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1 **Thermal and non-thermal processing of red-fleshed apple: how are (poly)phenol**
2 **composition and bioavailability affected?**

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27 **ABSTRACT**

28 The present study evaluated the impact of different thermal (infrared-drying, hot air-
29 drying and purée pasteurization) and non-thermal (freeze-drying) processing
30 technologies on red-fleshed apple (poly)phenolic compounds. We further investigated
31 the processing effect on the (poly)phenol bioavailability in a crossover postprandial
32 study where three subjects consumed three apple products (freeze-dried snack, hot air-
33 dried snack and pasteurized purée). (Poly)phenolic compounds present in the apple
34 products and their biological metabolites in urine were analyzed using liquid
35 chromatography coupled to mass spectrometry (UPLC-MS/MS). When comparing
36 different processes, infrared-drying caused important losses in most of the apple
37 (poly)phenolics, while hot air-drying and purée pasteurization maintained
38 approximately 83% and 65% of total (poly)phenols compared with the freeze-dried
39 snack, respectively. Anthocyanins in particular were degraded to a higher extend, and
40 hot air-dried apple and pasteurized purée maintained respectively 26% and 9%
41 compared with freeze-dried apple snack. The acute intake showed that pasteurized purée
42 exhibited the highest (poly)phenol bioavailability, followed by hot air-drying and
43 freeze-dried snack, highlighting the impact of processing on (poly)phenols absorption.
44 In conclusion, for obtaining affordable new red-fleshed apple products with enhanced
45 (poly)phenol bioavailability, purée pasteurization and hot air-drying represent viable
46 techniques. However, to obtain a red-fleshed apple snack with high anthocyanin
47 content, freeze-drying is the technique that best preserves them.

48

49

50 **KEYWORDS:** anthocyanins, bioavailability, (poly)phenolic compounds, red-fleshed
51 apple, thermal-processing, UPLC-MS/MS.

52 1. INTRODUCTION

53 Numerous *in vivo* and *in vitro* studies have demonstrated that (poly)phenol
54 compounds exhibit a wide range of biological effects such as lowering cholesterol,¹
55 antiproliferative activity on cancer cells,² and reducing the risk of suffering heart
56 diseases, asthma and type-2 diabetes,³⁻⁵ among others. However, the dietary habits of
57 the majority of people do not guarantee an adequate intake of fruits and vegetables, and
58 the intake of (poly)phenols is below the amounts found to have significant health
59 effects.⁶

60 In recent years, due to the global interest in phytochemicals and in developing
61 functional foods that are fortified, enriched, or enhanced with improved health-
62 promoting effects, there has been growing interest in the development of commercial
63 red-flesh apple cultivars. These apple cultivars have been obtained by traditional
64 breeding methods from new hybrids with red pulp.⁷ The flesh of these singular apple
65 varieties has an enhanced content of anthocyanins (red colour) with high antioxidant
66 activity and potential health-promoting effects.⁸ Besides, through crossbreeding
67 programs with good-flavoured white-fleshed apples, the poor taste of the wild red-flesh
68 varieties has been improved.⁹ So, these new apple varieties could make healthy eating
69 easier and more available, thus satisfying the increasingly widespread needs for high
70 food quality and diversity.

71 Since the production of apple fruit is seasonal, processing methods have been
72 developed and applied to obtain apple derived products with shelf-stability and increase
73 useful life while minimizing changes in the quality attributes.¹⁰ Dehydration is one of
74 the main and oldest techniques for preserving agricultural and food products, and in
75 recent years for producing ready-to-eat and healthy *snacks* from fresh fruit such as

76 apples, while retaining their nutrients and bioactive compounds and thus, being a
77 healthier alternative to salty or sugary snacks.^{11,12}

78 In the preparation of functional and ready-to-eat foods, freeze-drying method is
79 commonly used as there is a minimum loss of flavour, aroma and bioactive compounds,
80 with near-perfect preservation results. However, due to its expensive cost, other
81 dehydration methods such as hot air-drying or infrared-drying are employed to produce
82 high-quality dried fruits.^{11,13,14} Pasteurization of fruit purée is another commercial and
83 common thermal treatment used to increase shelf-life of fresh fruits. In addition fruit
84 purée can be used as an intermediate product for the production of other products such
85 as nectars, juices with solid particles or smoothies.¹⁵⁻¹⁷

86 In most dehydration methods, such as hot air-drying or infrared-drying, vegetables
87 are subjected to high temperatures at which highly thermosensitive and unstable
88 (poly)phenols can be readily degraded.¹⁸⁻²² However, it has also been shown that food
89 processing can induce chemical or physical modifications such as degradation or
90 modification of cell wall polysaccharides or proteins, molecular interactions between
91 components, and other food matrix factors that enhance (poly)phenol bioaccessibility
92 and bioavailability during digestion.^{23,24}

93 Therefore, the first aim of the present research was to evaluate the impact of four
94 different processing conditions (infrared-drying, hot air-drying, freeze-drying and
95 pasteurization of purée) on apple (poly)phenols stability. Secondly, we also investigated
96 the effect of the processing conditions on the apple (poly)phenol bioavailability in a
97 human pilot study in order to search the optimal conditions to obtain an apple snack
98 product with a higher (poly)phenol bioavailability and, thus, enhancing its functional
99 value.

