

ABSTRACT

22
23 Aim of the study was to evaluate color, total polyphenol content (TPC), antioxidant capacity
24 (ABTS, FRAP, DPPH), reducing sugars and heat damage (furosine, hydroxymethylfurfural,
25 glucosylisomaltol) of 21 commercial powder products obtained from South-American fruits
26 (mesquite, lucuma, camu camu), seeds (amaranth, purple maize), roots and tubers (yacon, maca,
27 mashua, tocosh), bark (cat's claw) and leaves (graviola). TPC and antioxidant capacity were
28 maximum in camu camu and cat's claw powders, and minimum in tocosh, amaranth, lucuma and
29 maca; graviola, mashua, purple maize and mesquite **are raw materials with unique antioxidant**
30 **properties that stand out from the others**. Yacon, mashua and lucuma powders had high reducing
31 sugars content (40.9, 34.4 and 21.2 g/100 g DM, respectively) and heat damage (HMF 146.6 mg/kg,
32 furosine 2399.8 and 2228.4 mg/100 g protein, respectively). Overall, camu camu powder and cat's
33 claw were the most interesting products, having high levels of total polyphenols and antioxidant
34 capacity together with very low heat damage.

35 **1. INTRODUCTION**

36 A major threat to human wellbeing is the oxidative stress, an “imbalance between oxidants and
37 antioxidants in favour of the oxidants” (SIES, 1997), which can lead to cellular damage and
38 facilitate the insurgence of cardiovascular and neurodegenerative diseases, diabetes mellitus, cancer
39 and inflammatory illness (UTTARA *et al.*, 2009). An effective approach to prevent oxidative stress
40 is to include in the daily diet products rich in antioxidants **which can quench the oxygen free**
41 **radicals, preventing** the oxidation of the cell membrane.

42 Plants and plant-derived ingredients have been used as medical remedies from prehistoric ages, and
43 still are a major source of health-promoting elements. In recent years, the interest in the
44 identification and utilization of plants rich in antioxidant compounds to limit the oxidative stress
45 (ALMEIDA *et al.*, 2011; KRISHNAIAH *et al.*, 2011) has been steadily growing, because they may
46 behave as preventive medicine. Several authors have reviewed the beneficial uses of underexploited
47 and little-known plant species used in food production but also in traditional medicine (e.g. BIEL *et*
48 *al.*, 2017; CAMPOS *et al.*, 2013; CHIRINOS *et al.*, 2013; CONTRERAS-CALDERÓN *et al.*,
49 2011; KRISHNAIAH *et al.*, 2011). Peru, thanks to its widely diversified climatic zones, is home to
50 a broad array of endemic plants, which show huge differences in the content and type of nutrients
51 and that are potential sources of valuable bioactive compounds (CAMPOS *et al.*, 2018).

52 The antioxidant capacities of plant-derived products vary depending on their content in
53 polyphenols, vitamin C, tocopherols and carotenoids, (SAURA-CALIXTO and GOÑI, 2006), as well as
54 on the different processing conditions. While in some cases the plant products are consumed fresh,
55 most often they undergo some type of transformation and/or drying to improve shelf life, to lower
56 transport costs and to reach far off consumers (CINAR, 2018). Accordingly, powders from South-
57 American plants with known health-promoting features (Supplementary Table 1) are manufactured
58 by several industries to find new market niches and to foster the consumption of health-promoting
59 natural products. These innovative powder products, obtained from fruits (mesquite, lucuma and
60 camu camu), seeds (amaranth and purple maize), roots and tubers (yacon, maca, mashua and
61 tocosh), bark (cat’s claw) and leaves (graviola), are currently used for the preparation or enrichment
62 of infusions, juices, shakes/smoothies, yogurts, desserts, as well as ingredients in cosmetic and
63 pharmaceutical recipes.

64 The aim of our study was to evaluate some **characteristics** of these powder products for their
65 possible utilization as enhancing ingredients in wheat-based oven products. To achieve this goal, 21
66 commercial powder samples of the above-mentioned species were assessed for color, total
67 polyphenol content, antioxidant capacity, reducing sugars and heat damage.

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69 **2. MATERIALS AND METHODS**

70 **2.1. Samples**

71 The powders analyzed were acquired in 2016 at an industrial fair dedicated to Peruvian export
72 products (Expoalimentaria, Lima, Peru; www.expoalimentariaperu.com) except amaranth, obtained
73 from the Peruvian market, and two maca samples, bought from the Italian market. Several samples
74 (3-5) of each powder product were collected. A detailed list of the products tested is presented in
75 Table 1.

