

# UHPLC-MS/MS analysis of cocoa bean proteomes from four different genotypes

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Scollo, E., Neville, D. C. A., Oruna-Concha, M. J., Trotin, M. and Cramer, R. (2020) UHPLC-MS/MS analysis of cocoa bean proteomes from four different genotypes. Food Chemistry, 303. 125244. ISSN 0308-8146 doi: https://doi.org/10.1016/j.foodchem.2019.125244 Available at http://centaur.reading.ac.uk/85765/

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To link to this article DOI: http://dx.doi.org/10.1016/j.foodchem.2019.125244

Publisher: Elsevier

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1	UHPLC-MS/MS analysis of cocoa bean proteomes from four
2	different genotypes
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18	Running title: Comparative UHPLC-MS/MS analysis of cocoa bean proteomes
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20	Nonstandard abbreviations: emPAI (exponentially modified protein abundance
21	index); FDR (false discovery rate); BSA (bovine serum albumin)
22	
23	Keywords: Theobroma cacao, cocoa beans, plant proteomics, storage proteins,
24	cocoa bean proteome, cocoa flavour

#### 25 Abstract

26 In this study the proteomic profiles of cocoa beans from four genotypes with different 27 flavour profiles were analysed by bottom-up label-free UHPLC-MS/MS. From a total of 430 28 identified proteins, 61 proteins were found significantly differentially expressed among the 29 four cocoa genotypes analysed with a fold change of  $\geq 2$ . PCA analysis allowed clear 30 separation of the genotypes based on their proteomic profiles. Genotype-specific 31 abundances were recorded for proteases involved in the degradation of storage proteins 32 and release of flavour precursors. Different genotype-specific levels of other enzymes, 33 which generate volatiles compounds that could potentially lead to flavour-inducing 34 compounds, were also detected. Overall, this study shows that UHPLC-MS/MS data can 35 differentiate cocoa bean varieties.

#### 36 **1. Introduction**

Chocolate as commonly sold and consumed is made from the beans of the cocoa tree *Theobroma cacao* (family *Sterculiacae*). This tree is native to the Amazon and Orinoco
valley and requires hot and humid weather conditions to grow. Traditionally, *Theobroma cacao* has been divided into three main genetic groups that are of commercial interest,
Forastero, Criollo and Trinitario, the latter being a hybrid of the first two genetic groups.

42 Most of the cocoa beans produced in the world comes from Forastero varieties, which are 43 considered "bulk in trade" (Lima, Almeida, Nout, & Zwietering, 2011). Two other cultivars 44 have also been described: Amelonado, which is considered a subvariety of Forastero and 45 mainly cultivated in West Africa, and Nacional, a cultivar native to Ecuador. However, this 46 general classification is broad as hybridisation has occurred over time, which has given 47 rise to differentiation within the same genetic groups, especially Forastero.

A study carried out by Motamayor *et al* (Motamayor, Lachenaud, da Silva e Mota, Loor, Kuhn, Brown, et al., 2008) has resulted in the identification of ten genetically distinct clusters. Based on these results the Forastero group has been differentiated into eight subvarieties: Amelonado, Contamana, Curaray, Guiana, Iquitos, Maranon, Nanay, and Purus. This new classification has not affected the Criollo and Nacional varieties, as they have maintained their original terms. The term Trinitario is commonly used to describe hybrids of Forastero and Criollo.

55 Cocoa beans from Trinitario and Criollo generate the "fine cocoa flavour" and make up 56 only 5% of cocoa's total worldwide production (Lima, Almeida, Nout, & Zwietering, 2011). 57 Ivory Coast is the main cocoa-producing country in the world with a total worldwide 58 production share of 42%, followed by Ghana and Nigeria whose shares are 19% and 7%, 59 respectively ("Quarterly Bulletin of Cocoa Statistic," 2017).

60 In general, shortly after harvest cocoa beans undergo natural fermentation which results in the release of free peptides and amino acids from storage proteins (Voigt, Biehl, 61 62 Heinrichs, Kamaruddin, Marsoner, & Hugi, 1994). These compounds are important flavour 63 precursors which contribute to the generation of cocoa aroma during roasting, and that is 64 why poorly fermented beans have a low amount of flavour precursors and do not generate 65 the typical cocoa aroma upon roasting. The autolysis of storage proteins extracted from 66 unfermented cocoa beans generate flavour precursors which produce the typical cocoa 67 aroma when roasted in the presence of the cocoa butter and reducing sugars (Voigt, Biehl, Heinrichs, Kamaruddin, Marsoner, & Hugi, 1994). The LC-ESI MS/MS analysis of these 68 69 autolysis products revealed the presence of mainly hydrophilic peptides whose sequence 70 could be linked to cocoa globulins (Voigt, Janek, Textoris-Taube, Niewienda, & 71 Wostemeyer, 2016). The proteome of Theobroma cacao beans has recently been 72 characterised using ultrahigh performance liquid chromatography (UHPLC) coupled to 73 electrospray ionisation (ESI) tandem mass spectrometry (MS/MS) (Scollo, Neville, Oruna-74 Concha, Trotin, & Cramer, 2018). The highest proportion of the identified proteins could 75 be linked to 'metabolism and energy' and 'proteins and synthesis' functions (Scollo, 76 Neville, Oruna-Concha, Trotin, & Cramer, 2018). The most abundant proteins were 77 albumin and vicilins (Scollo, Neville, Oruna-Concha, Trotin, & Cramer, 2018).

The proteomic profile of cocoa beans during development was previously evaluated by LC-ESI MS/MS using a 'bottom-up shotgun' approach (Wang, Nagele, Doerfler, Fragner, Chaturvedi, Nukarinen, et al., 2016). Cell division, ATP synthesis, RNA processing, amino acid synthesis and activation, protein synthesis, sucrose transportation and degradationassociated proteins were upregulated in young beans compared to mature beans (Wang, et al., 2016). Proteins involved in defence and stress were present at a higher level in mature beans (Wang, et al., 2016).

