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1 **Advanced characterization of polyphenols from *Myrciaria jaboticaba* peel and**
2 **lipid protection in *in vitro* gastrointestinal digestion**

3

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11

12 **Abstract**

13 Ultrasound-assisted and solvent extractions led to similar levels in hydrolyzable tannins (10.3-6.0
14 mg/g), anthocyanins (7.8-10.2 mg/g) and flavonols (0.24-0.32 mg/g) for dried *Myrciaria jaboticaba*
15 peel (DJP). Ultrasound was efficient for the extraction of poorly soluble hydrolyzable tannins but
16 affected the stability of anthocyanins and flavonols. UPLC-DAD-MSⁿ allowed the identification of
17 44 hydrolyzable tannins as single and mixed hexosides bearing galloyl, HHDP and tergalloyl units.
18 Twelve mixed HHDP-galloylgluconic acids and tergalloylated hexosides were newly discovered in
19 this work. Acid hydrolysis of both ultrasonic extract and DJP yielded five major compounds, *i.e.*
20 gallic acid, ellagic acid, gallic acid-C-hexoside, valoneic acid dilactone and sanguisorbic acid
21 dilactone and pointed to higher contents in hydrolyzable tannins than by summing individual
22 polyphenols after UPLC. Last, cyanidin-3-*O*-glucoside and hydrolyzable tannins from the ultrasonic
23 extract inhibited lipid peroxidation of a Western type meal in *in vitro* digestion, suggesting a health
24 benefit for these jaboticaba polyphenols.

25

26

27 **Key Words:** anthocyanins, hydrolyzable tannins, lipid oxidation, *in vitro* digestion,
28 bioaccessibility, ultrasound-assisted extraction

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32 1. Introduction

33 Jabuticaba (*Myrciaria jaboticaba* (Vell.) O. Berg, also known as *Plinia cauliflora* (Mart.)
34 Kausel), is a typical Brazilian berry from the *Myrtaceae* family. Fruits, presenting a dark violet peel
35 and a white gelatinous pulp at maturity, are consumed *in natura* but also largely after processing.
36 Because of its stiffness and astringent taste, the peel (up to 35% of the fruit weight) is not consumed
37 leading to wastes rich in phenolic compounds, mainly anthocyanins, flavonols, and hydrolyzable
38 tannins (ellagitannins, gallotannins) (Neves, Stringheta, Gomez-Alonso, & Hermosin-Gutierrez,
39 2018; Morales et al., 2016; Plaza et al., 2016; Pereira, Barbosa, da Silva, Ferri, & Santos 2017; Wu,
40 Dastmalchi, Long, & Kennelly, 2012). These studies suggest the contribution of galloyl and
41 hexahydroxydiphenoyl (HHDP) groups to jabuticaba hydrolyzable tannins although full
42 identification and quantification of these tannins remain elusive owing to a large structural diversity
43 and a lack of standards. Anthocyanins are present in high concentrations (0.14-3.2 g/100 g dry
44 peel), mainly as cyanidin and delphinidin glucosides (Albuquerque et al., 2020; Quatrin et al., 2019;
45 Plaza et al., 2016; Alezandro, Dube, Desjardins, Lajolo, & Genovese, 2013; Peixoto et al., 2016).
46 Phenolic compounds appear to contribute to the *in vitro* antioxidant, antimicrobial, anti-proliferative
47 and anti-inflammatory properties described for jabuticaba extracts (Albuquerque et al., 2020; Leite-
48 Legatti et al., 2012). Moreover, jabuticaba fruit, tea and peel improve plasma lipid profile and
49 insulin sensitivity, reduce lipid peroxidation in plasma and brain, and increase antioxidant enzyme
50 activity in various organs of animal models for diabetes and obesity (Lenquiste et al., 2019;
51 Alezandro, Granato & Genovese, 2013). A human study by Plaza et al. (2016) supported the
52 preventive role of jabuticaba peel in metabolic diseases decreasing insulin and glucose levels. Acute
53 and chronic human supplementations with anthocyanin-rich foods and extracts are clearly in favor
54 of a health benefit in vascular protection (Fairlie-Jones et al., 2017). Of further interest,
55 cardiometabolic risk biomarkers such as LDL-cholesterol, blood pressure or flow-mediated-dilation
56 responded differently to ellagitannin-rich foods (pomegranate and nuts) and anthocyanin-containing
57 products (berries and grape products) (Garcia-Conesa et al., 2018). Oxidative stress is a key factor

58 of inflammation and antioxidant phenolic compounds are expected to interfere with the production
59 of reactive oxygen species even in locations such as the artery wall.

60 Only anthocyanin glucosides are bioavailable and the gastric mucosa was found to play an
61 important role in absorption (Peixoto et al., 2016). Other anthocyanin glycosides and ellagitannins
62 are not absorbed in the upper gastrointestinal tract and are further metabolized in the colon by the
63 microbial flora. Ellagitannins mainly yield urolithins (Quatrin et al., 2020) while anthocyanins are
64 cleaved to small phenolic acids such as protocatechuic acid and phloroglucinaldehyde (de Ferrars et
65 al., 2014). The resulting microbial metabolites are partly deoxygenated and subjected to phase II
66 metabolism resulting in a partial masking of residual hydroxyl groups. Bioavailable forms of
67 anthocyanins and ellagitannins are thus less likely to play an antioxidant role in the circulation
68 compared to their native forms.

69 As a matter of fact, the gastrointestinal tract has been proposed as a major site for diet-related
70 oxidative stress and antioxidant activity of plant polyphenols (Gobert et al., 2014). Red meat iron,
71 as provided by the Western diet, can trigger the oxidation of dietary polyunsaturated lipids leading
72 to the formation of 4-hydroxyalkenals. These reactive lipid oxidation products are absorbable and
73 further recovered in lipoproteins. LDL covalent modification is a key event in atherogenesis.
74 Besides, the presence of cytotoxic and genotoxic 4-hydroxynonenal (4-HNE) in feces and plasma
75 was correlated to the development of colonic preneoplastic lesions and atherosclerotic plaques in
76 rodents fed with diets enriched in heme iron and polyunsaturated lipids. Interestingly, polyphenol-
77 rich red wine and pomegranate extracts were found to decrease heme-induced luminal peroxidation
78 and the promotion of colonic lesions (Bastide et al., 2017). On the other hand, both apple puree and
79 extract inhibited the formation of 4-HNE in feces and plasma and largely reduced aortic plaque
80 formation (Bolea et al., 2021).

81 Because anthocyanins display a large spectrum of coloring properties, they are nowadays widely
82 used as natural food additives in dairy products and beverages. Ultrasound-assisted extraction of the
83 jabuticaba peel and further extract stabilization by encapsulation are currently investigated to take

84 advantage of polyphenol nutritional and sensory properties (Tarone, da Silva, Cazarin & Marostica
85 Junior, 2021). Besides, peel extracts were shown to display antioxidant and antimicrobial activities
86 in sausages (Baldin et al., 2016).

87 The first objective of this work was to assess the efficiency of high-intensity ultrasound for
88 polyphenol extraction from dried jaboticaba peel by comparison with solvent extraction using full
89 structural elucidation and quantification by UPLC/DAD/ESI-MSⁿ. Next, the ultrasonic extract was
90 compared to pure cyanidin-3-*O*-glucoside for its capacity to inhibit lipid oxidation in the *in vitro*
91 gastrointestinal digestion of a Western type diet. Anthocyanin recovery was further evaluated.

92

93 **2. Materials and Methods**

94 **2.1. Materials**

95 Cyanidin-3-*O*-glucoside (C3G), gallic acid, ellagic acid and quercetin 3-*O*-glucoside were
96 purchased from Extrasynthese (Genay, France). Methanol, acetonitrile, hexane and 2-propanol were
97 HPLC-MS grade from Fisher Scientific (Illkirch, France); formic acid was HPLC-MS grade from
98 Merck (Darmstadt, Germany). Ultrapure water (resistivity 18.2 M Ω /cm at 25 °C) was obtained with
99 a Millipore OPak 2 (Millipore Corporation, Bedford, MA, USA).

100 Horse heart myoglobin (M1882, type II), pepsin from porcine gastric mucosa (P6887, 2829
101 U/mg according to Minekus et al. (2014)), porcine pancreatin (P7545, trypsin activity 2.8 U/mg
102 according to Minekus et al. (2014)), porcine bile extract (B8631) and 2,4-dinitrophenylhydrazine
103 (DNPH) were purchased from Sigma-Aldrich (Saint-Quentin Fallavier, France). 4-Hydroxy-2-
104 nonenal and 4-hydroxy-2-nonenal-D3 were purchased from Bertin Pharma (Montigny le
105 Bretonneux, France). Commercial sunflower oil, from Auchan (lot No. A07611), was stored at -20
106 °C after purchase. L- α -phosphatidylcholine from dried egg yolk (PL) (P3556) was from Fluka
107 (Buchs, Switzerland). Oil and PL compositions are detailed in the Supplementary Material.

108

109 **2.2. Jaboticaba peel extracts**

110 *2.2.1. Jaboticaba peel processing*

111 Jaboticaba (*Myrciaria jaboticaba* (Vell.) O. Berg) fruits were kindly donated by “Indústria e
112 Comércio Lagoa Branca Ltda”, located at Boa Vista II Farm, in White House city (São Paulo,
113 Brazil) under the authorization of the Brazilian Management System of National Genetic Heritage
114 and Traditional Associated Knowledge (#A72354F) (August 2016). The species used in this work
115 was assessed by comparison with species deposited in the herbarium of the Agronomic Institute of
116 Campinas, Santa Elisa farm, numbers 48093 and 48094. Twenty kg of fresh fruits were washed,
117 manually peeled and the peel was dried in a stove with air circulation (Marconi, Piracicaba, SP,
118 Brazil) at 40 °C for 72 h until constant weight. The dried peel (ca. 1 kg) was transformed in a fine
119 powder with an electrical mill (Marconi, MA 630/1, Piracicaba, SP, Brazil) and sifted (mesh 20).
120 The dried jaboticaba peel powder (DJPP) was packed in plastic bags under vacuum and stored at -
121 20 °C until use.