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77 **2.2. Physical and chemical analyses**

78 **2.2.1. Color**

79 The color coordinates L^* (luminosity), a^* (red-green) and b^* (yellow-blue) of the samples were
80 scored with a tristimulus colorimeter (Chroma meter CR-300, Minolta Italia S.p.A., Italy) using the
81 standard-white reflector plate and illuminant C. Four measurements for each sample were
82 performed.

83 Table 1. Samples analyzed: species, brands, codes, average dry matter and protein contents (g/100
84 g).

Product	Species	Brand	Sample	Dry	Protein
<i>Bark</i>					
Cat's claw	<i>Uncaria tomentosa</i> L.	A	Cat's claw 1	91.8	0.3
Cat's claw bio	<i>Uncaria tomentosa</i> L.	B	Cat's claw 2	92.8	2.7
Cat's claw tea	<i>Uncaria tomentosa</i> L.	B	Cat's claw tea	92.5	3.0
<i>Seeds</i>					
Amaranth flour	<i>Amaranthus caudatus</i> L.	C	Amaranth FR	90.4	11.5
Amaranth flakes	<i>Amaranthus caudatus</i> L.	D	Amaranth FS	90.4	8.8
Purple maize	<i>Zea mays</i> L.	B	Purple maize	90.3	7.0
<i>Roots</i>					
Yacon	<i>Smallanthus sonchifolius</i>	B	Yacon	87.8	1.8
Maca gluten free	<i>Lepidium meyenii</i> Chacon	E	Maca 1	85.9	10.6
Maca bio	<i>Lepidium meyenii</i> Chacon	B	Maca 2	87.3	9.0
Maca HP	<i>Lepidium meyenii</i> Chacon	B	Maca 3	90.3	8.5
Maca extract	<i>Lepidium meyenii</i> Chacon	A	Maca 4	85.4	9.3
Maca	<i>Lepidium meyenii</i> Chacon	A	Maca 5	86.6	7.7
Maca root	<i>Lepidium meyenii</i> Chacon	F	Maca 6	92.9	12.0
Maca energia	<i>Lepidium meyenii</i> Chacon	G	Maca 7	90.4	7.0
<i>Tubers</i>					
Tocosh	<i>Solanum</i> spp.	H	Tocosh	83.4	2.2
Mashua	<i>Tropaeolum tuberosum</i> Ruiz &	E	Mashua	84.7	9.0
<i>Leaves</i>					
Graviola bio	<i>Annona muricata</i> L.	B	Graviola	91.9	10.8
Graviola tea	<i>Annona muricata</i> L.	B	Graviola tea	91.0	11.0
<i>Fruits</i>					
Mesquite	<i>Prosopis</i> spp.	B	Mesquite	89.3	8.7
Lucuma	<i>Pouteria lucuma</i> Ruiz & Pav.	B	Lucuma	88.9	3.4
Camu camu	<i>Myrciaria dubia</i> (Kunth)	B	Camu Camu	87.4	5.4

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86 2.2.2. Dry matter and protein content

87 Dry matter was determined following the gravimetric method, drying 2 g of product at 130 °C for
88 90 min; protein content was assessed by Kjeldahl (N x 6.25). These and all the following analyses
89 were performed in triple.

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91 2.2.3. Samples preparation for total polyphenols content and antioxidant capacity analysis

92 All the reagents, of analytical grade, were purchased from Sigma-Aldrich Co. (Milan, Italy). Two
93 different solvents were tested for the extraction of total polyphenols and the evaluation of the
94 antioxidant capacity, i.e. ethanol:H₂O (EtOH:H₂O; 80:20) and methanol:H₂O:acetic acid
95 (MeOH:H₂O:acet; 50:42:8).

96 Exactly 0.15 g of powdered product were weighed in 2 mL tubes and subjected to three extractions,
97 adding 1 mL of an EtOH:H₂O solution each time. In the first extraction, the samples were stirred
98 with a Vortex (Reax 2000, Meindolph Heidolph, Schwabach, Germany) for 1 min and sonicated
99 (F5200b, Decon, UK) twice for 20 min; in the second extraction the samples were stirred with a
100 Vortex for 1 min, an orbital shaker (Multi-Rotator GRANT-BIO, Cambridge, UK) for 20 min and
101 sonicated for another 20 min; in the third extraction the samples were stirred with a Vortex for 1
102 min and sonicated for 5 min. After each extraction, the samples were centrifuged with a 4224
103 centrifuge (ALC Apparecchi per Laboratori Chimici Srl, Milan, Italy) for 5 min at 8048 g and all

104 the supernatants were mixed in a single tube. The extractions were performed at 10 °C and away
105 from light as far as possible.