85 The proteomic profiles of non-fermented cocoa beans from various origins and varieties have been characterised by 2D gel electrophoresis and subsequent analysis by MALDI-TOF 86 87 MS/MS from a total of 49 2D gel spots (Kumari, Grimbs, D'Souza, Verma, Corno, Kuhnert, et al., 2018). The authors reported differences in terms of numbers and intensities of 88 89 proteins between samples from different origins, and samples from the same varieties 90 grown in different countries (Kumari, et al., 2018). According to the authors a vicilin 91 subunit was specific to samples of CCN51 hybrids and the German Forastero variety CD03 92 (Kumari, et al., 2018). Two protein gel spots which revealed a degraded 17-kDa albumin 93 subunit and an internal 15-kDa vicilin subunit showed significant differences among the 94 samples analysed when selecting the geographical origin and protein intensities as 95 variables in a MANOVA analysis (Kumari, et al., 2018). The authors stated that these

- 96 proteins could be used as markers to assess the geographical origin and variety (Kumari,97 et al., 2018).
- Although there are cocoa varieties with different flavour characteristics, it is not fully
  understood whether there is a link between the proteomic profile and flavour development.
  In this work a UHPLC-MS/MS bottom-up label-free approach was employed to characterise
  qualitative and quantitative differences in the proteomic profiles of cocoa beans from four
- 102 genotypes, which show differences in both genetic background and flavour characteristics.

# 103 **2. Materials and methods**

#### 104 **2.1. Chemicals**

Petroleum ether 40-60 was obtained from Fisher Scientific, Loughborough, UK. All other
chemicals were obtained from Sigma-Aldrich, Gillingham, UK, except where stated
otherwise.

# 108 2.2. Plant materials

109 Cocoa beans were from four different genotypes of *Theobroma cacao*, namely ICS 1, ICS 110 39, SCA 6 and IMC 67 harvested at the Cocoa Research Centre of the University of West 111 Indies, St. Augustine, Trinidad, see Supplementary Table 1. Cocoa pods were harvested 112 from 6 different trees for each genotype. A total of six pods were harvested from each 113 tree. Each pod of the same cocoa genotype was considered a biological replicate within 114 the specified cocoa genotype.

115 Pods were stored refrigerated for no longer than 3 days after being harvested. The beans 116 were removed from the pods and the pulp manually removed with the aid of a scalpel. 117 Depulped beans were stored at -20° C and subsequently freeze-dried for 24 hours. 118 Following the freeze-drying step, the beans were stored at -20° C prior to shipping. The 119 freeze-dried beans were air-freighted without temperature control to the University of 120 Reading, UK. The shipment took less than 96 hours. Upon arrival, the beans were stored 121 at -20° C prior to analysis. To obtain a representative sample for each cocoa genotype, 122 approximately 2 g of beans from each biological replicate within the same genotype were 123 combined, and the remainder of the beans were stored in their original container.

#### 124 **2.3. Fat and polyphenols removal**

The freeze-dried beans were snap-frozen using liquid nitrogen and subsequently ground using a mortar and pestle. Fat from aliquots of approximately 160 mg were extracted with 3.5 ml of petroleum ether (boiling point 40-60° C) for 20 minutes in a vertical shaker. The suspensions were subsequently centrifuged at 3100 g for 5 minutes and the supernatants were discarded. The extraction was repeated twice and the precipitates were dried under a stream of nitrogen.

131 In order to prevent the formation of polyphenol-protein complexes during extraction (Voigt, Biehl, & Wazir, 1993), polyphenols were removed following a slight modification of 132 a published method (Voigt, Wrann, Heinrichs, & Biehl, 1994). In brief, polyphenols were 133 134 extracted from the defatted samples with 3.5 ml of a solution made up of cold (~4° C) aqueous acetone (80%; v/v), containing 5 mM sodium ascorbate. The suspensions were 135 136 vortexed for 1 minute and centrifuged at 3100 g for 10 minutes at 4° C. The supernatant 137 was discarded and the extraction repeated twice. Residual water was removed by 138 extraction with 3.5 ml of cold acetone. The sample was then dried under a stream of 139 nitrogen, resulting in acetone-dried powder (ACDP).

# 140 **2.4. Protein extractions and Bradford assay**

Proteins from the ACDP were extracted with 3.5 ml of a solution consisting of aqeous 7 M urea, 2 M thiourea and 20 mM dithiothreitol. The suspensions were placed on a vortexer for 1 minute and subsequently extracted for 1 hour at room temperature in a vertical shaker at 700 rpm. The suspension was subsequently centrifuged at 3100 g for 10 minutes at 20° C. The supernatant was removed and stored at -80° C prior to analysis. The protein concentration in each sample was assessed with the Bradford assay [4]. Bovine serum albumin (BSA) was used as reference standard for quantitation purposes.

# 148 **2.5. Trypsin digestion**

149 Aliquots of proteins extracts (35-47 µl) containing approximately 160 µg of proteins based 150 on the Bradford assay were transferred into 0.5-ml microcentrifuge tubes and spiked with 151 30 µl of an aqueous 10 mg/l BSA solution. A volume of 20 µl of an aqueous 200-mM 152 dithiothreitol (DTT) solution was then added to each tube, and the final concentration of 153 DTT was adjusted to 10 mM by adding 290 µl of 77 mM ammonium bicarbonate. The 154 solutions were incubated for 30 minutes at 37° C. A volume of 43 µl of an aqueous 200-155 mM iodoacetamide (IAA) solution was then added to each sample solution in order to 156 obtain a final IAA concentration of 20 mM. By adding small aliquots of a 2-M urea solution 157 the samples were adjusted to a final urea concentration of 0.6-0.7 M. To each sample 158 tube, a volume of 20  $\mu$ l of a 0.15- $\mu$ g/ $\mu$ l trypsin (Promega, Southampton, UK) solution was 159 added to obtain a 1:50 trypsin-to-protein ratio, and the solutions were incubated for approximately 16 hours at 37° C. After incubation the digestion was stopped by lowering 160 161 the pH to below 3 with the addition of 20  $\mu$ l of a 5% (v/v) solution of aqueous trifluoroacetic 162 acid (TFA) to each sample tube.