100

101

2. MATERIALS AND METHODS

102 **2.1. Chemicals and Reagents**

103 Pelargonidin-3-*O*-glucoside, cyanidin- 3-*O*-glucoside, cyanidin-3-*O*-galactoside,
104 delphinidin-3-*O*-glucoside, malvidin-3-*O*-glucoside, hydroxytyrosol, luteolin,
105 kaempferol, eriodictyol, quercetin, luteolin-7-*O*-glucoside, kaempferol-3-*O*-glucoside,
106 quercetin-3-*O*-rhamnoside, quercetin-3-*O*-glucoside, isorhamnetin-3-*O*-glucoside,
107 procyanidin dimer B₂, quercetin-3-*O*-rutinoside (rutin), myricetin, and phloretin-2-*O*-
108 glucoside were purchased from Extrasynthese (Genay, France). *p*-Hydroxybenzoic acid,
109 3,4-dihydroxybenzoic acid (protocatechuic acid), *p*-coumaric acid, gallic acid, caffeic
110 acid, ferulic acid, chlorogenic acid (5-caffeoylquinic acid), naringenin, catechin, and
111 epicatechin were acquired from Sigma-Aldrich (St. Louis, MO, USA). The vanillic acid
112 was from Fluka (Buchs, Switzerland). Methanol (HPLC grade), acetonitrile (HPLC
113 grade), acetic acid, and formic acid were purchased from Scharlab Chemie (Sentmenat,
114 Catalonia, Spain). The water was of Milli-Q quality (Millipore Corp., Bedford, MA,
115 USA).

116 Stock solutions of standard compounds were prepared by dissolving each
117 compound in methanol at a concentration of 1 g/L and storing it in dark flasks at -80 °C.

118

119 **2.2. Plant Material**

120 The red-fleshed apple variety used was ‘Redlove Era’. The apples were provided
121 by NUFRI SAT (Mollerussa, Lleida, Spain) and planted in the “La Rasa” experimental
122 plot (La Rasa, Soria, Spain).

123

124 **2.3. Non-thermal apple processing: freeze-drying**

125 Before drying, the apples were washed, wiped with paper towels and cut into 1
126 cm-sized cubes. The apple cubes were frozen in liquid nitrogen and the freeze-drying

127 was then performed at 0.6 bar with a temperature ramp of -20 to 0°C over 25 hours,
128 followed by a second complete vacuum drying with a temperature ramp of 0 to 20°C
129 over 40 hours (Lyophilizer TELSTAR Lyobeta 15, Terrassa, Spain).

130

131 **2.4. Thermal apple processing**

132 *2.4.1. Hot air-drying:* apples were washed, wiped with paper towels and cut into 1
133 cm-sized cubes. The apple cubes were immediately dried in a pilot dryer composed of a
134 cylindrical stainless-steel basket where the sample was introduced, an electric resistance
135 of 8 kW to heat the air to the desired temperature and a Casals model MA 26 M 2H fan
136 ($K_v = 0.865 C_v$) (Casals, Girona, Spain) connected to the speed variation system. The
137 drier was equipped with a dashboard with an Eliwell thermostat (Protherm controls,
138 Atherstone, England) to control the air temperature and speed of the fan, and a Vaisala
139 hygrometer equipped with a HMP 125B probe (Vaisala, Vantaa, Finland) that
140 controlled the relative humidity of the air at the entrance and exit of the drying
141 equipment. Finally, the control for the velocity of the air entering the basket containing
142 the apple cubes consisted of a WM-DA 4000 turbine anemometer (Pacer instruments,
143 Keene, United States) and several valves. The apple cubes were placed in the basket and
144 hot air without recirculating was passed through it. Drying was carried out until a
145 constant weight (weighing the basket with the apples on an electronic balance mod.EM-
146 60 KAM, A&D Company, Tokyo, Japan) at 60°C for 80 minutes with an air velocity of
147 1.5 ms^{-1} and then the temperature was increased until 70°C and maintained for 40
148 minutes with the same air velocity.

149

150 *2.4.2. Infrared-drying:* the infrared-drying process was carried out in an infrared
151 moisture analyzer mod. IRCDi5 FIR dehydrator (Irconfort, Seville, Spain). A thin layer

152 of the apple cubes was placed inside an aluminium support equipped with a precision
153 balance to record the weight. The continuous weighing of the sample allowed the drying
154 kinetics to be determined. The infrared-drying experiments were automatically stopped
155 when the weight of the cubes remained constant. Several tests were performed for a
156 range of infrared-drying temperatures (35, 40, 50 and 60°C) and the sample mass versus
157 time was recorded. In all the cases, the product weight, initial moisture content and dry
158 matter content of the apple cubes were used to calculate the moisture content obtained at
159 any drying time.

160

161 After the completion of the different drying processes, the dried apple cubes were
162 immediately transferred to airtight plastic containers and kept at -80°C until the
163 (poly)phenol chromatographic analysis. Prior to analysis, a fine powder of the dried
164 apple samples was obtained with the aid of an analytical mill (A11, IKA, Germany).

165

166 *2.4.3. Pasteurized apple purée:* red-fleshed apples were supplied to a local
167 company (Anela Fruits, Girona, Spain) to be processed to pasteurized purée. Briefly,
168 apples were milled to a fine purée which was hermetically closed into sterile containers
169 and submitted to continuous pasteurization in a tubular system (94°C for 10 min).