122

123 *2.2.2. Dried jaboticaba peel extraction by ultrasound (DJP-US)*

124 DJPP prepared as above (1 g) was added into a falcon with 25 mL of 50% ethanol in
125 ultrapure water (v/v) and sonicated using a 13-mm ultrasonic probe (Unique, Desruptor, 800 W, 19
126 kHz, Indaiatuba, Brazil) at an ultrasound intensity of 3.7 W/cm² for 3 min. The probe contact height
127 within the dispersion was 40 mm and an ice bath was used (temp. < 35 °C). The solvent was
128 removed in vacuum at 37 °C. The extraction was made in triplicate. The dry residues were
129 reconstituted in the same volume of ultrapure water before freeze-drying and storage at - 20 °C.

130

131 *2.2.3. Dried jaboticaba peel extraction by a conventional solvent method (DJP-CM)*

132 In a 2 mL microtube with conical bottom, 100 mg (for analysis of phenolic compounds) or
133 10 mg (for anthocyanin analysis) of DJPP were placed with balls of zirconium and 1.5 mL of
134 extraction solvent (70% ethanol in ultrapure water (v/v) containing 1% of formic acid). After

135 shaking at 1200 rpm for 45 s in a 1600 MiniG[®] SPEX Sample Prep (New Jersey, USA), the
136 microtube was centrifuged at 16000 g and 4 °C for 5 min. The supernatant was collected and the
137 residue was extracted again twice in the same conditions. Supernatants were pooled, the solvent was
138 evaporated in vacuum at 37 °C and the sample was stored at - 20 °C until analysis. The extraction
139 was made in triplicate.

140

141 2.2.4. *Hydrolysis of phenolic compounds*

142 The hydrolysis of the phenolic extracts was conducted according to Garcia-Villalba et al.
143 (2015) with some modifications. DJP-US or DJPP (100 mg) was put into a glass tube with 3.34 mL
144 of ultrapure water and 1.66 mL of 37% HCl. The tube was vortexed for 1 min and incubated at 90
145 °C for 24 h. After cooling to room temperature, the pH was adjusted to 2.5 with 5 M NaOH and the
146 volume adjusted to 6 mL with water in a graduated syringe. After centrifugation (5 min, 16000 g
147 and 4 °C), the supernatant was recovered and filtered through a Minisart RC4 filter before injection
148 onto UPLC/DAD/ESI-MSⁿ. The residual pellet was vortexed twice with 1 mL of DMSO/MeOH
149 (50:50, v/v) for 2 min and centrifuged as above. The mixed supernatants were filtered through a
150 Minisart RC4 filter before analysis. All samples were prepared in triplicate.

151

152 **2.3 Identification and quantification of phenolic compounds and anthocyanins**

153 Freshly reconstituted solutions of DJP-US and DJP-CM extracts were prepared in 98 vol. of
154 1% formic acid in ultrapure water (v/v) and 2 vol. of 1% formic acid in acetonitrile (v/v) at
155 concentrations of 100 mg/mL and 1 mg/mL for UPLC/DAD/ESI-MSⁿ analysis of phenolic
156 compounds and anthocyanins, respectively. Separation and mass conditions are detailed in the
157 Supplementary Material.

158 The quantification was performed using 5 to 10 point-calibration curves with C3G,
159 quercetin-3-*O*-glucoside, gallic acid and ellagic acid. All the standards were prepared in methanol
160 except C3G in methanol acidified with 0.1% HCl (v/v). Gallic acid was quantified with its own

161 standard and gallic acid derivatives were quantified as gallic acid equivalent at 280 nm; ellagic acid
162 was quantified with its own standard and ellagic acid derivatives as ellagic acid equivalent at 370
163 nm; quercetin derivatives and myricetin derivatives as quercetin-3-*O*-glucoside equivalent at 370
164 nm, and anthocyanins as C3G equivalent at 520 nm.

165

166 **2.4 Simulated static *in vitro* gastrointestinal digestion**

167 *2.4.1 Preparation of the simulated digestion fluids and antioxidants*

168 The simulated gastric (SGF) and intestinal (SIF) fluids were made up as described in
169 Minekus et al. (2014). Enzymes were prepared prior to use. Porcine pepsin was prepared as an 8.08
170 mg/mL solution in SGF for a final concentration in the gastric phase of 1000 U/mL. Porcine
171 pancreatin was prepared as a 94 mg/mL solution in SIF for a final concentration in the intestinal
172 phase of 100 U trypsin/mL. The porcine bile extract was prepared as a 20.85 mg/mL solution in SIF
173 for a final concentration of 5 mg/mL (ca. 10 mM) in bile salts. CaCl₂ (H₂O)₂ (0.1 mM), HCl (0.1 M)
174 and NaOH (0.25 M) were prepared in ultrapure water. DJP-US (1 mg or 5 mg) was dissolved prior
175 to use in 1 mL of SGF to bring respectively 14.88 µg (US-1) or 74.4 µg (US-5) of total
176 anthocyanins with 89% of C3G and 11% of delphinidin-3-*O*-glucoside (D3G). C3G-1 and C3G-5
177 were prepared by adding 21 and 104 µL of C3G at 0.859 mg/mL in methanol acidified with 0.1%
178 HCl (v/v) to 1979 and 1896 µL of SGF, respectively.

179

180 *2.4.2 Preparation of oil-in water emulsions*

181 The physical state of lipids during digestion was simulated by a 12.5% oil-in-water emulsion
182 stabilized by egg yolk L- α -phosphatidylcholine (PL). In a 60 mL short-necked glass bottle, 5 g of
183 sunflower oil were added to 100 mg of PL previously dispersed in 35 mL of SGF under magnetic
184 stirring. The mixed oil and aqueous phases were then homogenized using a rotor stator
185 homogenizer (Silent Crusher M-01, Heidolph) at 24000 rpm for 2 min. The resulting coarse

186 emulsion was sonicated on ice for 8 periods of 30 s with rest intervals of 30 s and an amplitude of
187 60% (Q700, QSonica, 20 kHz). See Suppl. Material for emulsion characterization.

188

189 2.4.3 *Lipid oxidation in in vitro gastrointestinal digestion*

190 The physicochemical conditions for *in vitro* static gastrointestinal digestion were from
191 Minekus et al. (2014) with slight modifications. First, 12 mL of the fine emulsion were placed in a
192 50 mL round-bottom flask containing a magnetic stirrer. For the gastric phase, 1 mL of pepsin, 50
193 μL of 0.1 mM CaCl_2 , 1 mL of antioxidant solution (US-1, US-5, C3G-1 or C3G-5) or 1 mL of SGF
194 for the blank were added. If necessary, the pH was adjusted to 5 with 0.1 M HCl or 0.25 M NaOH.
195 The first sample was collected (G0). Lipid oxidation was initiated by adding 2.5 mL of 200 μM
196 MbFe^{III} prepared in SGF (using $\epsilon = 7700 \text{ L}\cdot\text{mol}^{-1}\cdot\text{cm}^{-1}$ at 525 nm (Bolea et al., 2019) to reach a final
197 concentration of 30 μM . Bottom flasks were protected by punched parafilm and incubated in an
198 oven at 37 °C under stirring at 280 rpm during 60 min with sampling at 30 and 60 min. After 60
199 min, the pH was adjusted to 3 with 0.1 M HCl and the digestion was carried on another 60 min
200 (sampling at 90 and 120 min). After 2 hours of gastric digestion, pancreatin (6.5 mL), bile (6 mL),
201 and CaCl_2 (75 μl) were then added to simulate the intestinal phase. pH was adjusted to 6.5 with 0.25
202 M NaOH and sampling continued for the next 2 h. Experiments were run at least in triplicate.

203

204 2.4.4 *Determination of lipid oxidation products*

205 Digesta samples (200 μL) were diluted in a microtube with 1000 μL of a 2-propanol/hexane
206 (2:3, v/v) mixture, vortexed and centrifuged (10 min, 16 200 g, 4 °C). The upper hexane phase (ca.
207 600 μL) was collected and 1200 μL of hexane were added to the lower phase, vortexed and
208 centrifuged as above. Pooled hexane phases were evaporated under nitrogen and taken up in 2 mL
209 of 2-propanol to yield extract S1. S1 was immediately used for measurement of conjugated dienes
210 (CD) then stored at -20 °C until the determination of 4-hydroxy-2-nonenal (4-HNE).

211

212 2.4.4.1 Measurement of lipid-derived conjugated dienes (CD)

213 After further dilution of S1 (300 μL) in 2-propanol (1700 μL), the concentration in CD was
214 determined by measuring the absorbance at 234 nm (HP 8453 diode-array spectrometer) using a
215 molar absorption coefficient of 27 000 $\text{M}^{-1}\text{cm}^{-1}$ for conjugated linoleyl hydroperoxides.

216

217 2.4.4.2 Measurement of 4-hydroxy-2-nonenal

218 The secondary lipid oxidation product 4-HNE was derivatized with DNPH before
219 quantification in the MRM mode in HPLC-MS (See Supplementary Material).

220

221 2.4.5 *Determination of anthocyanin bioaccessibility*

222 Anthocyanin bioaccessibility was defined as the content in free C3G and D3G in the aqueous
223 phase of the digesta. Digesta samples (600 μL) were collected and placed in a microtube with 50 or
224 75 μL of 0.1 M HCl for gastric and intestinal samples, respectively. After centrifugation (16 200 g
225 for 10 min at 4 $^{\circ}\text{C}$), the aqueous phase was removed via syringe, filtered (Phenex RC 0.45 μm) and
226 stored at - 20 $^{\circ}\text{C}$ until anthocyanin analysis by UPLC/DAD/ESI-MSⁿ as described in section 2.3.