# 163 **2.6. Desalting of tryptic digests**

164 The tryptic digest solutions were desalted with SOLAµ HRP 96 well plate 2 mg sorbent 165 mass SPE cartridges (Thermo Scientific, Waltham, MA USA). The cartridges were initially 166 conditioned with 0.2 ml of methanol and subsequently equilibrated with 0.2 ml of 0.2% 167 (v/v) TFA in 50 mM ammonium bicarbonate. After loading the sample solutions, the 168 cartridges were washed with 0.2 ml of 0.2% TFA in water:methanol 97:3 (v/v), and then 169 eluted with 3x 25  $\mu$ l of 0.2% TFA in acetonitrile:water 50:50 (v/v) solution. The SPE 170 eluates were diluted with 0.225 ml of 0.1% TFA in water and stored at -80° C prior to 171 UHPLC-MS/MS analysis.

## 172 2.7. UHPLC-MS/MS analysis of tryptic digests

173 The desalted tryptic digests were analysed on a UHPLC-ESI MS/MS system consisting of 174 an Orbitrap Q Exactive (Thermo Scientific) mass spectrometer coupled to a Dionex 175 Ultimate 3000 (Thermo Scientific) UHPLC system. The injection volume was 15 µl. The 176 UHPLC system was kept at 50° C and the column configuration included an Acquity Peptide 177 CSH C18 150 mm × 0.1 mm ID, 1.7 µm particle size analytical column (Waters, Elstree, 178 UK). The chromatographic separation of the digests was carried out under a linear gradient 179 elution using 0.1 % (v/v) formic acid in water as mobile phase A and 0.1 % (v/v) formic 180 acid in acetonitrile as mobile phase B with a flow rate of 0.1 ml/min. The gradient 181 conditions were as follows: 2% B at 0-5 minutes, 30% B at 80 minutes, 60% B at 90 182 minutes, 90% B at 100-110 minutes, 2% B at 115-125 minutes. MS analysis was carried 183 out in positive ion mode using the Orbitrap mass analyser, setting its resolution at 70,000 184 and its AGC (acquisition gain control) target at 1,000,000 with a maximum injection time 185 of 200 ms. The MS scan covered a range between m/z 200 and 2400. For MS/MS analysis 186 a data dependent experiment selecting the 10 most abundant precursor ions was 187 performed, using the quadrupole mass analyser as the initial filter, and setting the isolation 188 window width to m/z 2.0. For this experiment, the resolution of the Orbitrap was set to 189 17,500 with an AGC target of 20,000. The injection time for MS/MS acquisition was set to 190 300 ms. Fragmentation was performed by collision-induced dissociation (CID) with a 191 normalised collision energy of 28%. Dynamic exclusion was enabled, setting the filter to 192 15 seconds. The threshold for triggering a data-dependent scan was set to '100,000' and 193 only ions with a charge state between 2 and 5 were selected.

## 194 2.8. Data analysis

195 All MS/MS spectra were processed using Mascot Distiller software (Matrix Science Ltd, 196 London, UK; Version 2.5.1.0) in order to convert the raw UHPLC-MS/MS data into peak 197 lists suitable for database searching using the Mascot search routine. For the evaluation 198 of the effect of harvest time and tree, Mascot Server Version 2.4.1 was used, while the 199 analysis of the different cocoa genotypes was carried out employing the Mascot Server 200 Version 2.6 (Matrix Science Ltd). Mascot searches were carried out against the Cacao 201 Matina 1-6 Theobroma Genome v1.1 database cacao (http://www.cacaogenomedb.org/Tcacao\_genome\_v1.1#tripal\_analysis-downloads-box; 202

203 downloaded on 31st May 2015; 59,577 sequences; 23,720,084 residues), and a custom-204 made contaminants database (70 sequences; 31,845 residues). Searches were performed 205 using the following parameters: peptide mass tolerance, 10 ppm; MS/MS tolerance, 0.3 206 Da; peptide charge, +2, +3, +4; missed cleavages, 2; fixed modification, 207 carbamidomethyl (C); variable modification, oxidation (M) and acetyl (N); enzyme, 208 trypsin. The false discovery rate (FDR) for all searches was adjusted to 1%, which resulted 209 in various significance thresholds for the different searches. However, the p-value was 210 <0.05 for all searches. The amino acid sequence of BSA was added to the Theobroma 211 cacao database. Functional annotation was carried out by matching the proteins' accession 212 codes from the Cacao Matina 1-6 Genome v1.1 Theobroma cacao to the GoMapMan 213 database (<u>http://protein.gomapman.org</u>). For each entry the highest hierarchical 214 classification was used in this study. Label-free quantitation was carried out using replicate protocol with Mascot Distiller software. Normalisation of the proteins' intensities was 215 216 carried out against BSA. Protein quantitation was performed by employing the median of 217 the ion signal intensity ratios from all peptide for each protein, for which a minimum of 218 two peptides were detected. For statistical analyses, JMP Pro 13.0 and XLSTAT 2108.5 219 software were used.

# 220 3. Results

A total of four different genotypes (ICS 1, ICS 39, IMC 67 and SCA 6) were evaluated in this project. These cocoa genotypes were carefully selected in order to include varieties with differences in both genetic background and flavour profiles. The list of the selected cocoa genotypes with their genetic backgrounds and flavour profiles is shown in Supplementary Table 1.

In order to minimise variability of the protein expression due to external factors, the investigated cocoa varieties were grown in the ICGT (International Cocoa Genebank Trinidad) field in Trinidad under controlled conditions in terms of water intake, fertilisation, and soil structure. However, to evaluate the effect of a different location on the proteomic profile of cocoa beans, trees from the genotype IMC 67 were also grown in a different field called "Campus" located within 5 Km off the ICGT field.