170

171 **2.5. Apple phenol bioavailability human study**

172 *2.5.1. Subjects and study design*

173 The protocol of the study was approved by the Ethical Committee of the Human
174 Clinical Research Unit at the Arnau Vilanova University Hospital, Lleida, Spain
175 (Approval Number: 13/2016). In this study the kinetics of phenolic metabolites in 24 h
176 urine in different interval times (0-2, 2-4, 4-8, and 8-24 h) were monitored in healthy

177 volunteers after the acute ingestion of freeze-dried apple, hot air-dried apple and
178 pasteurized apple purée. The study was performed in a crossover design to reduce
179 interpersonal variability. The volunteers comprised three healthy women, aged from 24
180 to 38 years, with body mass index between 19.4 and 25.1 kg/m². They declared no
181 gastrointestinal alterations and reported no antibiotic use over the last months before the
182 study. To standardize the baseline point, subjects were asked to follow a low-
183 (poly)phenol diet during the 2 days preceding each dietary intervention and during the
184 dietary intervention day. For this, subjects were asked to limit fruit and vegetables
185 consumption, and to avoid cherries, strawberries, blueberries, tea, coffee, wine, beer,
186 chocolate and all their derived products.

187 Each volunteer received in random order the three apple products in a crossover
188 design with a washout period of 14 days between interventions. The amount of each
189 administered product was: 60 g of freeze-dried apple snack, 66 g of hot-air apple snack
190 and 500 g of apple pasteurized purée, which represented a similar phenolic dose
191 (134±18 mg total phenols) (**Supplemental Table 1**).

192 Urine samples were collected 24 h before (basal conditions) and at the interval
193 times of 0-2, 2-4, 4-8, and 8-24 h after the apple products intake. The volume of urine in
194 each interval was measured and aliquots were stored at -80°C prior to the (poly)phenol
195 chromatographic analysis.

196

197 **2.6. Analysis of phenolic compounds in apple products and phenolic** 198 **metabolites in urine**

199 *2.6.1. Sample Pre-treatment*

200 The pre-treatment for the analyses of the phenolic compounds in apple products
201 was carried out according to Bars-Cortina et al. ⁽⁹⁾ with some modifications. Briefly, 0.4

202 g of dried apple powder or 1.6 g of lyophilized purée (the lyophilization parameters
203 were the same as those detailed in section 2.3.) was weighed and extracted with 10 mL
204 of methanol/Milli-Q water/formic acid (79.9:20:0.1, v/v/v). The samples were vortexed
205 for 10 min and then centrifuged at 8784g for 10 min. The extraction was repeated three
206 times and the supernatants were collected, combined and filtered through a 0.22 µm
207 PVDF filter (Scharlab, Barcelona, Spain) prior to the chromatographic analysis.
208 Samples were analyzed in triplicate.

209 The urine samples were pre-treated by µSPE. The micro-cartridges and their
210 conditioning and equilibration steps were the same as reported in our previous study.²⁵
211 In this case, 100 µL of phosphoric acid at 4% were added to 100 µL of the urine sample,
212 and this solution was loaded into the micro-cartridge. The retained phenolic compounds
213 were then eluted with 2 x 50 µL of methanol. Each sample was prepared in triplicate.

214

215 *2.6.2. Ultra-performance Liquid Chromatography Coupled to Tandem Mass* 216 *Spectrometry (UPLC-MS/MS)*

217 Liquid chromatography analyses were carried out on an AcQuity Ultra-
218 Performance liquid chromatograph coupled to a tandem mass spectrometer from Waters
219 (Milford, MA, USA).

220 Two chromatographic methods using UPLC-MS/MS were used for the analysis
221 of 1) anthocyanins and their metabolites, and 2) the rest of the phenolic compounds and
222 their metabolites. In both methods, the flow rate was 0.3 mL/min, and the injection
223 volume 2.5 µL. The UPLC-MS/MS conditions were the same used in our previous
224 study.²⁵ Briefly, for the analysis of (poly)phenolic compounds (including anthocyanins),
225 the analytical column used was an AcQuity BEH C18 (100 mm × 2.1 mm i.d., 1.7 µm)
226 equipped with a VanGuard PreColumn AcQuity BEH C18 (5 × 2.1 mm, 1.7 µm), also

227 from Waters. For the analysis of anthocyanins and their metabolites, the mobile phase
228 was 10% acetic acid (eluent A) and acetonitrile (eluent B). The elution gradient was
229 0–10 min, 3–25% B; 10–10.10 min, 25–80% B; 10.10–11 min, 80% B isocratic;
230 11–11.10 min, 80–3% B; 11.10–12.50 min, 3% B isocratic. For the analysis of the rest
231 of the (poly)phenolic compounds and their metabolites, the mobile phase was 0.2%
232 acetic acid (eluent A) and acetonitrile (eluent B). The elution gradient for the analysis of
233 these (poly)phenolic compounds was 0–5 min, 5–10% B; 5–12 min, 10–12.4% B;
234 12–18 min, 12.4–28% B; 18–23 min, 28–100% B; 23–25.5 min, 100% B isocratic;
235 25.5–27 min, 100–5% B; and 27–30 min, 5% B isocratic.

