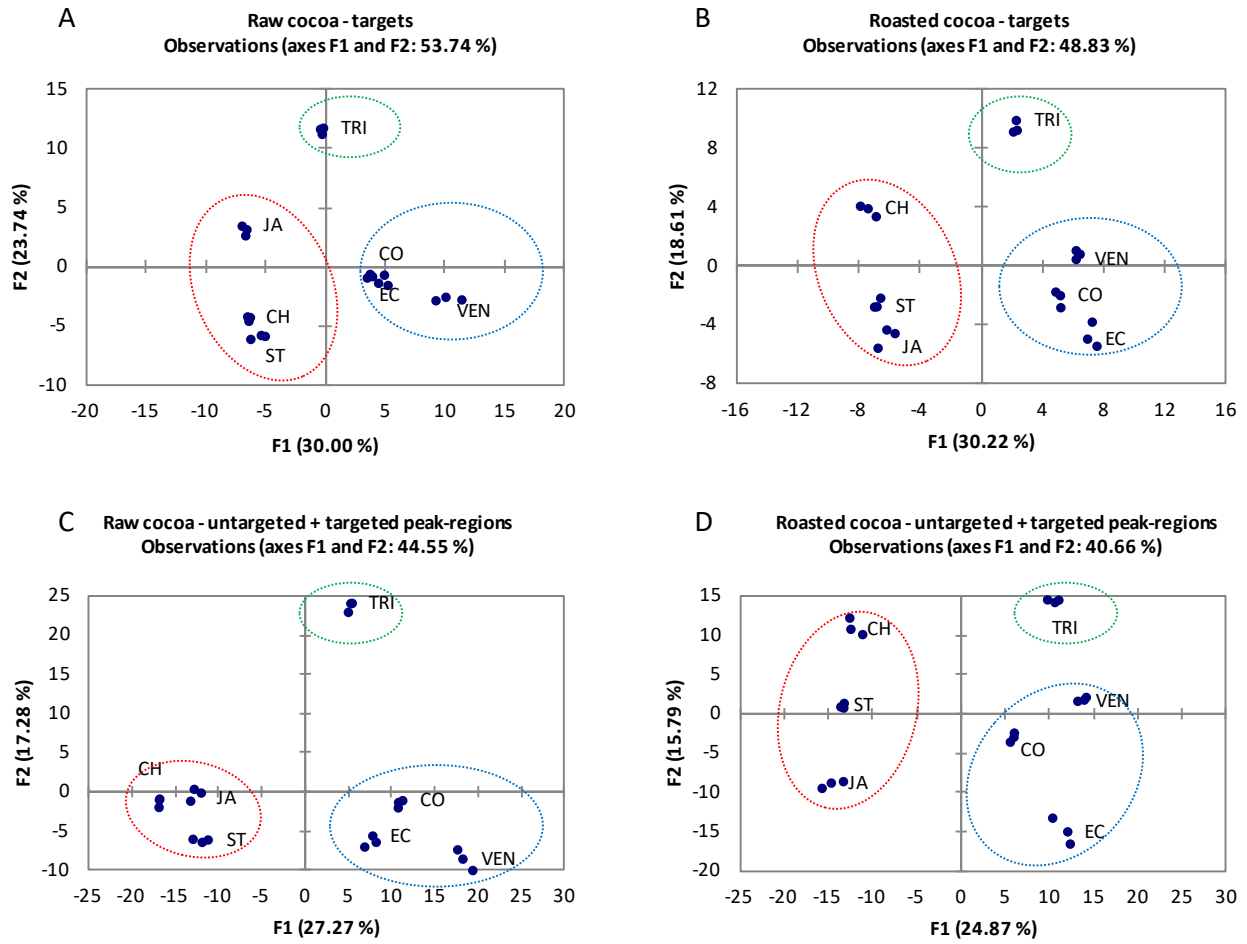


Figure 3



**Figure 4**