# 232 **3.1. Effect of harvest time and different trees**

To evaluate the effect of different trees on the proteomic profile of cocoa beans, four pods of the cocoa genotype IMC 67 were harvested on the same day, one from each of four different trees grown in the ICGT field. Three preparative replicates were prepared for each of these four biological replicates, and each preparative replicate was analysed by a single UHPLC-MS/MS run. UHPLC-MS/MS reproducibility was previously checked and constantly monitored by quality control samples of the same standard cocoa bean protein extract analysed alongside the preparative replicates. For each quantified protein, the mean of the intensities in the three preparative replicates for each biological replicate (preparative sample mean) was calculated, and subsequently the average of the preparative sample means of the four biological replicates was calculated (overall mean). For each biological replicate the fold increase/decrease from the overall mean expressed as the ratio between the sample mean and the overall mean was calculated.

245 A total of 511 proteins were detected in the four biological replicates and only six proteins S-adenosyl-L-methionine-dependent 246 (Thecc1EG042578t1: methyltransferases; 247 Thecc1EG025391t1: beta-amylase 6; Thecc1EG000326t1: salicylate 0-248 Thecc1EG026589t1: methyltransferase; eukaryotic aspartyl protease; 249 Thecc1EG027146t1: HSP20-like chaperones; Thecc1EG041163t1: glycosyl hydrolase 250 family protein) showed a fold increase/decrease from the overall mean of >2 in at least 251 one biological replicate, while none showed any increase/decrease from the overall mean 252 of >2.7. Of these six proteins only beta-amylase 6 showed this increase/decrease for two 253 biological replicates, while the other five proteins showed this differential abundance for 254 exactly one biological replicate, covering the entire set of the biological replicates.

The harvest time in this study covered a period of six months. Therefore, to evaluate the effect of harvest time on the proteomic profile of cocoa beans, four pods from the same tree (genotype IMC 67; grown in the ICGT field) but harvested at different times (20<sup>th</sup> Dec 2016, 21<sup>st</sup> Feb 2017, 23<sup>rd</sup> March 2017, 17<sup>th</sup> May 2017) were analysed. As before three preparative replicates were prepared for each of the four biological replicates, and each preparative replicate was analysed by a single UHPLC-MS/MS run.

261 A total of 502 proteins were detected in the four biological replicates analysed. Among 262 these proteins, only nine entries showed a fold increase/decrease from the overall mean 263 of >2 in at least one biological replicate (see Supplementary Table 2). In this case, two 264 proteins (Thecc1EG042149t1: serine carboxypeptidase-like 48; Thecc1EG047098t1: 265 uncharacterised) fluctuated far more than any protein in the tree comparison experiment 266 with a fold increase/decrease of >3 and up to 11. The number of proteins with a fold 267 increase/decrease from the overall mean of >2 in each biological replicate was between 268 three and five.

# **3.2. Investigation of proteome changes dependent on the genotype and field**

Cocoa pods were harvested from six different trees for each genotype grown in the ICGT field and for the six IMC 67 trees that were grown in the Campus field. A total of six pods were collected from each tree. Pooled samples containing an equal amount of all of the biological replicates from the same cocoa variety (Campus and ICGT grown IMC 67 pods were pooled separately) were prepared as described in the Materials and Methods section.

275 To evaluate the proteome changes, which are dependent on the genotype, a UHPLC-276 MS/MS label-free proteomic analysis was carried out on each of the cocoa genotypes. A 277 total of four preparative replicates were prepared for each genotype sample, and each 278 preparative replicate was analysed by UHPLC-MS/MS. A reference sample was prepared 279 by combining equal aliquots of all 20 preparative replicates. The Distiller software 280 calculated the ratios of the intensities of the proteins in each preparative replicate against 281 the same proteins in the reference sample. Only proteins which were identified and 282 quantified in at least three preparative replicates of a cocoa genotype were selected for 283 comparative label-free quantitative proteomic analysis. With this requirement a total of 284 430 proteins were identified and quantified (see Supplementary Table 3). The mean of the 285 ratios for the preparative replicates of the same cocoa genotype was calculated for each 286 quantified protein. The fold differences between the cocoa genotypes are reported as the 287 ratio of the highest mean versus the lowest mean for each quantified protein.

288 Almost all of the 430 proteins were detected in all genotypes apart from a 60S acidic 289 ribosomal protein (accession number Thecc1EG005040t1) that was not detected in the 290 genotype SCA 6. However, the abundance of this protein was not significantly different in 291 the other genotypes. From all other identified and quantified proteins, a total of 61 proteins 292 showed a significant fold difference of >2 (p-value <0.05) within at least one pairwise 293 comparison among the four cocoa genotypes. Among these proteins, those with a sum of 294 the four sample-to-reference ion signal ratios that is outside the range of 75-125% from 295 the theoretical value of 4 were further evaluated to assess their peptide ion signal 296 intensities. In this case a total of four proteins showed a signal too weak for reliable 297 quantitation, and therefore these proteins were not further investigate. A list of the 298 differentially expressed proteins, which showed an acceptable ion signal intensity, 299 including their biological process and function, is provided in Table 1. A graphical 300 representation of the proteins' classification based on their biological processes and 301 functions is provided in Figure 1. Biological processes, for which only one protein was 302 identified and quantified, are labelled as "Others" in Figure 1.

**Table 1.** List of differentially abundant proteins with a fold difference of >2, obtained from the four cocoa genotypes analysed by label-free 304 LC-MS/MS.