236 Tandem MS analyses were carried out on a triple quadrupole detector (TQD)
237 mass spectrometer (Waters, Milford, MA, USA) equipped with a Z-spray electrospray
238 interface. Ionization was achieved using the electrospray (ESI) interface operating in the
239 positive mode $[M - H]^+$ for the analysis of anthocyanins and in the negative mode $[M -$
240 $H]^-$ for the other compounds. The data were acquired through selected reaction
241 monitoring (SRM). The ionization source parameters were as follows: capillary voltage,
242 3 kV; source temperature, 150 °C; desolvation gas temperature, 400 °C, with a flow rate
243 of 800 L/h. Nitrogen (99.99% purity, N₂LCMS nitrogen generator, Claind, Lenno, Italy)
244 and argon ($\geq 99.99\%$ purity, Aphagaz, Madrid, Spain) were used as the cone and
245 collision gases, respectively. The dwell time established for each transition was 30 ms.
246 Data acquisition was carried out with MassLynx 4.1 software.

247 Due to the lack of commercial (poly)phenolic standards and their generated
248 metabolites, some of the compounds were tentatively quantified by using the calibration
249 curve of their precursor or of a (poly)phenolic compound with a similar structure.
250 **Supplemental Table 2** shows the selected reaction monitoring (SRM) conditions as
251 well as its cone voltage and collision energy used for the quantification of these

252 (poly)phenolic compounds. This table also shows in which (poly)phenolic standard
253 compound, these (poly)phenolics have been quantified.

254

255 **2.7. Statistical Analysis**

256 Concentration values of the (poly)phenolic compounds and their metabolites
257 studied were reported as means \pm standard deviation (SD). One-way analysis of
258 variance (ANOVA) and Tukey's test at a level of 0.05 were used to determine the
259 significance of differences among the different apple processing techniques, among the
260 urine excretion of the different (poly)phenolic families after the intake of the three red-
261 fleshed apple products studied, and among intra- and inter-individual differences in this
262 (poly)phenol excretion. Moreover, General Linear Model and One-way ANOVA at a
263 level of 0.05 was used to determine the significance of the % excretion of total
264 (poly)phenols after the intake of the three products derived from red-fleshed apple. All
265 data were analyzed with the Minitab Statistical Software, version 17.2.1 (Minitab Inc.,
266 State College, Pennsylvania, United States).

267

268 **3. RESULTS AND DISCUSSION**

269 **3.1. Effect of apple processing on the phenol stability**

270 The aim of the present study was to explore the potential of new red-fleshed
271 apple varieties, as anthocyanin enriched alternatives (biofortified) to common white-
272 fleshed apples for the development of an apple snack product with an enhanced
273 functional value. So, the first objective of this research was to evaluate the impact of
274 different processing technologies on the (poly)phenols stability of red-fleshed apple
275 and, therefore, to search the optimal conditions to develop an apple snack product with a

276 high shelf-stability maintaining at the same time a high concentration of bioactive
277 (poly)phenols. The different products are shown in **Figure 1**.

278 A total of 26 (poly)phenols were identified in the red-fleshed apple products
279 using UPLC-MS/MS (**Table 1**) and included compounds from the following 6
280 (poly)phenol classes (**Figure 2**): i) Phenolic acids, ii) Flavan-3-ols, iii) Flavonols, iv)
281 Anthocyanins, v) Flavanones, and vi) Dihydrochalcones.

282 The impact of the different processes on the stability of the (poly)phenols in red-
283 fleshed apple products was compound-dependent and some (poly)phenols were
284 degraded more than others (**Table 1**). In general terms as seen in **Figure 2**, when
285 comparing the different processes, infrared-drying caused important losses in most of
286 the apple (poly)phenolics, while hot air-drying and purée pasteurization allowed the
287 maintenance of approximately 83% and 65% of the (poly)phenols compared with the
288 freeze-dried snack, respectively. In accordance with the literature^{9,26-28} 5-*O*-
289 caffeoylquinic acid (chlorogenic acid) (**Table 1**) was the most abundant phenolic acid
290 and the main phenolic compound detected in the apple samples (representing 90%-95%
291 of the total phenolic acids and 60-70% of the total phenolic content). The group of
292 phenolic acids was significantly influenced by the dehydration technique used, freeze-
293 drying and hot air-drying being the techniques that preserves them best (**Figure 2**). The
294 total content of phenolic acids was 1661 mg/kg d.w. in freeze-dried apple samples,
295 which was reduced to 1587 mg/kg d.w. in hot air-dried and followed by the values
296 found in the pasteurized purée (1187 mg/kg d.w.), while infrared-dried samples
297 presented the lowest amounts ranging from 474 to 690 mg/kg d.w. (**Table 1**).

298 In this study, four flavan-3-ols were detected and quantified in the apple
299 samples. The freeze-dried samples contained the highest amounts (197 mg/kg d.w.),
300 followed by the hot-air dried (101 mg/kg d.w.) and the pasteurized purée (80.9 mg/kg

301 d.w.) (**Table 1**). In accordance with Bars-Cortina et al. ⁽⁹⁾, the most abundant flavan-3-ol
302 was a epicatechin dimer followed by epicatechin and a trimer. In the infrared-dried
303 apple samples, all flavan-3-ols were found in significantly lower concentrations and the
304 trimer was not detected in any of the infrared temperatures studied. Compared to
305 phenolic acids, flavan-3-ols showed higher losses due to thermal treatment with losses
306 of approximately 49% and 59% in hot air-dried apples and pasteurized purée,
307 respectively and between 87% and 93% in infrared-dried apples with respect to freeze-
308 dried apples.