227

228 **2.5 Statistical analyses**

229 All the results are expressed as mean \pm standard deviation (SD). One-way analysis of variance
230 (ANOVA) with repeated measures was performed to test the effect of variation factors. If
231 significant effects were found at a 95% confidence level, ANOVA was followed by a Tukey-
232 Kramer post hoc test to identify differences among groups (XLStat software (2020), Addinsoft,
233 Paris, France).

234

235 3. Results and discussion

236 3.1. Profile and content in phenolic compounds and anthocyanins of dried jaboticaba 237 peel by UPLC/DAD/ESI-MSⁿ analyses

238 In order to assess the effect of ultrasound on polyphenol extraction, the extracts produced by
239 ultrasound-assisted extraction and conventional solvent extraction of dried jaboticaba peel were
240 compared through their profiles in phenolic compounds and anthocyanins (Fig. 1 & Fig. S1 in
241 Suppl. Mat.). For both extracts, the same 58 phenolic compounds were tentatively identified (Table
242 1). Maximum absorption wavelength, molecular ion and fragmentation pattern in MS were used in
243 the absence of standards for structure assessment.

244

245 3.1.1. Hydrolyzable tannins

246 Hydrolyzable tannins are polyesters between a sugar moiety and hydroxylated benzoic acid
247 derivatives. In the DJP-US and DJP-CM extracts, gallic acid and ellagic acid were the main
248 contributors. Gallic acid was not found free but esterified by a hexose unit which could be glucose
249 according to Pereira et al. (2017). Galloylhexose (**1**) displays a parent ion at m/z 331 with major
250 fragment ions at m/z 271 and 169, which are typical of galloylglucose fragmentation (Quatrin et al.,
251 2019). Di- (**4**, **22**; m/z 483), tri- (**34**; m/z 635), tetra- (**44**, **46**; m/z 787) and pentagalloylhexoses (**50**;
252 m/z 939) were recovered as several isomers, as found in jaboticaba (Quatrin et al., 2019) and in
253 pomegranate (Garcia-Villalba et al., 2015).

254 Ellagic acid was found free (**42**; m/z 301) and bound to a hexose moiety (Pereira et al., 2017;
255 Plaza et al., 2016; Morales et al., 2016; Fischer, Carle, & Kammerer, 2011). Ellagic acid pentoside
256 (**39**) displays a parent ion at m/z 433 and a major fragment at m/z 301 indicating the loss of a
257 pentose unit. The hexahydroxydiphenoyl (HHDP) group related to ellagic acid appears in five
258 isomeric di-HHDP-hexosides (**6**, **9**, **10**, **17**, **40**). Two of them, **10** and **17**, could be quantified. They
259 display a parent ion at m/z 783 and major fragments at m/z 481 and 301 indicating the loss of a
260 HHDP group except **40** which gave a dehydrated form first (m/z 765). Previous studies (Quatrin et

261 al., 2019; Plaza et al., 2016; Wu et al., 2012) reported 2 isomers in jabuticaba peel while Silva et al.
262 (2016) isolated pedunculagin from jabuticaba seed and Garcia-Villalba et al. (2015) found one
263 isomer in pomegranate.

264 A larger group of hydrolyzable tannins is constituted by mixed esters containing both galloyl
265 and HHDP groups. Quantified HHDP-galloylhexose isomers (**2**, **5**, **16**, **25**) display a parent ion at
266 m/z 633 and a major fragment ion at m/z 301 indicating the loss of HHDP. HHDP-digalloylhexose
267 isomers (**21**, **29**, **37**) were evidenced as previously described in *Myrciaria* species (Quatrin et al.,
268 2019; Plaza et al., 2016; Fracassetti, Costa, Moulay, & Tomas-Barberan, 2013). They present a
269 parent ion at m/z 785 and fragment ions at m/z 633 and 483 resulting from the loss of a galloyl or a
270 HHDP group. They were all quantified. For di-HHDP-galloylhexoses (**26**, **27**, **36**), with a parent ion
271 at m/z 935, two different fragmentation patterns are observed. Isomer **26** fragments to yield a major
272 ion at m/z 917 ($-H_2O$) while the two others yielded m/z 633 in agreement with the loss of a HHDP
273 group. Isomeric casuarictin, stachyurin, potentillin and casuarinin have been tentatively identified in
274 jabuticaba (Quatrin et al., 2019; Pereira et al., 2017; Plaza et al., 2016), camu-camu (Fracassetti et
275 al., 2013) and pomegranate (Fischer et al., 2011; Garcia-Villalba, et al., 2015). Casuarictin was
276 formerly identified by NMR in *Myrciaria cauliflora* by Pereira et al. (2017). Fast elution, close
277 retention times but different fragmentation schemes are observed for **26** and **27** suggesting that
278 these compounds could be casuarinin and casuarictin (Plaza et al., 2016). Di-HHDP-
279 digalloylhexose (**55**) displays a parent ion at m/z 1087 with a major fragment ion at m/z 917 (loss of
280 gallic acid) as well as a fragment ion at m/z 749 (loss of hexahydroxydiphenic acid). This
281 compound is tentatively reported for the first time for the jabuticaba species. Finally, HHDP-
282 trigalloylhexose (**41**) is characterized by a parent ion at m/z 937 and displays losses of gallic acid
283 (m/z 767), HHDP (m/z 635), gallic acid + HHDP (m/z 465) and formation of ellagic acid (m/z 301)
284 (Plaza et al., 2016; Fracassetti et al., 2013). They were all quantified except **55**.

285 HHDP-galloylgluconic acid (**3**), HHDP-digalloylgluconic acids (**8**, **12**) and di-HHDP-
286 galloylgluconic acids (**13**, **18**, **19** and **31**) are tentatively reported for the first time in the jabuticaba

287 species although they were previously described Tanaka et al. (1992) in pomegranate and
288 *Lagerstroemia speciosa*. The parent ions at m/z 649, 801 and 951 differ by 16 amu from those
289 found for HHDP-galloylhexose (m/z 633), HHDP-digalloylhexose (m/z 785) and di-HHDP-
290 galloylhexose (m/z 935). Major fragment ions at m/z 605, 757 and 907 indicate the loss of CO₂
291 which is in agreement with the presence of a carboxylic acid function. HHDP (m/z 301) could be
292 observed in MS³ for **3** while the loss of HHDP was observed in the MS² fragmentations of **13**, **18**,
293 **19** and **31** with m/z 605 resulting from the major fragment at m/z 907. They proved to be in smaller
294 amounts compared to related hexose derivatives. Additionally, the fragmentation of compound **31**,
295 proposed to be di-HHDP-galloylgluconic acid, totally differs from that reported for DHHDP-
296 HHDP-galloylglucose in pomegranate (Calani et al., 2013).

297 Isomeric nonahydroxyterphenic acid dilactones (**14** and **30**) were detected as previously
298 found in jabuticaba (Wu et al., 2012) and camu-camu (Fracassetti et al., 2013). They presented a
299 parent ion at m/z 469 and a major fragment ion at m/z 425 (loss of CO₂) and differed remarkably by
300 their retention times. Compound **14** was not further fragmented and it was thus assigned as tergallic
301 acid dilactone owing to likely strong C-C bonds between gallic acid units. Additional fragmentation
302 for **30** revealed fragments at m/z 407, 301 and 167. Fragment m/z 301 corresponds to ellagic acid
303 and is in favor of an additional labile ether linkage between 2 aromatic units, as found for valoneic
304 acid dilactone in pomegranate (Garcia-Villalba et al., 2015). Two compounds (**20** and **23**), in trace
305 amounts, presented a parent ion at m/z 631 and a major fragment ion at m/z 451. The loss of 180
306 amu suggests the presence of a hexose moiety linked through a C- or O-glycosidic linkage to a
307 tergalloyl group as found in jabuticaba (Quatrin et al., 2019) and in hydrolyzed pomegranate
308 (García-Villalba et al., 2015). Pereira et al. (2017) identified 4,6-*O*-tergalloyl-D-glucose
309 (Cauliflorin) by NMR in jabuticaba.

310 Six compounds with a parent ion at m/z 933 were identified although only 3 of them could
311 be quantified. Fragment ions at m/z 451 and 631 are observed for four compounds (**15**, **33**, **38** and
312 **52**) corresponding respectively to the tergalloyl group and the loss of HHDP from the parent ion.

313 This fragmentation pathway is consistent with structures such as vescalagin, castalagin, and α - or β -
314 alnusiins as reported in jabuticaba (Albuquerque et al., 2020; Quatrin et al., 2019). According to
315 Pereira et al. (2017) who secured by NMR experiment the structures of two these compounds,
316 vescalagin was only present in peel of *Myrciaria cauliflora* whereas vescalagin and castalagin were
317 both present in seeds and pulp. The two other compounds (**7**, **11**) eluted faster and display common
318 fragment ions at m/z 915 (- H₂O), 613 and 569, as observed for vescalagin and castalagin standards
319 (Tavares et al., 2016). Standards are thus required for the unequivocal attribution of these different
320 HHDP-tergalloylhexoses.

321 High-molecular weight compounds likely bearing a tergalloyl or a gallagyl group appeared
322 in trace amounts. They are reported for the first time in the jabuticaba species. Isomers **51** and **57**
323 display m/z at 1085 and 542, corresponding to singly and doubly charged ions, respectively, and a
324 fragment ion at m/z 633 which could be HHDP-galloylhexose. Isomers **53** and **56** displayed m/z at
325 1083 and 541 (doubly charged ion) and yielded m/z 631 upon fragmentation as found for
326 tergalloylhexose. It is worth noting that HHDP-galloyltergalloylhexose (MW 1086) exhibits the
327 same molecular weight as digalloylgallagylhexose and ditergalloylhexose (MW 1084) as HHDP-
328 digallagylhexose. However, only dilactones of valoneic acid and sanguisorbic acid (tergalloyl
329 derivatives) were quantified after acid hydrolysis and no gallagic acid dilactone could be recovered.
330 Hydrolyzable tannins **51**, **53**, **56** and **57** are thus clearly tergalloyl derivatives.