ID	Accession	Description	Biological process	Function	ICS 1	ICS 39	IMC 67	SCA 6	Fold diff.
1	Thecc1EG029400t1	N-terminal nucleophile aminohydrolases	Protein degradation	ME	4.80	4.02	0.70	0.69	6.95
2	Thecc1EG029392t1	Glutathione S-transferase family protein	Gluthatione S-transferase	ME	0.47	0.48	2.46	0.85	5.28
3	Thecc1EG025391t1	Beta-amylase 6	Carbohydrate metabolism	ME	0.49	1.30	1.14	2.39	4.90
4	Thecc1EG017184t1	Sulfite oxidase	S-assimilation	ME	0.96	4.08	2.12	1.06	4.26
5	Thecc1EG038258t1	Molybdenum cofactor sulfurase	Co-factor and vitamin metabolism	ME	0.78	1.46	0.92	0.35	4.12
6	Thecc1EG030320t1	Ethylene-forming enzyme	Hormone metabolism	ME	0.44	1.79	0.50	1.10	4.03
7	Thecc1EG021639t1	PEBP	Unspecified biological process	UN	1.72	2.22	1.22	0.66	3.36
8	Thecc1EG020604t1	Primary amine oxidase	Oxidase	ME	1.89	0.78	1.27	2.54	3.25
9	Thecc1EG047098t1	Uncharacterized protein	Unspecified biological process	UN	0.93	0.89	0.61	1.94	3.18
10	Thecc1EG036433t1	HSP20-like chaperones protein	Stress	DFS	1.19	0.38	1.11	1.00	3.15
11	Thecc1EG026543t1	Lipoxygenase 1	Hormone metabolism	ME	0.47	0.45	1.23	1.37	3.01
12	Thecc1EG026589t1	Eukaryotic aspartyl protease	Protein degradation	ME	1.69	1.67	0.58	1.10	2.91
13	Thecc1EG042578t1	S-adenosyl-L-methionine-dependent methyltransferases protein	Hormone metabolism	ME	0.66	1.53	0.54	1.53	2.85
14	Thecc1EG019372t1	Bifunctional inhibitor/lipid-transfer protein/seed storage 2S albumin protein	Protease inhibitor/seed protein/lipid transfer	SP	1.91	1.43	2.33	0.84	2.78
15	Thecc1EG027146t1	HSP20-like chaperones protein	Stress	DFS	1.46	0.53	0.98	0.72	2.77
16	Thecc1EG026193t1	Threonine aldolase 1	Amino acid metabolism	ME	1.53	0.85	0.58	0.91	2.66
17	Thecc1EG037345t1	17.6 kDa class II heat shock protein	Stress	DFS	1.35	0.51	1.13	0.60	2.65
18	Thecc1EG012673t1	21 kDa seed protein <sup>*</sup>	Stress	DFS	0.83	1.56	0.59	1.16	2.65
19	Thecc1EG025860t2	Uncharacterized protein	Unspecified biological process	UN	1.41	0.88	2.00	0.76	2.64
20	Thecc1EG012662t1	21 kDa seed protein <sup>*</sup>	Stress	DFS	0.79	1.56	0.59	1.16	2.63
21	Thecc1EG038931t1	Xyloglucan endotransglycosylase 6	Cell wall degradation	ME	0.58	0.79	0.59	1.48	2.57
22	Thecc1EG006471t1	Flavin-dependent monooxygenase 1	Oxidase	ME	1.14	0.62	1.57	0.81	2.55
23	Thecc1EG030938t1	Cc-nbs-Irr resistance protein	Unspecified biological process	UN	0.51	1.28	1.22	0.53	2.53
24	Thecc1EG036938t1	Aldolase-type TIM barrel	Nucleotide metabolism	ME	1.95	0.88	0.92	0.79	2.47
25	Thecc1EG006154t1	Glycinamide ribonucleotide synthetase	Nucleotide metabolism	ME	1.06	0.68	0.43	0.84	2.45
26	Thecc1EG041496t1	Stress responsive A/B Barrel Domain	Unspecified biological process	UN	1.27	1.75	0.76	0.72	2.44
27	Thecc1EG030354t1	Fumarylacetoacetase	Amino acid metabolism	ME	1.06	0.88	0.75	1.80	2.40
28	Thecc1EG019909t2	Carrot EP3-3 chitinase	Stress	DFS	0.77	0.52	0.76	1.25	2.39
29	Thecc1EG040975t1	Alpha/beta-Hydrolases protein•	Glucogalamannosidase	ME	0.64	0.81	0.98	1.52	2.37
30	Thecc1EG020603t2	Primary amine oxidase	Oxidase	ME	0.91	0.52	1.02	1.22	2.36
31	Thecc1EG016747t1	Acyl-CoA-binding protein 6	Lipid metabolism	ME	0.94	0.80	0.49	1.13	2.33
32	Thecc1EG022426t1	Thioredoxin protein	Redox	ME	1.31	0.99	0.56	1.25	2.33