309 Regarding flavonols, the most abundant were those derived from quercetin, in
310 accordance with the literature,²⁶ being quercetin-rhamnoside the most abundant in all
311 the apple products. Apart from these, other flavanols, such as dihydroquercetin and
312 quercetin arabinoside were detected in all samples (**Table 1**). Quercetin was only
313 detected in the hot air-dried and pasteurized purée samples. Furthermore, quercetin
314 glucoside and dihydrokaempferol glucoside were not detected in the infrared-dried
315 samples. As in the previous (poly)phenolic groups, the treatments that best preserved
316 flavonol group, when comparing with the non-thermal freeze-drying method, were hot
317 air-drying and purée pasteurization.

318 Regarding anthocyanins, the most distinctive (poly)phenolics in red-fleshed
319 apple cultivars, cyanidin-3-*O*-galactoside and cyaniding arabinoside were detected in all
320 the apple samples (**Table 1**). According to the literature,^{9,27,29} cyanidin-3-*O*-galactoside
321 was the most abundant anthocyanin detected and quantified in the red-flesh apple
322 samples (around 90% of the total anthocyanin content). Concerning the impact of the
323 different treatments, results showed that anthocyanins were degraded to a higher extend
324 compared to other (poly)phenolic compounds. Results showed that freeze-drying is the
325 technique that statistically best preserves these compounds (285 mg/kg d.w) followed

326 by hot air-drying (73.2 mg/kg d.w.) and purée pasteurization (26.2 mg/kg d.w.). In the
327 infrared-dried samples, these compounds were degraded almost completely with
328 observed losses of > 98% when comparing with freeze-drying.

329 Other groups of (poly)phenols influenced by the processing technique applied
330 were the flavanones and dihydrochalcones (**Table 1**). In the flavanone group, only
331 naringenin-glucoside and eriodictyol-hexoside were detected as reported in the literature
332 ²⁶ and the processes that statistically best preserved them were freeze-drying and purée
333 pasteurization with total values of 15.7 and 15.2 mg/kg d.w., respectively. Within the
334 group of the dihydrochalcones, phloretin-xylosil-glucoside was the predominant
335 compound. At a general level, this (poly)phenol group is the least affected by the apple
336 transformation, and the total values of non-thermal (freeze-drying), and some thermal
337 (hot air-drying, infrared-drying at 40°C and pasteurized purée) treatments show no
338 statistically significant differences. Our results are in accordance with a previous
339 study,³⁰ confirming that apple dihydrochalcones are more stable against the application
340 of high temperatures than other (poly)phenol classes, such as flavan-3-ols and
341 anthocyanins.

342 Considering the freeze-dried samples as reference, infrared-drying (at all
343 temperatures) caused significant ($p < 0.05$) losses in most of the apple (poly)phenols,
344 while (poly)phenol losses by hot air-drying and by the purée pasteurization were
345 considerably lower. This fact is probably a consequence of the intense time/temperature
346 treatment applied in the infrared processes and to a lesser extent in hot air-drying or in
347 the production of the pasteurized purée, in which the treatment at high temperatures
348 lasts only a few minutes. Thermal treatment can cause severe degradation of
349 (poly)phenolic compounds as it is well known that these compounds are temperature
350 sensitive.¹⁸⁻²² This fact is also reflected in infrared-dried samples where those subjected

351 to higher temperatures (50°C and 60°C) showed the greatest phenol degradation/losses.
352 Moreover, the losses could also be due to the presence of oxygen in the case of infrared-
353 drying and hot air-drying producing an oxidative degradation of the (poly)phenols and
354 consequently the browning of the apple samples (**Figure 1**). (Poly)phenols are the
355 desirable substrates of oxidoreductive enzymes such as phenoloxidases, whose main
356 function is to oxidized phenols.^{18,28} These enzymes catalyze the oxidation of *o*-
357 diphenols into quinones, which polymerize to form brown melanin pigment.³¹ In
358 freeze-dried samples, these reactions do not occur because the freezing and subsequent
359 sublimation of water under vacuum conditions prevents the action of these enzymes and
360 consequently the browning of the samples.³² However, although this process efficiently
361 preserves bioactives, freeze-drying costs can be 2-5 times higher than hot air-drying in
362 order to achieve the same final moisture content. This fact justifies the need to find
363 alternatives for the production of economically affordable healthy food for everyone.³³
364 In this sense, when comparing the different dehydration techniques for preparing
365 healthy apple snacks preserving the content of (poly)phenols (**Figure 2**), infrared-drying
366 showed high losses of 57-74% depending on the temperature. On the other hand, hot
367 air-drying and purée pasteurization allow the maintenance of 83% and 65% respectively
368 of (poly)phenols quantified in the freeze-dried snacks and may be good alternatives to
369 the costly freeze-drying technique.

370

371 **3.2. Effect of apple processing on the phenolic bioavailability: human pilot** 372 **study**

373 The second aim of the present work was to study how apple processing impacts
374 on the release and absorption of (poly)phenols present in the apple products and
375 determine the optimal processing technique that improves (poly)phenol bioavailability.

376 For this, the apple products with the highest (poly)phenolic contents (freeze-dried
377 apples, hot air-dried apples and pasteurized apple purée) were chosen to conduct a
378 crossover acute intervention study with three subjects. The amounts of each product
379 administered to the subjects were selected to match as best as possible the total content
380 of phenolic and the contents of the different phenolic subclasses (**Supplemental Table**
381 **1**).