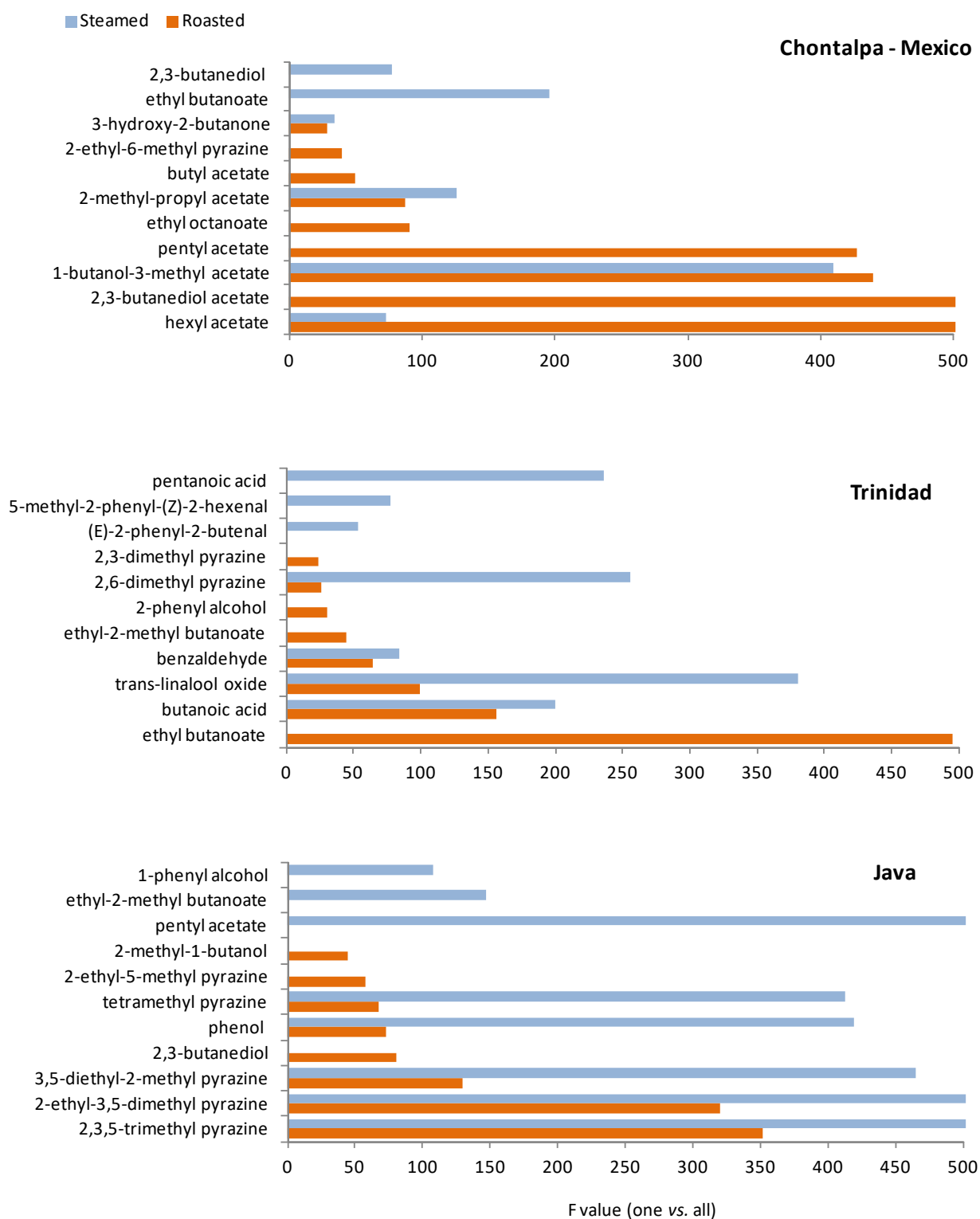
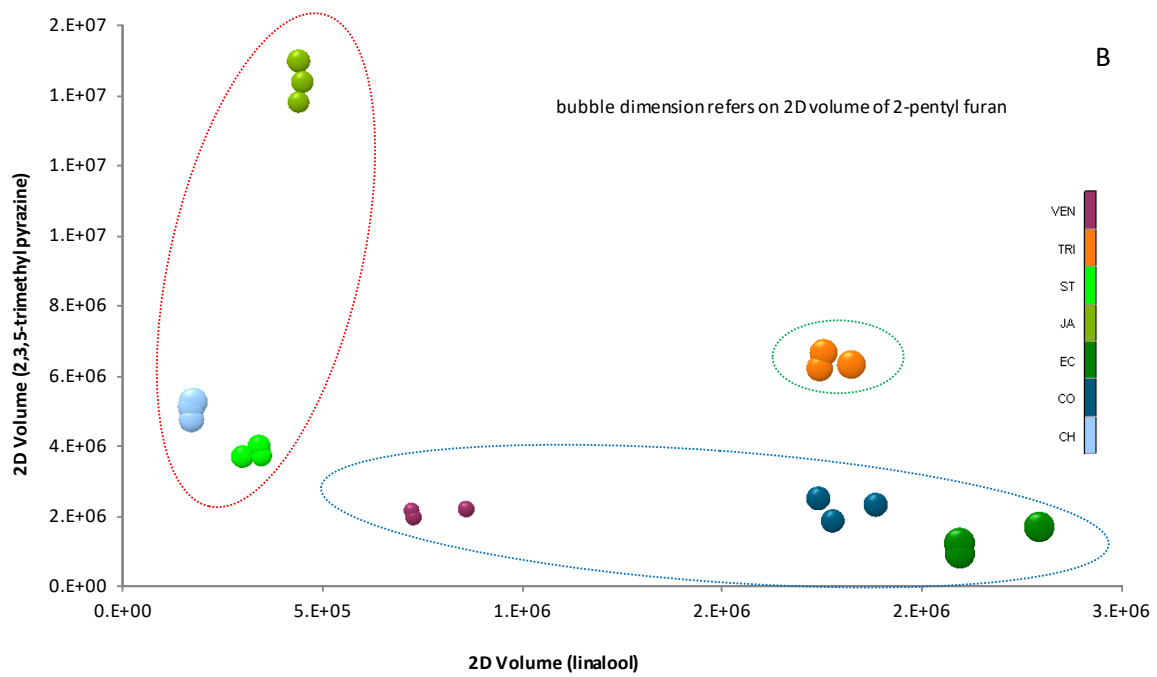