33	Thecc1EG008318t1	Aldolase-type TIM barrel	Oligopeptide transport system permease protein	ME	1.26	1.27	0.57	1.32	2.30
34	Thecc1EG022506t1	Monodehydroascorbate reductase seedling isozyme	Redox	ME	1.25	1.03	1.44	0.63	2.28
35	Thecc1EG047057t1	Cystathionine beta-synthase	Unspecified biological process	UN	2.21	1.29	1.78	0.97	2.27
36	Thecc1EG029923t1	Larreatricin hydroxylase	Unspecified biological process	UN	1.36	2.26	1.21	1.02	2.21
37	Thecc1EG000245t1	Serine carboxypeptidase S28	Protein degradation	ME	1.47	3.01	1.37	1.51	2.20
38	Thecc1EG021820t1	Tau class glutathione transferase GSTU45	Gluthatione S-transferase	ME	1.75	0.96	1.08	0.79	2.20
39	Thecc1EG025715t1	Uncharacterized protein	Unspecified biological process	UN	0.87	1.03	0.47	0.92	2.20
40	Thecc1EG043707t1	Anti-oxidant 1	Metal handling	ME	1.05	1.19	0.80	1.76	2.19
41	Thecc1EG016386t1	6-Phosphogluconate dehydrogenase	Oligopeptide transport system permease protein	ME	1.36	0.90	0.62	1.08	2.18
42	Thecc1EG014591t1	Malate synthase glyoxysomal	Gluconeogenesis	ME	2.08	0.96	1.05	1.37	2.17
43	Thecc1EG042584t1	S-adenosyl-L-methionine-dependent methyltransferases protein	Hormone metabolism	ME	0.90	1.34	0.63	1.36	2.16
44	Thecc1EG034339t1	Dehydrin 2	Stress	DFS	1.37	0.91	0.72	0.63	2.16
45	Thecc1EG035433t1	Alcohol dehydrogenase 1 <sup>+</sup>	Fermentation	ME	1.14	0.90	0.63	1.37	2.16
46	Thecc1EG010364t2	Carbonic anhydrase 2 CA2	TCA/organic transformation	ME	1.20	0.98	0.79	0.56	2.13
47	Thecc1EG006694t2	Triosephosphate isomerase	Photosynthesis	ME	1.40	1.35	0.71	1.51	2.11
48	Thecc1EG006498t1	Basic chitinase	Stress	DFS	0.58	0.92	1.15	1.22	2.10
49	Thecc1EG026326t2	Pathogenesis-related protein P2	Stress	DFS	1.33	1.49	0.71	0.87	2.10
50	Thecc1EG029913t1	Alpha/beta-Hydrolases protein•	Glucogalamannosidase	ME	1.55	1.16	0.75	0.81	2.07
51	Thecc1EG036604t1	Secretory laccase	Secondary metabolism	ME	0.75	1.55	1.10	0.83	2.05
52	Thecc1EG015253t1	RNA binding Plectin/S10 domain-containing protein	Protein synthesis	PSP	1.02	1.35	0.72	1.48	2.05
53	Thecc1EG005533t1	Transketolase	Photosynthesis	ME	1.36	1.14	0.81	1.65	2.05
54	Thecc1EG000770t1	Acetamidase/Formamidase	Photosynthesis	ME	2.12	1.52	1.04	1.49	2.03
55	Thecc1EG014683t1	Hydroxysteroid dehydrogenase 1	Dehydrogenase	ME	1.30	0.73	1.08	0.65	2.01
56	Thecc1EG001447t1	Alcohol dehydrogenase 1 <sup>+</sup>	Fermentation	ME	1.32	1.11	0.70	1.41	2.01
57	Thecc1EG001141t1	Lipase/lipooxygenase PLAT/LH2	Unspecified biological process	UN	1.61	1.11	1.02	0.81	2.00

DFS, defence and stress; ME, metabolism and energy; PSP, protein synthesis and processing; SP, storage proteins; UN, unclassified. In the genotype columns the average abundance ratio values relative to the reference sample of the preparative replicates are reported.\*These proteins entries have a 99.5% homology and can therefore be considered to be proteoforms of the same gene. <sup>+</sup>These protein entries have a 87% homology. <sup>•</sup>These protein entries have a 35% homology.



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**Figure 1.** Classification of the differentially abundant proteins listed in Table 1 based on their biological process (upper pie chart) and their function (lower pie chart). 'Others' in the upper pie chart refers to all biological process, for which only one protein was found. The function group labels are as follows: DFS, defence and stress; ME, metabolism and energy; PSP, protein synthesis and processing; SP, storage proteins; UN, unclassified.

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321 For each genotype, the number of proteins listed in Table 1 whose intensity was highest

322 and lowest compared to the other genotypes are graphically represented in a histogram

in Figure 2.

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Figure 2. Number of differentially expressed proteins (fold difference of >2) detected at
 the higher (UP) and lower (DOWN) level for each genotype (see text for further details).

A pairwise comparison between the genotypes for each protein listed in Table 1 was also carried out, using the non-parametric Whitney Mann test to assess the significance of the differential expression (p <0.05). The result of this comparison can be found in Supplementary Table 4.

332 To evaluate whether the proteomic data would allow a graphical differentiation of the four 333 cocoa genotypes analysed, PCA analysis loading the ratios of the differentially expressed 334 proteins listed in Table 1 as variables and the genotypes as observations was performed. 335 In this case the data from all analytical replicates were used. The PCA score plot of the 336 first two components clearly separates the four cocoa genotypes (see Figure 3). Each point 337 in this graph represents a preparative replicate, and the replicates from the same genotype 338 are displayed with the same colour. In order to assess which proteins were positively correlated to each genotype, a PCA loading plot of the differentially expressed proteins 339 340 listed in Table 1 is also shown in Figure 3 (lower plot). Using this plot, variables should be 341 positively correlated to observations which are located in similar regions of the score plot. 342 For instance, the proteins with the ID 2, 19 and 22 are closest to the region in the score 343 plot where the genotype IMC 67 is located and are greatly more abundant in the same 344 genotype.



Figure 3. PCA score plot (upper plot) of the 57 differentially abundant proteins listed in
Table 1. Preparative replicates of the same genotype are displayed with the same colour.
The lower plot shows the PCA loading plot. Each variable is labelled with the corresponding
ID number as listed in Table 1. The blue and yellow oval in the loading plot indicate clusters
related to IMC 67 and SCA 6, respectively.

Comparing the proteomic profiles of the cocoa genotype IMC 67 grown in two different fields allowed the identification and quantitation of 430 proteins in the two biological replicates. Among these proteins, only four proteins were significantly different with a fold change of >2 between the two samples, while a ribosomal protein and a secretory laccase were detected only in the IMC 67 genotype grown in the ICGT field (see Supplementary Table 5). The latter two proteins were detected at low levels while the others had a fold change of <3.4.

385 Data supporting the results of this work are available in the PRIDE (Proteomics
386 Identifications Database) partner repository at the European Bioinformatics Institute,
387 PXD011984 (<u>http://www.ebi.ac.uk/pride/</u>).

# 388 4. Discussion

In the experiments carried out to assess the tree-to-tree, harvest time and field-to-field effects, the same genotype was used and only a few proteins, i.e. less than ten in each case, were detected with a fold increase/decrease of >2. Amongst these only three showed a fold increase/decrease of >3 and two were solely detected in one biological replicate but at a low level. Thus, given the total number of identified and quantified proteins, these results indicate that the variability in the detected proteome is very low between the biological replicates analysed to assess these effects.