382

383 *3.2.1. Effect of apple processing on phenolic compounds bioavailability*

384 A total of 59 phenol metabolites were identified and quantified in urine samples
385 after the intake of the three red-fleshed apple derived products, including derivatives of
386 benzoic acids, phenylpropionic acids, phenylvalerolactones and catechols (major
387 groups) (**Figure 3 a, b, c and Supplemental Table 3**), and anthocyanins, flavan-3-ols,
388 dihydrochalcones and hydroxycinnamic acids (minor groups) (**Figure 3 d, e, f and**
389 **Supplemental Table 3**). The phenolic metabolites were mainly phase-II sulfated,
390 glucuronided and/or methylated conjugates of parent compounds present in the apples
391 as well as of microbial catabolites resulting from colonic degradation (**Supplemental**
392 **Table 4**). Excretion kinetics for each of these (poly)phenolic groups expressed as
393 $\mu\text{mol/h}$ are shown in **Supplemental Figure 1** and **Supplemental Figure 2**.

394 Our data show intra-individual, inter-individual and processing-derived
395 differences among the three products. Large inter-individual differences in urine
396 metabolite concentration were found between the three volunteers, with highest
397 excretion found in volunteer 1 (V1) followed by volunteer 3 (V3) and volunteer 2 (V2)
398 (**Figure 3 and Supplemental Table 3**). This trend was repeated in each of the three
399 apple products (**Supplemental Table 3**) and for almost all phenolic groups (except for
400 anthocyanins which are discussed separately).

401 Regarding the processing effect, pasteurized purée showed significantly higher
402 total (poly)phenol bioavailability in all three volunteers, followed by the hot air-dried
403 red-fleshed apple and finally, the freeze-dried red-fleshed apple showed the lowest.
404 **Figure 4** shows the % excretion of (poly)phenols in urine, which have been calculated
405 as the average between the three volunteers comparing the sum of (poly)phenols
406 ingested and the sum of the moles excreted in 24 h urine. This was observed for almost
407 all (poly)phenolic groups except for anthocyanin and flavan-3-ols derivatives. It is of
408 interest to note that although freeze-drying is the method that best preserved
409 (poly)phenolic compounds during processing, it was the product with the lowest %
410 bioavailability, while pasteurized purée, with the highest losses during processing,
411 showed the highest % bioavailability.

412 The results obtained corroborate the notion that the apple processing could
413 enhance (poly)phenols bioavailability. Chemical structure, concentration and matrix
414 interactions are the three basic pillars that govern bioaccessibility and bioavailability of
415 (poly)phenolic compounds from fruits. It has been shown that food processing can
416 influence and alter all these three factors.³⁴ Regarding thermal processing, it has been
417 reported that it can promote the (poly)phenol release disrupting intracellular barriers,
418 thus modulating (poly)phenol bioaccessibility for example in tomatoes,²⁴ in which it
419 was found that human plasma levels of naringenin and chlorogenic acid, increased
420 several times after the intake of cooked tomatoes compared with levels observed after
421 consumption of fresh tomatoes. This fact was also verified in a recent study³⁵ in which
422 authors observed that after the administration of similar doses of flavan-3-ols through
423 three apple products (phenolic extract, raw apple or apple purée) in
424 hypercholesterolemic pigs, higher number of genes were modulated by purée than raw
425 apples, which suggests that the processing of apples into purée increased the

426 bioavailability of some phytochemicals, as flavan-3-ols, that could contribute to the
427 postprandial nutrigenomic response.

428 Apart from this, it has also been shown that liquid foods possess a lower
429 viscosity and pass through the stomach more rapidly than solid food. This is due to a
430 higher water content, and typically lower content in proteins or complex carbohydrates
431 that may bind to (poly)phenols.³⁶ Several studies have reported rapid absorption of
432 (poly)phenols from liquid foods such as coffee³⁷ or apple products.³⁸ This latter study
433 reported 3-fold higher quercetin plasma concentration after consumption of apple sauce
434 than after vacuum-impregnated apple chips or apple peel extract capsules. Moreover, in
435 this study the different treatments also resulted in a high inter-individual variability of
436 all plasma pharmacokinetic parameter after equal intake of quercetin equivalents of the
437 apple product types.³⁸

438 The high inter- and intra-individual variation in the response to (poly)phenolic
439 intake, which in many cases leads to contradictory results in human trials, could result
440 from intrinsic aspects such as genetics, age, sex and physiological or pathological states,
441 in addition to the matrix effect already mentioned. For example, it has been shown that
442 genetic polymorphisms between individuals affect the efficacy of bioactive
443 compounds,³⁹ since they differentially affect genes that encode enzymes involved in the
444 metabolism of these compounds.