106 Following the same procedures, 0.3 g of powdered product underwent three extraction cycles,
107 adding respectively 1.5, 1.5 and 1.0 mL of a MeOH:H₂O:acet solution.

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109 **2.2.3.1. Total polyphenol content**

110 Total polyphenol content (TPC) in samples extracted with EtOH:H₂O and MeOH:H₂O:acet was
111 assessed with the Folin-Ciocalteu method as described by BRANDOLINI *et al.* (2013) using a Du-
112 62 Beckman spectrophotometer (Beckman Coulter, Nyon, VD, Switzerland). The TPC, in mg gallic
113 acid equivalent (GAE)/kg DM, was computed from a reference curve obtained from six gallic acid
114 concentrations (range: 0-150 mg/L).

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116 **2.2.3.2. Assessment of antioxidant capacity using the ABTS method**

117 The 2,2'-azino-bis(3-ethylbenzothiazoline-6-sulphonic acid (ABTS) radical scavenging capacity
118 was analysed as described by YILMAZ *et al.* (2015). A stable stock solution of the ABTS radical
119 cation was prepared by reacting 10 mL of an aqueous solution of 2-2'-azinobis-3-
120 etilenbenzotiazoline 7 mM and 176 µL potassium persulfate 140 mM in the dark at room
121 temperature for 12-16 h. The EtOH:H₂O or MeOH:H₂O:acet extracts (150 µL) were reacted with 5
122 mL of a diluted ABTS radical solution in ethanol (absorbance: 0.70±0.02 AU at 734 nm); the
123 absorbance was measured at 734 nm, after 6 min at 30 °C, with a V650spectrophotometer (Jasco,
124 Japan), using ethanol as blank. The antioxidant capacity was evaluated as percentage of absorbance
125 decrease (inhibition percentage). A reference curve was built with 11 concentrations (from 0.05 to
126 0.72 mM) of Trolox. The results are expressed as mmol Trolox equivalents (TE)/kg DM.

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128 **2.2.3.3. Assessment of the reduction power using the FRAP method**

129 The ferric reducing antioxidant power (FRAP) was determined as described by YILMAZ *et al.*
130 (2015). Briefly, 200 µL of EtOH:H₂O or MeOH:H₂O:acet extracts were mixed with 4.5 mL FRAP
131 reagent. Absorption was measured with a V650 spectrophotometer (Jasco, Japan) at a wavelength
132 of 593 nm after 60 min incubation at 37 °C; acetate buffer 0.3 M pH 3.6 was used as blank. The
133 FRAP reagent, prepared daily, consisted of 0.3 M acetate buffer (pH 3.6), 10 mM 2,4,6-Tris(2-
134 pyridyl)-s-triazine (TPTZ) in 40 mM HCl and 20 mM FeCl₃ (10:1:1 v/v/v). FRAP values were
135 obtained by comparing the results to a calibration curve built with 18 concentrations (0.06 - 0.90
136 mM) of Trolox. The antioxidant capacity was expressed as mmol TE/kg DM.

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138 **2.2.3.4. Assessment of antioxidant capacity using the DPPH method**

139 The 2,2-diphenyl-1-picrylhydrazyl (DPPH) radical cation scavenging capacity of EtOH:H₂O and
140 MeOH:H₂O:acet extracts was evaluated according to BRANDOLINI *et al.* (2013) using a DU-62
141 spectrophotometer (Beckman, USA). For each extract five different dilutions were analysed. A
142 dose-response line was computed for each sample and the powder quantity needed to scavenge 50%
143 of the radical (I50) was determined. A reference regression line was computed for the antioxidant
144 Trolox, with concentrations between 3 and 50 µM. The antioxidant capacity was expressed as ratio
145 between I50 of Trolox and I50 of the sample, i.e. mmol TE/kg DM.

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147 **2.2.4. Sugars content**

148 Fructose, glucose, maltose and sucrose were assessed by HPLC, following HIDALGO and
149 BRANDOLINI (2011). For peak quantification, sugars calibration curves were constructed using 15
150 different concentrations (between 0 and 155 mg/L) of fructose, 19 different concentrations (between
151 0 and 428 mg/L) of glucose, 19 different concentrations (between 0 and 385 mg/L) of maltose, and
152 15 different concentrations (between 0 and 153 mg/L) of sucrose standards (Sigma, St. Louis, MO,
153 USA). The calibration curves, after log transformation, were linear ($r^2 = 1.00$; $p \leq 0.001$) in the
154 concentration ranges considered. The results are reported as g/100 g DM.