331

332 3.1.2. Anthocyanins

333 Two main anthocyanins, delphinidin-3-*O*-glucoside (D3G) and cyanidin-3-*O*-glucoside
334 (C3G) were identified, as described before in the jabuticaba peel (Figure 1B). D3G (**24**) displayed a
335 parent ion in the positive mode at m/z 465 with a major fragment ion at m/z 303 (loss of a hexose
336 moiety) (Plaza et al., 2016; Mena et al., 2012). C3G (**28**) ($[M]^+$ at m/z 449 and 287 in MS²) was
337 identified by comparison with the standard compound. Minor pelargonidin-*O*-hexoside (**32**, $[M]^+$ at

338 m/z 433 and 271 in MS²) and peonidin-*O*-hexoside (**35**, [M]⁺ at m/z 463 and 301 in MS²) were also
339 identified as previously in jabuticaba peel by Quatrin et al. (2019).

340

341 3.1.3 Flavonols

342 Myricetin deoxyhexoside (**43**; m/z 463), quercetin hexosides (**45**, **47**; m/z 463) and quercetin
343 deoxyhexoside (**54**; m/z 447) display daughter ions at m/z 317 or 301, allowing to assign their
344 aglycone part (Figure 1C). For quercetin pentosides (**48**, **49**), their parent and fragment ions at m/z
345 433 and m/z 301 as well as their maximal absorption in the range 350-360 nm were not sufficient to
346 discriminate them from ellagic acid pentoside. However, they yielded in MSⁿ subsequent
347 fragmentation at m/z 273, 257, 179 and 151 which are typical of the benzopyrane moiety of
348 flavonols as evidenced for quercetin (**58**) (Neves et al., 2018)

349

350 3.1.4 Others

351 No caffeoylquinic acids and flavan-3-ols ((epi)catechin, type-B procyanidin dimers,
352 (epi)catechin gallate, gallo(epi)catechin) could be evidenced in jabuticaba peel by search of their
353 parent ions. Thioacidolysis was additionally conducted on both peel extract and peel powder
354 indicating the absence of flavan-3-ol oligomers in these materials.

355

356 3.1.5 Polyphenol quantification by UPLC/DAD

357 Quantification was possible for 32 and 27 compounds out of the 58 tentatively assigned for
358 the DJP-US and DJP-CM extracts, respectively (Table 2).

359 Gallic acid derivatives are the largest class of compounds found for both extracts. They comprise
360 single and mixed esters containing at least a galloyl or HHDP group. The DJP-US extract presented
361 the highest amount in gallic acid derivatives (9.2 ± 0.3 mg/g DJP), differing statistically from the
362 DJP-CM extract (4.9 ± 0.7 mg/g DJP). Contents between 3.5 and 8.2 mg/g DJP were previously

363 reported using different extraction methods and standards (Albuquerque et al., 2020; Quatrin et al.,
364 2019; Plaza et al., 2016).

365 Similar levels in ellagic acid were quantified in both DJP-US and DJP-CM extracts (1.1 mg/g DJP)
366 and these contents are in the same range as those found by Inada et al. (2015) (1.8 mg/g DJP), Plaza
367 et al. (2016) (1.4 mg/g DJP) and Quatrin et al. (2019) (0.5 mg/g DJP) although without ellagic acid
368 as standard for the latter two groups.

369 Anthocyanins, quantified as both C3G and D3G, are the second most abundant group. Contents in
370 anthocyanins were found to be significantly lower for the DJP-US extract (7.8 ± 0.1 mg/g DJP) than
371 for the DJP-CM extract (10.2 ± 1.2 mg/g DJP). Plaza et al. (2016) and Albuquerque et al. (2020)
372 found higher amounts of anthocyanin derivatives (32.2 and 24.5 mg/g DJP, respectively) while
373 Quatrin et al. (2019), Peixoto et al. (2016) and Alejandro, Dube et al. (2013) found relatively
374 similar or lower amounts (11.5, 1.7, and 3.5 mg/g DJP).

375 Additionally, the DJP-CM extract presented a statistically higher level of flavonol derivatives (0.32
376 ± 0.02 mg/g DJP) than the DJP-US extract (0.24 ± 0.01 mg/g DJP). Both Quatrin et al. (2019) and
377 Plaza et al. (2016) found similar amounts (0.6 mg/g DJP) with solvent and pressurized hot water
378 extractions. Finally, our work did not reveal any differences for the overall contents in phenolic
379 compounds whatever the extraction method used. Quantification differences between classes can be
380 explained by the behavior of specific polyphenols towards ultrasound. The acoustic cavitation
381 produced by ultrasound promotes a very efficient cell wall disruption. This may have considerably
382 increased the transfer of galloyl, HHDP and tergalloyl hydrophobic esters to the liquid phase.
383 However, this process promotes the creation of shear forces that generate critical temperature and
384 pressure, locally generating hydroxyl radicals which cause the degradation of sensitive compounds
385 such as anthocyanins and flavonols (Wang, Cheng, Ma, & Jia, 2020).

386

387 3.2 Acid hydrolysis of DJP and DJP-US extract

388 Acid hydrolysis of the dried jabuticaba peel and ultrasonic extract was conducted to release
389 the phenolic depsides present as *O*-hexosides. Because of the low water solubility of the released
390 phenolic compounds, an additional step of pellet washing by MeOH/DMSO (1:1) was performed as
391 proposed by Garcia-Villalba et al. (2015). Identification and quantification of the compounds after
392 acid hydrolysis are presented in Table 3 while the chromatographic profiles are in Figure 1BC and
393 Suppl. Mat.

394 Gallic acid (**1**) was not present in the jabuticaba peel and appeared in large amounts after
395 acid hydrolysis. Digalloylhexose (**2**) presented a similar MS pattern as compound **4** before
396 hydrolysis (Table 1) with a parent ion at m/z 483 and fragment ions at m/z 331, 313 and 169. Its
397 resistance to acid hydrolysis points to the presence of C-glycosidic linkages. The dehydrated form
398 of gallic acid *C*-hexoside (**3**) was identified with a parent ion at m/z 313 and a fragment at m/z 169
399 for gallic acid. A parent ion at m/z 181 and fragments at m/z 137 and 109 as for the standard
400 allowed the identification of dihydroxyphenylpropionic acid (**4**).

401 Newly formed compounds **14** and **15** were tentatively assigned as valoneic acid dilactone
402 and sanguisorbic acid dilactone, respectively. Sanguisorbic acid dilactone presented a parent ion at
403 m/z 469 and fragments at m/z 425, 301 and 299 as observed from sanguisorbic acid dilactone
404 obtained from hydrolysis of strawberry (Mattila and Kumpulainen, 2002). Additionally, valoneic
405 acid dilactone eluted earlier than sanguisorbic acid dilactone and presented fragments at m/z 425,
406 407 and 301 as in pomegranate (Garcia-Villalba et al., 2015). Contents in valoneic acid dilactone
407 were 3 to 4-fold higher than contents in sanguisorbic acid dilactone in both DJP and the DJP-US
408 extract.

409 No gallagic acid dilactone was found to form after 24 h of hydrolysis in contrast to
410 pomegranate peel (García-Villalba et al., 2015). Compounds **7**, **11**, **15**, **33**, **38** and **52** (Table 1) were
411 without ambiguity assessed as HHDP-tergalloylhexoses rather than galagylgalloylhexoses and
412 amounted to 0.55 mg/g of DJP after conventional extraction and 1.03 mg/g after ultrasound

413 extraction. Therefore, hexose acylated with the gallagyl group as in pedunculagin III, punicalagin
414 and punicalin remains typical of pomegranate ellagitannins (García-Villalba et al., 2015). Similarly,
415 **51** and **57** should be HHDP-galloyltergalloylhexoses while **53** and **56** ditergalloylhexoses.
416 Additionally, various *C*-hexosides of valoneic/sanguisorbic acid dilactone or tergallic acid formed
417 in tiny amounts and could not be quantified in this work (**7**, **9**, **11**, **12** and **13**) (Table 2). Castalagin
418 and vescalagin are expected to release tergalloyl-*C*-glucose after acid hydrolysis while alnusiin will
419 give valoneic acid dilactone.

420 Ellagic acid (**16**) was the most intense peak from pellet washing. Four HHDP-ellagic acid
421 derivatives were identified (**5**, **6**, **8**, **10**) with parent ions at m/z 783 and fragmentation indicating the
422 loss of HHDP/ellagic acid ($[M-302-H]^-$ at m/z 481 and fragment at m/z 301). One of the substituent
423 must be ellagic acid rather than HHDP as these compounds absorb between 365 and 375 nm. They
424 could be *C*-hexosides rather than *O*-hexosides as they are resistant to acid hydrolysis and are likely
425 arising from a rearrangement of the five di-HHDP hexosides detected at different retention times in
426 peel (Table 1).

427 Most abundant hydrolysis compounds are in decreasing order: gallic acid *C*-hexoside (**3**),
428 ellagic acid (**16**), gallic acid (**1**) and valoneic acid dilactone (**14**). Gallic acid amounted to 11.4 mg/g
429 in DJP and 6.3 mg/g after ultrasound extraction. The content in ellagic acid increased upon
430 hydrolysis from 1.1 to 16.4 mg/g for DJP and from 1.1 to 10.9 mg/g after ultrasound treatment.
431 These values agree well with that found by Alejandro, Dube et al. (2013) after acid hydrolysis of
432 DJP (22.5 mg/g DJP). Largely lower contents in gallic acid (0.4 and 1.5 mg/g DJP, respectively)
433 and ellagic acid (0.3 and 2.8 mg/g DJP, respectively) were found by Quatrin et al. (2019) and Inada
434 et al. (2015) after polyphenol extraction and subsequent alkaline and acid hydrolyses of the residue
435 supporting the degradation of the oxidizable trihydroxyphenyl moiety in basic conditions.