445 Moreover, only small amounts of ingested (poly)phenols are absorbed in the
446 upper gastrointestinal tract (GIT), while most compounds reach the lower GIT
447 unmodified,⁴⁰ where they undergo extensive metabolism mediated by the colonic
448 microbiota. Thus, a key role is played by the gut microbiota, which may modify the
449 structure of (poly)phenols, releasing lower molecular weight colonic catabolites that can
450 be absorbed more easily.⁴⁰ Each human has a unique gut microbiota that changes

451 throughout life, and the environment, diet and lifestyle, all influence the
452 microbiome.^{41,42} In this sense, the existence of metabotypes (metabolic phenotype with
453 specific gut microbiome-derived metabolites that characterize the metabolism of the
454 parent compound) in the production of phenolic metabolites has been discussed almost
455 exclusively in recent year. It was observed that the benefits associated with the ingestion
456 of foods rich in ellagitannins, such as pomegranate and walnuts may be related to
457 specific metabotypes.⁴³⁻⁴⁵ In our study, most of the apple (poly)phenol metabolites are
458 of colonic origin while only a minority were parent compounds and their phase II
459 conjugates from upper GIT absorption (**Supplemental Figure 3 and Supplemental**
460 **Table 4**). Furthermore, despite having only three volunteers in our study, “low” (V2)
461 and “high” (V3) excretors can be observed. It should be stressed that although V2
462 showed in all three products the lowest (poly)phenol bioavailability, this was greatly
463 improved after the ingestion of apple purée.

464 **Supplemental Figure 3** shows the representation of the phenolic groups
465 detected after each ingestion for each volunteer, separated into those absorbed in the
466 upper GIT (dihydrochalcone, flavan-3-ol, and anthocyanin parent compounds and their
467 phase II metabolites) and in the lower GIT (simple phenolic acids). The major groups of
468 upper GIT absorption were dihydrochalcone and flavan-3-ol derivatives while
469 phenylpropionic acid derivatives were the major ones among the metabolites that are
470 absorbed in the lower GIT. Similar results were observed in our previous study in which
471 ten volunteers ingested 80g of freeze-dried red-fleshed apple snack.²⁵

472 It is of note that in the present study we observed intra-individual differences in
473 the proportions of the different metabolite groups after the ingestion of the three apple
474 products. If the observed differences depend on the apple processing or, more probable,
475 on intrinsic aspects of each individual should be the focus of further studies.

476 The inter-individual differences are also shown here, since for V1 and V3 the
477 (poly)phenolic groups that are absorbed in the upper GIT (dihydrochalcones, flavan-3-
478 ols and anthocyanins) are greater in the freeze-dried snack, while for V2 it is greater in
479 the hot air-dried red-fleshed snack (**Supplemental Figure 3**). Interestingly, in no case
480 was the bioavailability of the (poly)phenols absorbed in the upper GIT greater after
481 ingesting the purée as might be expected, since in the solid apple matrix of the freeze-
482 dried snack and the hot air-dried snack these (poly)phenolics are bound to cell walls by
483 covalent bonds between (poly)phenolics and polysaccharides possibly restricting
484 bioavailability in the small intestine.⁴⁶ Besides, genetic variation between individuals for
485 enzymes involved in the absorption and metabolism of these groups in the gut
486 epithelium and / or liver may result in large differences in the expression of a functional
487 enzyme which might explain the observed inter-individual differences.⁴⁷⁻⁴⁹

488 Regarding the metabolites found after the ingestion of the three apple products,
489 the majority were sulphate and methyl-sulphate conjugates representing between 61%-
490 83% and 7-27% of the total (poly)phenol metabolites detected, respectively (**Figure 5**
491 **and Supplemental Table 4**). The rest of the (poly)phenol metabolites (glucuronide,
492 methyl, glycine, sulphate-glucuronide and methyl-glucuronide conjugates, and free
493 acids/parent compounds) varied considerable between volunteers and between the type
494 of the ingested apple product showing again great inter- and intra-individual differences.
495 The differences in human subjects' genetics regarding digestive enzymes, intestinal
496 transporters, phase I and II metabolism or tissue carriers, and also differences in gut
497 microbiota composition and functionality affecting the catabolism of the not absorbed
498 (poly)phenols in the small intestine being responsible of the differences observed.⁵⁰

499 Although it is a preliminary study, our bioavailability results show large
500 differences in the concentrations of metabolites (inter-individual differences) as well as

501 in the metabolic profile depending on the type of product ingested for each person
502 (intra-individual differences), which could result in different effects on health. This fact
503 justifies the need to carry out future studies with a greater number of volunteers to be
504 able to address these differences. Nevertheless, an important limitation of this study is
505 the lack of some of the authentic standards to quantify more accurately the
506 concentrations of some (poly)phenol metabolites and thus the real bioavailability % of
507 red-fleshed apple products.

508

509 *3.2.2. Effect of apple processing on anthocyanin bioavailability*

510 In this study, the effect of apple processing on anthocyanin bioavailability was
511 of special interest since this phenolic group is the most characteristic of the red-fleshed
512 apple cultivars and is not found in the pulp of any common white-fleshed apple variety.
513 This (poly)phenol group was however, the most affected by the processing treatments
514 (**Table 1**) and, consequently, different amounts of anthocyanins were consumed (2, 4
515 and 16 mg) after the acute intake of pasteurized purée, hot air-dried and freeze-dried
516 apple products, respectively (**Supplemental Table 1**). The results showed (contrary to
517 what happens for the other (poly)phenolic groups, **Figure 4**) that a higher average
518 excretion in urine of anthocyanins was observed after the intake of hot air-dried apple
519 (0.07%) while for the freeze-dried apple and pasteurized purée it was 0.04% and 0.03%,
520 respectively (data not shown). These excretion rates have been calculated as the average
521 of the three volunteers and comparing the total ingested anthocyanins with the total
522 anthocyanins and their phase II metabolites excreted in 24 h-urine. It should be noted
523 that these observed differences are very small and probably, if more volunteers had
524 participated in this study, the differences would had shown no significant differences
525 between products. Moreover, although the bioavailability of anthocyanins is very low