396 In contrast, the analysis of the proteomic difference with respect to genotype revealed a 397 high variability with more than 60 proteins showing a significant fold change of >2 for at least one pairwise genotype comparison. The overall highest fold difference in this 398 399 comparison was found for an aminohydrolase (see Table 1). This protein was detected at 400 significantly higher levels in both ICS genotypes, while it was found at much lower 401 abundance in the genotypes IMC 67 and SCA 6. A blast search of the amino acid sequence 402 of this protein returned a 100% match to a 20S proteasome alpha subunit which is part 403 of the N-terminal nucleophile hydrolase superfamily. This class of proteins are involved in 404 the hydrolysis of the amide bonds in either proteins or small molecules (Marchler-Bauer, 405 Bo, Han, He, Lanczycki, Lu, et al., 2017). The active site is the N-terminal amino group which accepts a proton during the hydrolysis activating as a result either the nucleophilic 406 407 hydroxyl in a Ser or Thr residue or the nucleophilic thiol in a Cys residue (Marchler-Bauer, 408 et al., 2017).

The next highest fold change was recorded for a glutathione S-transferase (GST) family protein which was found at a much higher level in the genotype IMC 67 compared to all other genotypes. GST family proteins catalyse the conjugation of a variety of substrates to the reduced form of glutathione and therefore are involved in detoxification processes (Armstrong, 1997).

A 60S acidic ribosomal protein was not detected in any of the preparative replicates of the genotype SCA 6, while it was found in all other genotypes without any significant abundance differences. This class of proteins regulates the translation of mRNA in protein synthesis (Remacha, JimenezDiaz, Santos, Briones, Zambrano, Gabriel, et al., 1995).

With respect to the 57 proteins in Table 1, the highest number of less abundant proteins was found in the genotype IMC 67, and only 5 proteins were detected in this genotype at a higher level compared to the other genotypes (see Figure 2). The highest relative number of more abundant proteins compared to less abundant proteins (19 vs 9) was found for the genotype ICS 1.

423 The PCA score plot of the differentially expressed proteins for the four genotypes analysed 424 shows that the individual genotypes are located in different quadrants of the plot and can 425 be clearly separated from each other (see Figure 3). Both ICS 1 and ICS 39 belong to the same genetic group Trinitario, which originates from hybridizations between Criollo and 426 427 Forastero. Therefore, the positive correlation of these genotypes in the PCA score plot 428 could result from their closer genetic background compared to the other genotypes. IMC 429 67 and SCA 6 are genotypes from the genetically distant varieties Forastero and 430 Contamana, respectively, of which both have a different genetic background from 431 Trinitario (Motamayor, et al., 2008). Therefore, the separation pattern observed on the 432 PCA score plot reflects the differences in genetic background among the four genotypes 433 evaluated. Based on these findings, the PCA score plot of the differentially expressed 434 proteins could be used as a tool to differentiate cocoa genotypes.

435 Loading the differentially expressed proteins as variables on a PCA loading plot allows a 436 graphical visualisation of the proteins positively correlated to each genotype. The majority 437 of the proteins more abundant in IMC 67 and SCA 6 form respective clusters in the bottom 438 left and bottom right corner of the PCA loading plot (see Figure 3, lower plot), reflecting 439 the separation of these genotypes observable in the PCA score plot. The genotypes ICS 1 440 and ICS 39 show a high degree of correlation in the PCA score plot. Therefore, the proteins 441 found at a higher level in each of these genotypes cannot be separated in the PCA loading 442 plot and form a single large cluster located at the top centre of the PCA loading plot. The 443 location of this cluster is consistent with the position of these genotypes in the PCA score 444 plot.

The highest number of differentially expressed proteins can be associated to metabolism and energy. This function class generally encompasses the majority of the proteins expressed in cocoa beans as shown in a previous study (Scollo, Neville, Oruna-Concha, Trotin, & Cramer, 2018), and includes two primary amine oxidases (ID 30 and ID 8 in Table 1) and two alcohol dehydrogenases identifications (ID 45 and ID 56 in Table 1), of which the latter are highly homologous (87% homology). Primary amine oxidases catalyse

451 the oxidation of alkylamines to aldehydes with the release of ammonia and hydrogen 452 peroxide (Conklin, Prough, & Bhatanagar, 2007), while alcohol dehydrogenases catalyse 453 the oxidation of primary and secondary alcohols to the corresponding aldehydes and 454 ketones (Svensson, Hoog, Schneider, & Sandalova, 2000). It has been reported that both 455 aldehydes and ketones are formed during roasting of fermented cocoa beans as a result 456 of the Maillard reaction and Strecker degradation, and both classes of compounds 457 contribute to the cocoa flavour (Aprotosoaie, Luca, & Miron, 2016). These reactions are 458 endothermic as they require high temperatures to be activated and are not catalysed by 459 enzymes. In theory, aldehydes and ketones could also be produced from oxidation of 460 amines and alcohols during fermentation catalysed by amine oxidases and alcohol 461 dehydrogenases. However, it is not known whether these enzymes are activated during 462 this process, and whether there is a relation between their concentration and the 463 generation of cocoa flavour. The primary amine oxidase (ID 30) was significantly more 464 abundant in the genotype SCA 6 compared to ICS 39, while the other primary amine 465 oxidase was significantly higher in the genotype SCA 6 versus ICS 39, and in the genotype 466 ICS 1 versus ICS 39. Both alcohol dehydrogenase identifications IDs 45 and 56 were 467 significantly more expressed in the genotype SCA 6 compared to IMC 67, reflecting their 468 high homology and indicating that two proteoforms of the same gene were detected.