436 Overall, hydrolytic products of ellagitannins amounted to 70.3 mg/g DJP (no extraction) and
437 42.4 mg/g DJP after ultrasound extraction. This difference was likely due to incomplete extraction
438 by EtOH:water (1:1) of phenolic compounds which are rather hydrophobic or bound to cell wall

439 polysaccharides and proteins. These amounts are markedly higher than those obtained by summing
440 individual compounds from UPLC analysis (10.3 and 6.0 mg/g DJP after ultrasound and solvent
441 extractions, respectively, and subtraction of anthocyanins and flavonols in Table 2). Acid
442 hydrolysis, which relies on the titration of two standards ellagic acid and gallic acid, appears by far
443 more accurate than UPLC in the absence of individual standards. Subsequent washing of the pellet
444 with MeOH/DMSO provided one third of the hydrolysis products. It is worth noting that
445 hydrophobic molecules such as ellagic acid and valoneic acid dilactone were mainly recovered
446 using this additional step whereas sanguisorbic acid dilactone was only soluble in this solvent
447 system.

448

449 **3.3 Inhibition of lipid oxidation by the DJP-US extract and cyanidin-3-O-glucoside in *in*** 450 ***vitro* gastrointestinal digestion**

451 The lipid oxidation initiated by metmyoglobin (MbFe^{III}) of a 10% oil-in-water emulsion
452 stabilized by egg yolk phospholipids was used to simulate the digestion of a Western type diet rich
453 in ω -6 polyunsaturated lipids (Gobert et al., 2014). Initial gastric digestion is characterized by a fast
454 rise in pH after meal intake. Then pH almost linearly decays returning to a basal pH of 2 after
455 gastric emptying. Although static modeling cannot reproduce pH kinetics, two one hour-periods
456 were sequentially run in this work, one at pH 5 and the other at pH 3. Conjugated dienes (CD),
457 consisting mainly of the lipid-derived hydroperoxides formed in the propagation step of the lipid
458 oxidation process, were followed as primary markers. Secondary marker 4-hydroxy-2-nonenal (4-
459 HNE) was selected as a specific end-product of the oxidation of ω -6 lipids.

460

461 *3.3.1 Lipid oxidation in the absence of antioxidant*

462 In the control experiment, CD and 4-HNE accumulated in a similar linear pattern yielding 28
463 μ moles of CD/g of lipids and 86 nmoles of 4-HNE/g of lipids after 2 hours of gastric digestion (Fig.
464 2AB). The low 4-HNE/CD ratio of 3 to 1000 observed at this stage can be explained by the

465 formation of various other secondary oxidation products such as short-chain aldehydes, epoxides
466 and alcohols as well as by the high reactivity of 4-HNE. Electrophilic 4-HNE reacts rapidly with
467 nucleophilic cysteine, histidine and lysine residues in proteins as already evidenced in pig gastric
468 digesta (Delosiere et al., 2016). Increase of pH at 6.5 and addition of bile and pancreatin induced an
469 apparent drop in lipid oxidizability in the early intestinal phase. A CD faster degradation or an
470 incomplete extraction of lipids from the mixed micelles by the 2-propanol/hexane system could
471 explain this apparent CD loss. As to 4-HNE, it disappeared almost totally. Then, CD and 4-HNE
472 exhibited different patterns in the intestinal step. CD kept accumulating reaching 40 μ moles/g lipids
473 at the end of the intestinal step. In contrast to the concomitant formation of 4-HNE and CD in the
474 gastric phase, free 4-HNE only weakly accumulated reaching 7.5 μ moles per g lipids after 240 min.
475 Additional proteins and enzymes from bile and pancreatin may have reacted with continuously
476 formed 4-HNE.

477

478 *3.3.2 Lipid oxidation in the presence of the DJP-US extract and cyanidin-3-O-glucoside*

479 The DJP-US extract and C3G were evaluated as inhibitors of lipid oxidation at similar anthocyanin
480 concentrations although anthocyanins in the ultrasonic extract were constituted by 89% of C3G and
481 11% of D3G. Five-fold higher levels in anthocyanins were brought through US-5 and C3G-5
482 compared to US-1 and C3G-1.

483 In the gastric step, the addition of C3G slowed down CD accumulation in a concentration-
484 dependent manner with a significant effect ($p = 0.05$) for both levels between 90 and 120 min. After
485 2 h of gastric digestion, CD inhibition was of 52% and 65% for C3G-1 and C3G-5, respectively.
486 Interestingly, US-1 and US-5 extracts afforded a largely higher protection with inhibition rates of
487 87% and 100%, respectively. This difference was also outlined for the accumulation of 4-HNE. As
488 a matter of fact, C3G-1 and C3G-5 significantly inhibited 4-HNE formation between 90 and 120
489 min while being significantly less antioxidant than the two concentrations used for the DJP-US
490 extract. Inhibition rates were 28% and 55% for C3G-1 and C3G-5 while 91% and 97% for US-1

491 and US-5. With pure C3G, 30 and 60 min-long lag phases were evidenced and found consistent
492 with remaining C3G during these periods (Fig. 3).

493 After 30 min of intestinal digestion, the DJP-US extracts appeared unexpectedly unable to inhibit
494 the accumulation of CD with apparent rates almost faster than for control. In contrast, the less
495 antioxidant C3G-1 and C3G-5 kept their inhibitory action throughout the intestinal phase. This
496 unexpected increase in CD with the DJP-US extracts may be ascribed to the extraction by the 2-
497 propanol/hexane mixture of amino acids or small peptides released through proteolysis of the
498 jabuticaba peel extract and absorbing at 234 nm. This apparent loss of activity has also been
499 observed for apple puree and extract (unpublished data). Epicatechin used for comparison strongly
500 inhibited CD accumulation as evidenced here with C3G. Primary lipid oxidation products should be
501 assessed with a different method because of the possible interferences at 234 nm from complex
502 plant products.

503 As observed earlier for control, no clear accumulation of 4-HNE occurred in the intestinal step in
504 agreement with the likely fast reaction of this marker with digestive proteins at a higher pH.

505

506 **3.4 Bioaccessibility of anthocyanins**

507 Only anthocyanins could be recovered during the digestion of phospholipid-stabilized emulsions
508 under oxidative stress. Other phenolic compounds from the DJP-US extracts were present at too
509 low levels to be quantified.

510 In the gastric step, pure C3G disappeared very fast from the aqueous phase since only 14% were
511 recovered for the highest concentration at 30 min and 0% for the lowest concentration. By contrast,
512 C3G from US-1 and US-5 was largely present at the end of the gastric step with recovery rates of
513 21% and 72%. Less abundant D3G could only be recovered from US-5 at a rate of 22%. This lower
514 recovery is mainly due to the higher oxidizability of the 1,2,3-trihydroxyphenyl moiety of
515 delphinidin when compared to that of the 1,2-dihydroxyphenyl nucleus of cyanidin. Similarly,

516 Quatrin et al. (2020) showed a higher recovery of C3G (85%) than D3G (71%) at the end of the
517 gastric step for a jaboticaba peel powder.

518 The 5 min-long transition from the gastric to the intestinal phase led to a complete disappearance of
519 C3G from US-1 and D3G from US-5. C3G from US-5 dropped to 26% during this short period
520 before totally disappearing. As sample acidification was conducted before anthocyanin
521 quantification, the decays observed suggest a degradation of the flavylum nucleus as a result of
522 both antioxidant action and unfavorable near neutral conditions.

523

524 **4. Discussion**

525 **Lipid oxidation and inhibition in gastrointestinal digestion**

526 Chemical reactions and physicochemical interactions occurring between bolus constituents
527 should be unraveled to shed some light on the bioavailability of plant secondary metabolites and
528 their health benefit directly in the gastrointestinal tract. Digestion modeling allows to take into
529 account both human food and physiological conditions of digestion. An average Western adult daily
530 consumes 100 to 150 g of triglycerides and 2 to 10 g of phospholipids. After gastric antral
531 contractions, dietary lipids are recovered in the emulsion form with most of the lipid droplets in the
532 1-50 μm diameter range (Lorrain, Dangles, Genot, & Dufour, 2010). A 12.5% oil-in water emulsion
533 stabilized by egg yolk PL and characterized by a single mode at $14.8 \pm 0.3 \mu\text{m}$ and a $D(3,2)$ at 1.09
534 $\pm 0.18 \mu\text{m}$ was thus prepared as food model. While simulated fluids, enzymes (1000 U pepsin/mL,
535 100 U trypsin/mL) and bile levels were as proposed by Minekus et al. (2014), the gastric pH was
536 more physiologically simulated with a pH jump from 5 to 3 after one hour.

537 The heme iron form of red meat, metmyoglobin, was added at a 30 μM level in the gastric
538 step simulating the consumption of 85 g of beef steak (chyme volume of 1 L). Metmyoglobin-
539 initiated oxidation of polyunsaturated fatty acid led to the formation of lipid-derived conjugated
540 dienes and 4-HNE as primary and end markers, respectively (Fig. 2AB). The change in pH from 5
541 to 3 appeared to increase both rates of accumulation. As a matter of fact, the proteolysis of

542 metmyoglobin by pepsin has been evidenced at pH 5 although this process is faster at lower pHs
543 (Bolea et al., 2019). The resulting micro-metmyoglobin can accelerate lipid oxidation likely due to
544 a facilitated access for lipids to the iron center. Then, pH lowering leads to the formation of an
545 unfolded form of metmyoglobin which appeared to be stable during at least 30 min at pH 3. The
546 release of heme iron from proteolyzed unfolded metmyoglobin at pH 3, although suggested by some
547 authors, is still unclear by now. Whatever the heme iron form, lipid oxidation proceeded at both
548 gastric pHs suggesting that this deleterious process is susceptible to degrade essential ω -6 lipids *in*
549 *vivo*.