526 compared to other (poly)phenolic groups, most of these compounds pass to the colon
527 where they are degraded by the microbiota to simpler phenolic acids that are common to
528 other (poly)phenol groups and once reabsorbed they contribute to the pool of circulating
529 phenolic metabolites in the body.^{48,51}

530 The fact that anthocyanins were more bioavailable in the solid matrices (hot air-
531 dried and freeze-dried) than in the pureé, may be due to the fact that these compounds
532 are very unstable and more easily degraded. Thus, in the solid matrix, they could remain
533 more attached to the fiber that may stabilize them or offer protection against further
534 reactions until the site of absorption is reached. Our results could be in agreement with a
535 previous study reporting that when raspberry extract was digested *in vitro* with
536 foodstuffs (bread, breakfast cereals or ice cream) higher proportions of anthocyanins
537 were bioaccessible compared to the extract digested alone.⁵²

538 Finally, **Supplemental Figure 2** shows the urine excretion kinetics of
539 anthocyanins after the apple products intake expressed as total nmols of anthocyanins
540 and their phase II metabolites excreted per hour. The higher concentration observed in
541 the freeze-dried format is due to the fact that the anthocyanin dose administered with
542 this product was higher. In all cases, similar kinetics were observed with a maximum
543 excretion between 2-4 hours, which is in agreement with previous *in vivo* studies
544 reporting that anthocyanins are absorbed in the stomach and the small intestine with
545 rapid detection of intact anthocyanin glycosides in urine and plasma within 30 to 60 min
546 of ingestion.^{48,51}

547

548 **4. CONCLUSIONS**

549 Our findings revealed that, considering the freeze-drying as a reference
550 technology to preserve food bioactives, infrared-drying at all temperatures caused

551 significant losses in all the red-fleshed apple (poly)phenols, while (poly)phenol losses
552 by hot air-drying and the purée pasteurization were considerably lower. Anthocyanins in
553 particular were degraded to a higher extend after all thermal processing technologies.
554 So, we conclude that for obtaining red-fleshed apple products affordable for the
555 consumer, hot air-drying and purée pasteurization represent interesting technologies to
556 obtain apple products with a high shelf-stability maintaining at the same time a high
557 concentration of bioactive (poly)phenols. However, to obtain a product with the highest
558 anthocyanin content, the extra cost of freeze-drying would have to be assumed.

559 Results obtained from the human postprandial crossover study showed that when
560 comparing the total ingested (poly)phenol dose and the urinary excreted amount, the
561 pasteurized apple purée proved to be the processing with the highest bioavailability,
562 followed by the hot air-dried apple and the freeze-dried apple. Further, a great intra- and
563 inter-individual variability between the metabolites was found, which highlights the
564 importance of characterizing the metabotypes in future studies.

565 The present study is a proof of concept to select the most appropriate apple
566 processing to preserve the apple (poly)phenolic compounds and provides further
567 evidence on how food processing plays a significant role in the bioavailability of
568 (poly)phenols, which is a step forward towards the design of healthier foods.

569

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581

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779 **Figure Captions**

780 **Figure 1.** Different red-fleshed apple products used in this study obtained by: a) freeze-
781 drying, b) hot air-drying, c) infrared-drying d) or by a process of purée pasteurization.

782

783 **Figure 2.** Impact of the different thermal and non-thermal processes on the
784 (poly)phenolic groups in red-fleshed apple products. The letters express statistically
785 significant differences between the content of the total (poly)phenol classes content
786 among the processing technologies ($p < 0.05$).

787

788 **Figure 3.** Total major and minor metabolite excretion after intake of freeze-dried apple,
789 hot air-dried apple and apple pasteurized purée. Data expressed as mean values \pm
790 standard deviation. Different lowercase letters: indicates differences between volunteers
791 in excretion of total major or minor metabolites after the intake of freeze-dried red-
792 fleshed apple. Different capital letters: indicates differences between volunteers in
793 excretion of total major or minor metabolites after the intake of hot air-dried red-
794 fleshed apple. Different numbers: indicates differences between volunteers in excretion
795 of total major or minor metabolites after the intake of red-fleshed apple pasteurized
796 purée. The symbols *, +, and # indicate differences between the 3 intakes for the same
797 volunteer (One-way ANOVA, Tukey's test between all means, $p < 0.05$).

798

799 **Figure 4.** % Excretion in urine of total (poly)phenols after the intake of freeze-dried
800 red-fleshed apple, hot air-dried red-fleshed apple and red-fleshed apple pasteurized
801 purée (n=3). This % was calculated as the ratio of total moles of excreted
802 (poly)phenolic metabolites with respect to the total moles of ingested phenolic

803 compounds. Different letters indicate significant differences in excretion between
804 products (General Linear Model, and One-way ANOVA, $p < 0.05$).

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806 **Figure 5.** Schematic representation (% of each group over the total) of the main phase
807 II metabolites, free acids and parent compounds found in urine in each volunteer after
808 the intake of freeze-dried red-fleshed apple, hot air-dried red-fleshed apple and red-
809 fleshed apple pasteurized purée.

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