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2.2.5. Heat damage indices.

Furosine was determined by HPLC as described by HIDALGO and BRANDOLINI (2011). A calibration curve was built using nine different concentrations (between 0.33 and 5.13 $\mu\text{mol/L}$ of furosine dihydrochloride (NeoMPS, PolyPeptide Laboratories, Strasbourg, France) in 3 N HCl. The calibration curve was linear ($r^2 = 1.00$; $p \leq 0.001$) in the concentration ranges considered. The results are expressed as milligrams of furosine/100 g of protein.

Hydroxymethylfurfural (HMF) and glucosylisomaltol (GLI) were determined following the HPLC method of RUIÁN-HENARES *et al.* (2008) as described by HIDALGO and BRANDOLINI (2011). For peak quantification, a calibration curve was constructed using 13 different concentrations (between 0 and 6.25 mg/L) of HMF (Safc, St. Louis, MO, USA). The calibration curve was linear ($r^2 = 1.00$; $p \leq 0.001$) in the concentration range considered. GLI quantification was computed considering the response factor of HMF at 280 nm. The results are expressed as mg/kg DM.

2.3. Statistical analysis

The data were processed by one-way analysis of variance (ANOVA) considering the samples as factors. The distribution of the data was checked and, for normalization purposes, L^* and b^* values were squared, while the other parameters were \log_{10} -transformed; however, for easier comprehension, in Tables and Figures the original data are reported.

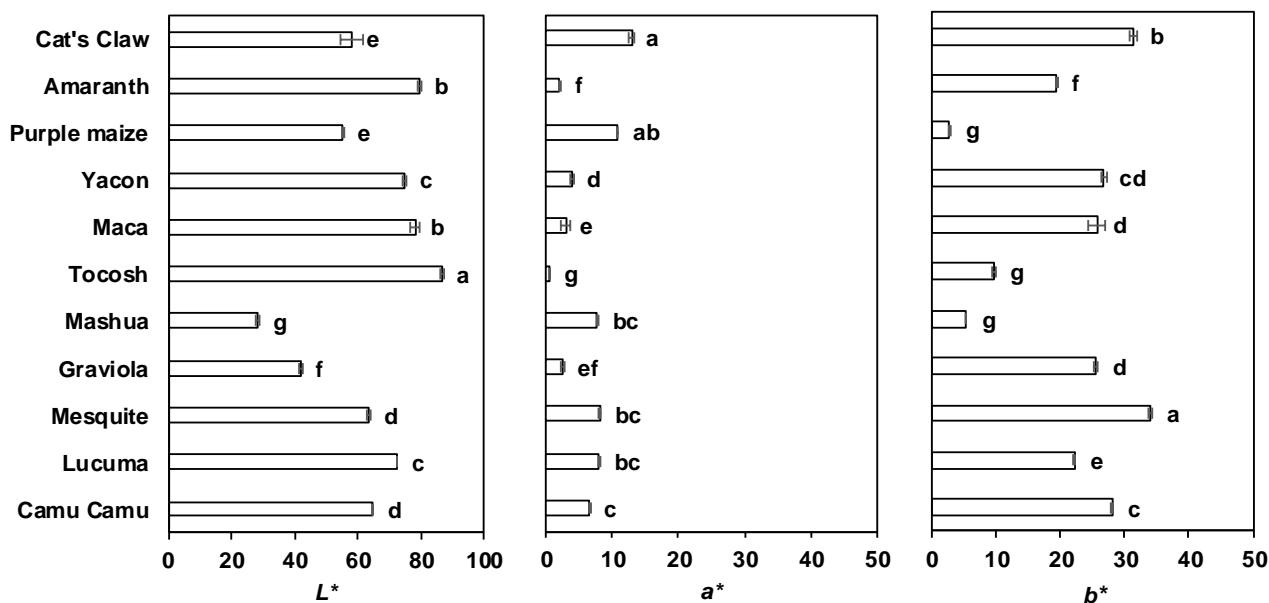
When significant differences were found ($p \leq 0.05$), Fisher's lowest significant difference (LSD) was computed at a 95% significance level. To compare the results of the two solvents used for the preparation of the extracts, the t-test was applied ($p \leq 0.05$). ANOVA, LSD test and t-test were conducted using the statistical program STATGRAPHICS® Centurion. Mean, standard error and coefficient of variation were computed using the program Excel (Microsoft® Office Excel 2007). Principal Components Analysis (PCA), performed considering the mean values of the 21 samples and all the parameters, was carried out with the software The Unscrambler X 10.2 (CAMO software AS, Norway).