550 Thus, it is critical to identify protective food that, when co-ingested, could limit lipid oxidation in
551 the GIT and the onset of cardiovascular diseases in a longer term. In this work, the effect of a
552 jaboticaba peel rich in anthocyanins and hydrolyzable tannins was compared to that of a pure
553 anthocyanin, C3G. At identical anthocyanin contents, the DJP-US extracts more extensively
554 inhibited the accumulation of CD and 4-HNE than C3G-1 and C3G-5. This difference was
555 particularly significant at 90 and 120 min of gastric digestion for 4-HNE ($p < 0.05$) and at 60, 90
556 and 120 min for CD (Fig. 2AB). C3G-1 and C3G-5 were found to inhibit the accumulation of the
557 primary marker CD in a concentration-dependent manner while totally inhibiting that of 4-HNE
558 during lag phases that correlate with the presence of C3G in the digesta. After the disappearance of
559 C3G, 4-HNE formed at rates that are lower than observed for control suggesting an additional
560 antioxidant effect of the degradation products of C3G. In the pH range of 2 to 4 or in oxidative
561 conditions at pH 7, cyanidin glycosides were reported to be degraded into protocatechuic acid,
562 phloroglucinaldehyde and 3,5,7-trihydroxycoumarin among others (Dangles & Fenger, 2018).
563 Protocatechuic acid, which is also a metabolite from quercetin oxidation, could contribute to the
564 antioxidant capacity of C3G as previously demonstrated for quercetin (Lorrain et al., 2010).

565 The almost total inhibition of lipid oxidation in the presence of US-5 is consistent with the high rate
566 of remaining anthocyanins, found to be 72% for C3G and 22% for D3G, after the gastric step (Fig.
567 3). The potent capacity of the DJP-US extract can also be attributed to the additional presence of

568 hydrolyzable tannins and flavonols which constitute 84% and 0.5% of the phenolic pool of the
569 jaboticaba peel extract (evaluation after acid hydrolysis). All of these compounds display the typical
570 1,2-dihydroxy- and 1,2,3-trihydroxyphenyl moieties that are critical to the reducing capacity of
571 phenolic compounds.

572 In the intestinal phase at pH 6.5, the formation of lipid-derived conjugated dienes was demonstrated
573 for the first time suggesting that the heme iron form maintains its prooxidant activity towards
574 triacylglycerols and their lipolytic products. Those are mainly monoacylglycerols and free fatty
575 acids solubilized in mixed micelles with biliary salts. Although this work did not allow to evidence
576 the accumulation of 4-HNE because of its rapid scavenging by peptides and proteins, primary lipid
577 oxidation products are known to be unstable in intestinal conditions. Indeed, the *in vitro* digestion
578 of a thermodegraded sunflower oil generated hexanal and nonanal as the major alkenals while 2-
579 heptenal and 2-octenal were recovered in higher rates than 4-HNE (Goicoechea et al., 2008).
580 Moreover, Kanazawa & Ashida (1998) identified uncleaved epoxyketones, 9-oxononanoic acid and
581 hexanal in the intestinal lumen of rats fed with linoleyl hydroperoxides. 4-HNE is absorbable and
582 can be recovered under various metabolized forms (Keller, Baradat, Jouanin, Debrauwer, &
583 Gueraud, 2015). By covalently reacting with the ApoB protein of LDL, 4-HNE is involved in the
584 formation of atherogenic oxLDL particles. Furthermore, 4-HNE appears to play a role in the
585 development of colonic preneoplastic lesions (Bastide et al., 2017).

586

587 **Bioaccessibility and reactivity of anthocyanins and hydrolyzable tannins**

588 Diet-induced oxidative stress led to the fast disappearance of C3G during gastric digestion.
589 Anthocyanins can play an antioxidant role in metmyoglobin-initiated lipid oxidation by reducing
590 the ferrylmyoglobin initiating species rather than the propagating peroxy radicals (Goupy, Bautista-
591 Ortin, Fulcrand, & Dangles, 2009). Interestingly, hydrolyzable tannins, which are present in a 5-
592 fold excess compared to anthocyanins in the DJP-US extract, had the ability to slow down their
593 degradation. Hydrolyzable tannins were constituted for half by HHDP-galloylhexose **2**, ellagic acid

594 **42**, HHDP-digalloylhexoses **29** and **37**, tetragalloylhexose **46**, pentagalloylhexose **50**, HHDP-
595 tergalloylhexose **38**, and di-HHDP-hexoses **10** and **17**, in decreasing order. This composition agrees
596 well with results from Plaza et al. (2016) and Quatrin et al. (2019). The possible interaction of these
597 high-molecular weight compounds with the iron-oxo center of ferrylmyoglobin is suggested by the
598 superior antioxidant activity of the extracts. Furthermore, ellagitannins and gallotannins are prone to
599 degradation in *in vitro* gastrointestinal digestion, yielding mainly gallic acid and ellagic acid for a
600 jabuticaba peel powder (Quatrin et al., 2020). Their stability was mostly affected by intestinal
601 conditions which also led to the total disappearance of anthocyanins while only lightly affecting
602 flavonols. A matrix effect should also be taken into consideration when investigating extracts and
603 peel powders (Quatrin et al., 2020; Peixoto et al., 2016). Binding to cell wall polysaccharides and
604 proteins is expected to slow polyphenol diffusion out of the matrix increasing thus their life time.

605

606 **Conclusion**

607 Jabuticaba peel hydroalcoholic extracts exhibit high potency in limiting the formation of deleterious
608 lipid hydroperoxides and aldehydes arising from diet-induced oxidation of essential ω -6 lipids in
609 the gastrointestinal tract. This inhibitory capacity can be ascribed to high levels in antioxidant
610 anthocyanins (1% DM) and hydrolyzable tannins (7% DM), thus pointing to promising future uses
611 for jabuticaba extracts as food ingredient or colorant.

612

613 **Conflicts of interest**

614 There are no conflicts of interest to declare.

615

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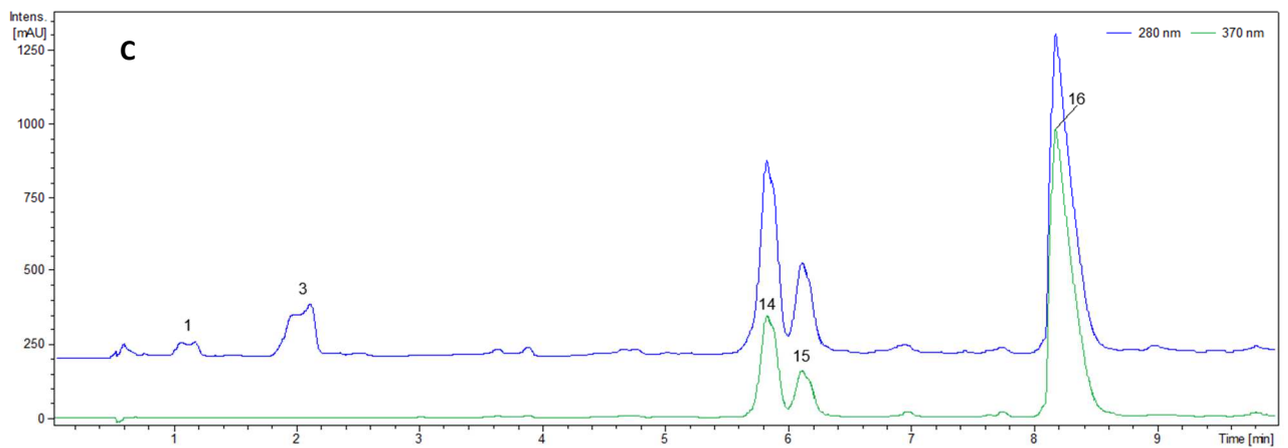
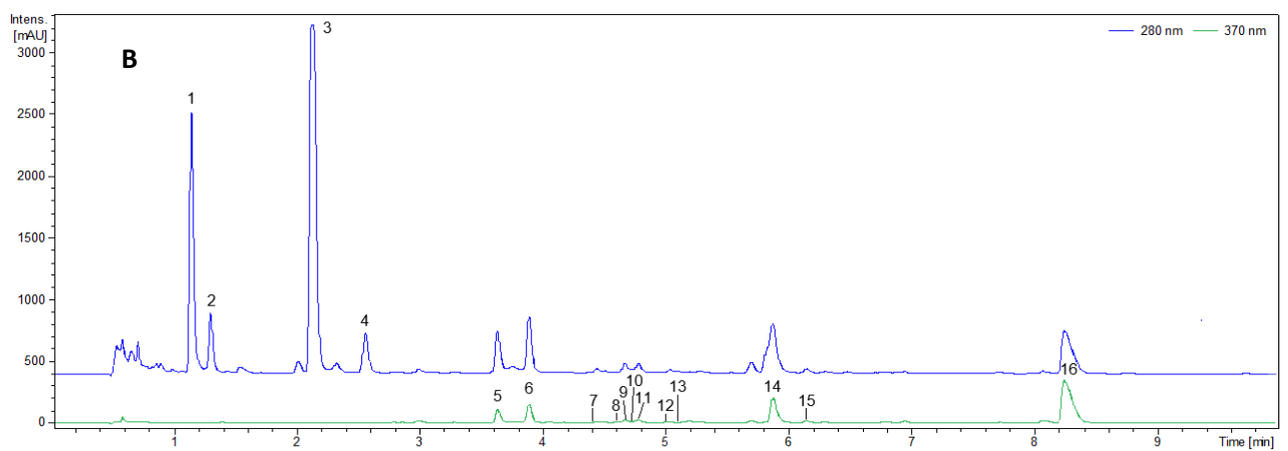
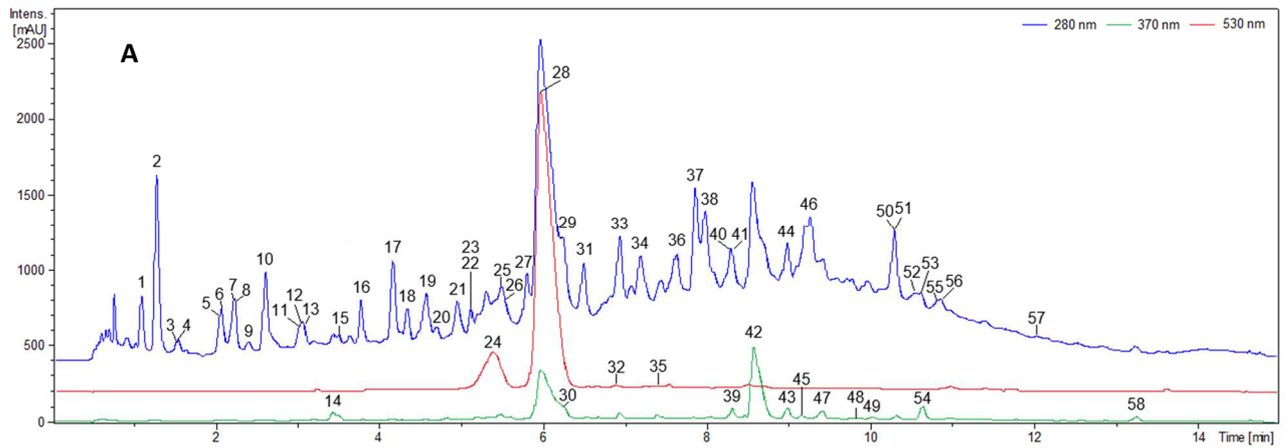