469 A total of 9 proteins involved in stress response were differentially expressed. Four of 470 these proteins (ID 10, 15, and 17 in Table 1) were heat shock proteins by name which are 471 linked to the response of the plant to stress conditions (Al-Whaibi, 2011). There are no 472 significant differences in the abundances of these proteins in ICS 1 versus IMC 67 and SCA 473 6 versus IMC 67 but they were significantly more abundant in ICS 1 compared to ICS 39. 474 Both these genotypes belong to the Trinitario variety which is originally from Trinidad and 475 includes all hybridisation combinations of the Criollo and Forastero varieties. Criollo 476 varieties are more susceptible to disease and adverse environmental factors. The genotype 477 ICS 39 has a stronger Criollo ancestry compared to ICS 1, which could explain why heat 478 shock proteins are more abundant in ICS 1 compared to ICS 39.

479 A eukaryotic aspartyl protease (ID 12 in Table 1) was significantly more abundant in the 480 genotypes ICS 1 and ICS 39 compared to IMC 67 (fold difference of 2.9). Eukaryotic 481 aspartyl protease is a cocoa endogenous protease which has an optimum pH of around 482 3.8 and is active during early stage of fermentation, cleaving internal peptides bonds with 483 the release of mainly hydrophobic peptides (Voigt, Biehl, Heinrichs, Kamaruddin, 484 Marsoner, & Hugi, 1994). The abundance of this protease was not consistent in the 485 biological replicates of IMC 67 harvested from different trees on the same day. Therefore, 486 the low amount found in the pooled sample may be due to natural variations amongst 487 biological tree replicates.

488 A serine carboxypeptidase (ID 37 in Table 1) was detected at a significant higher level in 489 ICS 39 compared to the other genotypes. Carboxypeptidase is an exopeptidase which 490 cleaves off C-terminal amino acids from mainly hydrophobic oligopeptides formed by the 491 action of aspartyl protease during fermentation with the preferential release of 492 hydrophobic amino acids and hydrophilic peptides (Bytof, Biehl, Heinrichs, & Voigt, 1995). 493 These compounds are important flavour precursors which react with sugars during roasting 494 to form volatiles compounds which contribute to the cocoa aroma. A higher amount of 495 aspartyl protease and carboxypeptidase could result in an increase in the generation of 496 flavour precursors during fermentation, which could lead to changes in the flavour profiles 497 of roasted cocoa beans.

498 A beta-amylase was detected at a significantly higher level in the genotype SCA 6 compared to ICS 1 and IMC 67 (ID 3 in Table 1). Beta-amylases are part of the glycoside 499 500 hydrolase family, which are a group of enzymes catalysing the cleavage of the glycosidic 501 bond in polysaccharides with release of maltose units (Rejzek, Stevenson, Southard, 502 Stanley, Denyer, Smith, et al., 2011). This disaccharide can react with nitro compounds 503 such as amino acids and peptides during roasting through the Maillard reaction which 504 results in the generation of volatile compounds (Kramholler, Pischetsrieder, & Severin, 505 1993). Therefore, the release of maltose can be affected by the levels of beta-amylase 506 present in cocoa beans, which in turn could have an effect on the flavour profile of roasted 507 cocoa beans. However, the abundance of this specific beta-amylase was not consistent in 508 the biological replicates of IMC 67 harvested from different trees on the same day. 509 Therefore, the low amount found in the pooled sample may be due to natural variations 510 amongst biological tree replicates.

511 Two 21-kDa seed albumin identifications were obtained at a significant higher level in the 512 genotype ICS 39 compared to IMC 67 (ID 18 and 20 in Table 1). These albumins are 513 storage proteins with endopeptidase inhibitor activity, which contain 219 amino acids 514 residues, and can be considered to originate from the same gene as they are 99.5% 515 homologous. The main 21-kDa seed albumin in cocoa beans is a protein with 221 residues 516 which shares a homology of 80% with the albumins ID 19 and 21 listed in Table 1. The 517 221-residues albumin was not differentially expressed in the cocoa genotypes analysed 518 (see Supplementary Table 3). LC-MS/MS identification of free peptides released from this 519 protein during fermentation have been reported by several authors (Caligiani, Marseglia, 520 Prandi, Palla, & Sforza, 2016; D'Souza, Grimbs, Grimbs, Behrends, & Corno, 2018; 521 Marseglia, Sforza, Faccini, Bencivenni, Palla, & Caligiani, 2014). However, so far there is 522 no evidence that the 219-residues albumins are also degraded during this process. As a 523 result, the shorter chain albumins may not play a role in the generation of cocoa flavour.

A 2S albumin was significantly more abundant in the genotypes ICS 1 and IMC 67 compared to SCA 6 (see ID 14 in Table 1). This albumin is a seed storage protein with protease inhibitor activity which is also involved in the transfer of phospholipids and fatty acids through the cell membrane (Kader, 1996). Degradation of this protein during fermentation has not been reported in the literature.

529 In addition to cocoa endogenous enzymes, the amount of pulp in the cocoa pod and the 530 surrounding microflora can also play a role in the generation of cocoa flavour.

# 531 **5. Conclusions**

532 This work has shown that UHPLC-MS/MS can be employed to characterise qualitative and 533 quantitative differences in the proteomic profiles of cocoa beans from various genotypes. 534 The PCA analysis has allowed separation of the cocoa genotypes from different varieties 535 and has shown a correlation between close genotypes and their genetic background. Using 536 this approach, it was possible to graphically visualise proteins positively correlated with 537 each genotype, and assess which proteins contribute most to the separation of the 538 genotypes in the PCA plot. This methodology could be employed as a platform to build 539 larger datasets of proteins which could allow traceability of cocoa beans from different 540 varieties. Proteases which degrade storage proteins during fermentation with the release 541 of flavour precursors have been found differentially expressed in some of the genotypes 542 analysed. Changes in the amount of these proteases could be related to variation in the 543 flavour profiles of cocoa varieties. Different genotype-specific levels of other enzymes that 544 could potentially lead to flavour-inducing compounds have also been detected. Thus, 545 further experiments could be performed to assess whether the different amounts of these 546 enzymes, present during fermentation, affect the final flavour profiles obtained

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# 548 Acknowledgements

549 The authors are grateful to the University of West Indies Cocoa Research Centre for 550 providing the cocoa bean samples. This work has been financially supported by Mondelēz 551 International.

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