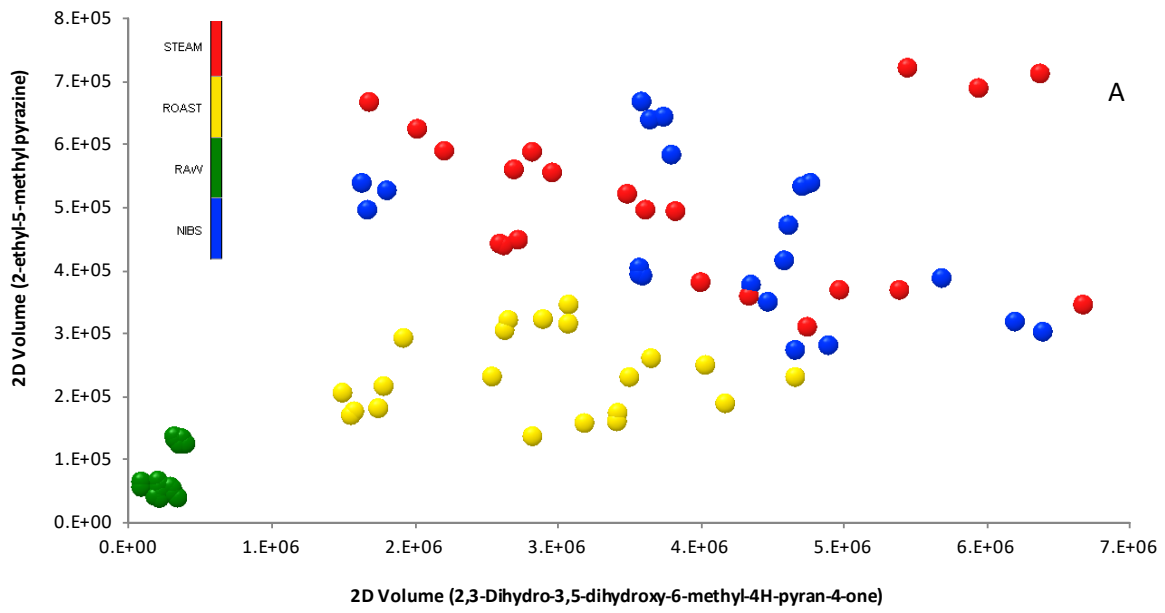


Figure 5



TOC graphics