Figure 1. Chromatographic profile of the phenolic compounds for the DJP-US extract before hydrolysis at 280 nm, 370 nm and 530 nm (A) and after hydrolysis at 280 nm and 370 nm (B – Supernatant and C – Pellet washing).

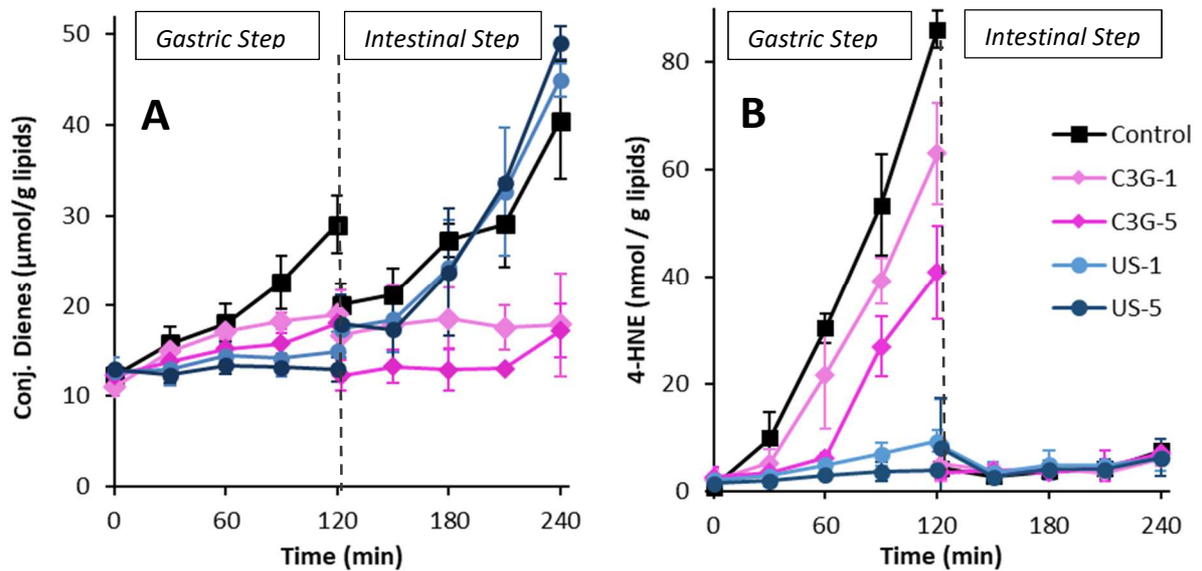


Figure 2: Inhibition by the DJP-US extract and pure C3G of the accumulation of (A) lipid-derived conjugated dienes (CD) and (B) 4-HNE during the *in vitro* simulated digestion of phospholipid-stabilized emulsions. US-1 and C3G-1 bring 14.9 µg total anthocyanins while US-5 and C3G-5 74.4 µg. Lipid oxidation was initiated by metmyoglobin (30 µM). Values represent mean ± SD ($n = 3-7$).

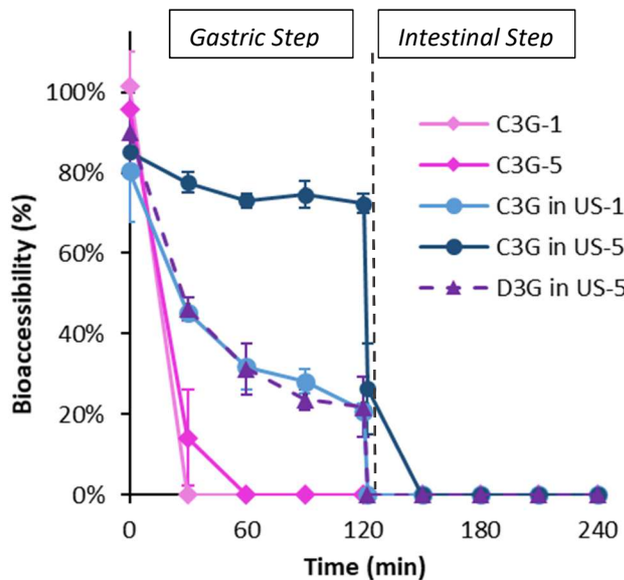


Figure 3: Bioaccessibility of C3G and D3G during the *in vitro* simulated gastrointestinal digestion of phospholipid-stabilized emulsions in the presence of metmyoglobin (30 µM) and the DJP-US extract or pure C3G. Values represent mean ± SD ($n = 3$).

Table 1. Phenolic compounds identified by UPLC/DAD/ESI-MSⁿ in dried jabuticaba peel extracts obtained by ultrasound technology and conventional solvent extraction.

N ^o	RT (min)	λ_{\max} (nm)	$[M - H]^- / [M]^+$ (<i>m/z</i>) [*]	MS ² fragments (<i>m/z</i>)	Tentative identification
1	1.1	277	331	271-169	Galloylhexose ^a
2	1.3	270	633	301-275-249	HHDP-galloylhexose (1) ^{abh}
3	1.6	266	649	605-481-479-425	HHDP-galloylgluconic acid ^c
4	1.6	266	483	331-313-169	Digalloylhexose (1) ^{ad}
5	2.1	264	633	301-275-249	HHDP-galloylhexose (2) ^{abh}
6	2.1	264	783	765-721-507-481-419-341-301-275	Di-HHDP-hexose (1) ^{abdhj}
7	2.2	282	933	915-871-613-569	HHDP-tergalloylhexose (1) ^{defg}
8	2.2	282	801	783-757-633	HHDP-digalloylgluconic acid (punigluconin) (1) ^{cd}
9	2.5	260	783	481-301-275	Di-HHDP-hexose (2) ^{abdhj}
10	2.7	260	783	481-301-275	Di-HHDP-hexose (3) ^{abdhj}
11	3.1	260	933	915-631-613-569	HHDP-tergalloylhexose (2) ^{adefg}
12	3.1	260	801	757-633	HHDP-digalloylgluconic acid (punigluconin) (2) ^{cd}
13	3.1	260	951	907-783-605-481-405-347-301	Di-HHDP-galloylgluconic acid (1)
14	3.5	370	469	425	Tergallic acid dilactone (1)
15	3.6	260	933	631-571-451-425-301	HHDP-tergalloylhexose (3) ^{adefj}
16	3.8	260	633	615-481-301-275	HHDP-galloylhexose (3) ^{abh}
17	4.2	260	783	481-421-301-275	Di-HHDP-hexose (4) ^{abdhj}
18	4.4	260	951	907-783-605-481-425-361-299	Di-HHDP-galloylgluconic acid (2)
19	4.6	260	951	907-783-605-481-425-361-299	Di-HHDP-galloylgluconic acid (3)

20	4.7	260	631	613-571- 451 -407-351	Valoneic (sanguisorbic) acid dilactone C-hexoside or Tergalloyl-O-hexose (Cauliflorin) ^e or Tergalloyl-C-hexose ^a (1)
21	5.0	270	785	633- 483 -419-331-301-275	HHDP-digalloylhexose (1) ^{abdfj}
22	5.1	272	483	331- 271 -169	Digalloylhexose (2) ^{ad}
23	5.1	272	631	613-571- 451 -407-351	Valoneic (sanguisorbic) acid dilactone C-hexoside or Tergalloyl-O-hexose (Cauliflorin) ^e or Tergalloyl-C-hexose ^a (2)
24	5.4	526	465*	303*	Delphinidin-3-O-glucoside ^{abh}
25	5.6	260	633	615-463- 301	HHDP-galloylhexose (4) ^{abhj}
26	5.7	260	935	917 -873-855-573	Di-HHDP-galloylhexose (1) (Casuarinin/Stachyurin/Casuarictin/Potentillin) ^{abdefhj}
27	5.9	265	935	917-873-783-659- 633 -571-301	Di-HHDP-galloylhexose (2) (Casuarinin/Stachyurin/Casuarictin/Potentillin) ^{abcdefh}
28	6.2	514	449*	287*	Cyanidin-3-O-glucoside (std) ^{abh}
29	6.3	270	785	633- 483 -419-301-249	HHDP-digalloylhexose (2) ^{abdfj}
30	6.3	372	469	451- 425 -407-301-167	Valoneic acid dilactone ^{cfh}
31	6.5	260	951	907 -783-605	Di-HHDP-galloylgluconic acid (4)
32	6.9	516	433*	271*	Pelargonidin-O-hexoside ^a
33	7.0	260	933	763-631-481- 451 -425-301	HHDP-tergalloylhexose (4) ^{adefj}
34	7.2	264	635	617-483- 423 -271	Trigalloylhexose ^a
35	7.4	524	463*	301*	Peonidin-O-hexoside ^a
36	7.7	260	935	917-765- 633 -615-481-451-301	Di-HHDP-galloylhexose (3) (Casuarinin/Stachyurin/Casuarictin/potentillin) ^{abdf}
37	7.9	264	785	633- 483 -301	HHDP-digalloylhexose (3) ^{abdfj}
38	8.0	260	933	631-481- 451 -301	HHDP-tergalloylhexose (5) ^{adefj}
39	8.3	355	433	301 -287-273-209	Ellagic acid pentoside ^{abdfh}
40	8.3	-	783	765 -613-465-451-419-301	Di-HHDP-hexose (5) ^{abdh}
41	8.3	-	937	785- 767 -741-635-465-419-301-275	HHDP-trigalloylhexose ^{bfhj}

42	8.6	367	301	<i>229-185</i>	Ellagic acid (std) ^{abcdeh}
43	9.0	350	463	317-237-179-151	Myricetin deoxyhexoside ^{ahi}
44	9.0	264	787	635- 617 -467-447-403	Tetragalloylhexose (1) ^{ad}
45	9.2	350	463	301-273-245-213-179-151	Quercetin hexoside (1) ^{ai}
46	9.3	272	787	617-465-449	Tetragalloylhexose (2) ^a
47	9.4	350	463	301-273-257-229-179-151	Quercetin hexoside (2) ^{ai}
48	9.8	350	433	301-271-257-179	Quercetin pentoside (1) ^{ai}
49	10.0	350	433	301-271-253-225-179-151-125	Quercetin pentoside (2) ^{ai}
50	10.3	275	939	787- 769 -617-599-447	Pentagalloylhexose ^{abj}
51	10.3	275	1085	783-633-451-301	HHDP-galloyltergalloylhexose (1) ^{ad}
52	10.5	260	933	631-481- 451-301	HHDP-tergalloylhexose (6) ^{adefj}
53	10.5	260	1083(Nf), 541[M-2H] ²⁻	[541]: 631-466-451-301	Ditergalloylhexose (trace) (1) ^d
54	10.7	350	447	301-255--217-207-179-151	Quercetin deoxyhexoside (quercetin-3-rhamnoside ^b) ^{ahij}
55	10.8	260	1087	935- 917 -749-451	Di-HHDP-digalloylhexose
56	10.9	260	1083(Nf), 541[M-2H] ²⁻	[541]: 631-452-301	Ditergalloylhexose (trace) (2) ^d
57	12.2	260	1085	783-633-481-451-301	HHDP-galloyltergalloylhexose (2) ^{ad}
58	13.3	366	301	<i>179-151</i>	Quercetin ⁱ

*Ionization in negative mode for all compounds except for anthocyanins in positive mode. The main fragment in MS² is given in boldface, the minor fragments are given in normal font and the ultra-minor fragments are given in italics. HHDP: hexahydroxydiphenoyl group. Std: identified with authentic standard. Nf: not fragmented. [M-2H]²⁻: doubly charged ion.

^aQuatrin et al., 2019; ^bPlaza et al., 2016; ^cFischer et al., 2011; ^dGarcia-Villalba et al., 2015; ^ePereira et al., 2017; ^fFracassetti et al., 2013; ^gTavares et al., 2016; ^hWu et al., 2012; ⁱNeves et al., 2018; ^jAlbuquerque et al., 2020; ^kMorales et al., 2016.

Table 2. Contents in various phenolic classes for dried jabuticaba peel after extraction by ultrasound technology and conventional solvent extraction.

	Gallic acid derivatives (mg/g DJP)	Ellagic acid (mg/g DJP)	Flavonol derivatives (mg/g DJP)	Anthocyanin derivatives (mg/g DJP)	Sum of phenolic compounds (mg/g DJP)
Ultrasound extraction	9.23 ± 0.32 (a)	1.08 ± 0.02 (a)	0.24 ± 0.01 (b)	7.81 ± 0.06 (b)	18.36 ± 0.36 (a)
Solvent extraction	4.89 ± 0.65 (b)	1.11 ± 0.07 (a)	0.32 ± 0.02 (a)	10.22 ± 1.17 (a)	16.53 ± 1.19 (a)

Values represent mean ± SD ($n = 3$). Sum of phenolic compounds is obtained from the different columns on the left (UPLC). DJP: dried jabuticaba peel; Different letters indicate a significant difference between both extracts at $p < 0.05$.

Table 3. Acid hydrolysis of dried jabuticaba peel and DJP extract obtained after ultrasound extraction followed by UPLC/DAD/ESI-MSⁿ analysis of phenolic compounds.

N°	RT (min)	λ_{max} (nm)	[M - H] ⁻ (<i>m/z</i>)	MS ² fragments (<i>m/z</i>)	Tentative identification	DJP (mg/g DJP)			US extract (mg/g DJP)		
						Supernatant	Pellet	Sum	Supernatant	Pellet	Sum
1	1.2	271	169	125	Gallic acid (Std) ^{ab}	10.7 ± 0.3	0.75 ± 0.02	11.4 ± 0.3	6.27 ± 0.73	0.02 ± 0.01	6.29 ± 0.74
2	1.35	281	483	331-313-169	Digalloylhexose ^{ab}	0.95 ± 0.23	-	0.95 ± 0.23	0.89 ± 0.03	-	0.89 ± 0.03
3	2.2	284	313	295-245-169-125-107	Gallic acid C-hexoside (dehydrated form) ^c	26.9 ± 4.6	3.49 ± 0.09	30.4 ± 3.8	18.6 ± 1.5	0.77 ± 0.34	19.4 ± 1.8
4	2.55	276	181	137-109	DiOH-phenylpropionic acid ^d	5.07 ± 0.72	-	5.07 ± 0.72	0.67 ± 0.18	-	0.67 ± 0.18
5	3.6	374	783	481-451-301-299-271	HHDP-ellagic acid-C-hexoside (1)	0.12 ± 0.04	-	0.12 ± 0.04	0.13 ± 0.04	-	0.13 ± 0.04
6	3.85	375	783	481-451-301-299-271	HHDP-ellagic acid-C-hexoside (2)	0.45 ± 0.07	-	0.45 ± 0.07	0.32 ± 0.05	-	0.32 ± 0.05

7	4.4	365	631	451 -407-299-271	Valoneic/sanguisorbic acid dilactone <i>C</i> -hexoside or Tergalloyl- <i>C</i> -hexose (1) ^b	-	-	-	-	-	-
8	4.6	365	783	481 -451-301-299-271	HHDP-ellagic acid- <i>C</i> -hexoside (3)	-	-	-	-	-	-
9	4.65	365	631	587-451-301- 299 -271	Valoneic/sanguisorbic acid dilactone <i>C</i> -hexoside or Tergalloyl- <i>C</i> -hexose (2) ^b	-	-	-	-	-	-
10	4.7	365	783	767- 483 -453-303-271	HHDP-ellagic acid- <i>C</i> -hexoside (4)	-	-	-	-	-	-
11	4.75	365	631	451-301- 299 -271	Valoneic/sanguisorbic acid dilactone <i>C</i> -hexoside or Tergalloyl- <i>C</i> -hexose (3) ^b	-	-	-	-	-	-
12	5	365	631	587-451- 299 -271	Valoneic/sanguisorbic acid dilactone <i>C</i> -hexoside or Tergalloyl- <i>C</i> -hexose (4) ^b	-	-	-	-	-	-
13	5.1	365	631	587-451-301- 299 -271	Valoneic/sanguisorbic acid dilactone <i>C</i> -hexoside or Tergalloyl- <i>C</i> -hexose (5) ^b	-	-	-	-	-	-
14	5.9	370	469	451- 425 -407-301-167	Valoneic acid dilactone ^{be}	0.73 ± 0.06	3.44 ± 0.09	4.17 ± 0.09	0.99 ± 0.07	2.02 ± 0.10	3.02 ± 0.17
15	6.15	365	469	451- 425 -301-299	Sanguisorbic acid dilactone ^{be}	-	1.39 ± 0.04	1.39 ± 0.04	-	0.85 ± 0.04	0.85 ± 0.04
16	8.4	367	301	<i>258-229-186</i>	Ellagic acid (Std) ^{abf}	3.52 ± 0.14	12.9 ± 0.5	16.4 ± 0.4	3.62 ± 0.38	7.24 ± 0.61	10.9 ± 1.0
TOTAL PHENOLIC COMPOUNDS						48.4 ± 4.1	21.9 ± 0.9	70.3 ± 3.2	31.5 ± 1.3	10.9 ± 0.5	42.4 ± 1.6

The main fragment in MS² is given in boldface, the minor fragments are given in normal font and the ultra-minor fragments are given in italics. Values represent mean ± SD (*n* = 3). Sum of phenolic compounds is the addition of Supernatant and Pellet. Std: identified with authentic standard. —: Means below quantification limit or not present.

^aQuatrin et al., 2019; ^bGarcia-Villalba et al., 2015; ^cFischer et al., 2011; ^dFragmentation as for standard; ^eMorales et al., 2016; ^fNeves et al., 2018.