3. RESULTS AND DISCUSSION

3.1. Powders color

Supplementary Table 2 shows the average values and the results of the LSD test for the color coordinates L^* , a^* , b^* of the twenty-one samples. The results obtained grouping the samples by species are reported in Fig. 1.

The broad heterogeneity of the samples led to an ANOVA (not presented) showing significant differences for all the parameters. In fact, a preliminary visual control gave the following color characterization: mashua and purple corn were purple; maca, yellow-orange; cat's claw, mesquite and yacon, orange; graviola, green-brown; camu camu, yellow-brown; tocosh, white; amaranth, cream-white; lucuma, ocher.



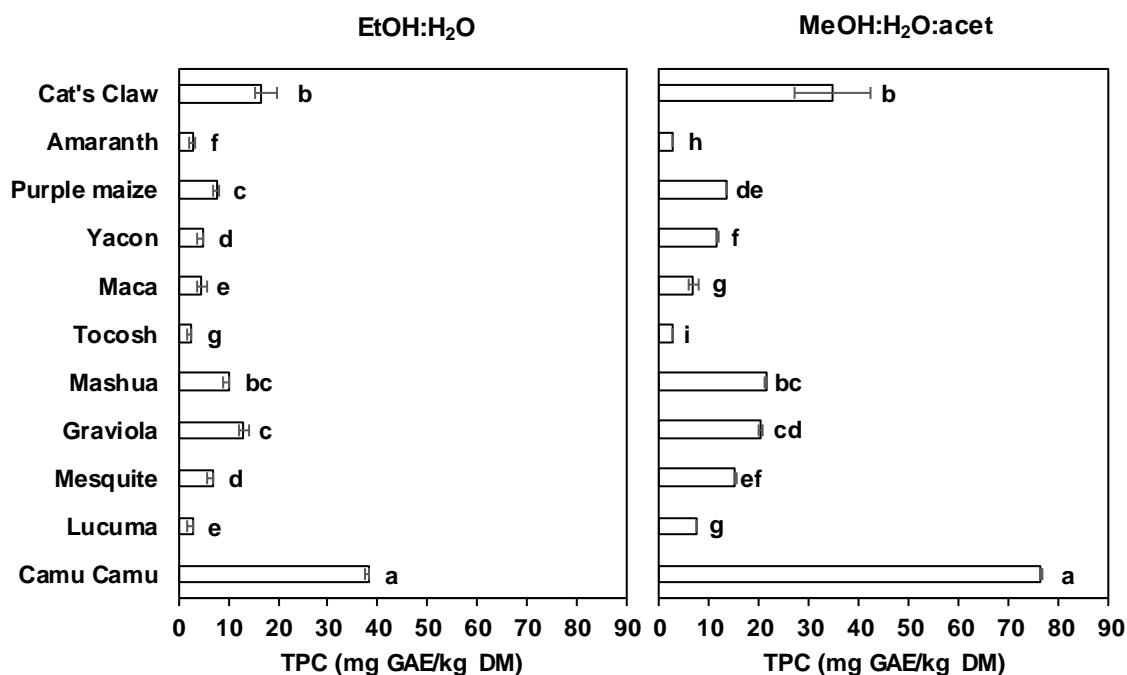
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196 **Figure 1.** Colour coordinates (L^* , a^* , b^*) of powdered products from 11 species. Different letters
 197 indicate significant differences (LSD, $p \leq 0.05$) among species.
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199 The tocosh powder was the brightest (L^* : 86.6), followed by most maca samples (76.2-83.9) and
 200 amaranth (78.5-80.3). One maca (maca 4) had an L^* of 72.0, lower than the other maca samples.
 201 Camu camu had a L^* like the lyophilized samples (60.45 ± 2.78) and higher than the spouted bed
 202 dried samples (36.6-40.8) described by FUJITA *et al.* (2013). Overall, mashua presented the lowest
 203 brightness (28.3), followed by graviola (43.3-41.0). Cat's claw and purple corn scored the highest
 204 a^* red component values (12.1-13.5 and 10.8, respectively), while tocosh presented the lowest
 205 (0.5). The variation among the different maca samples was quite limited, ranging from 1.2 (Maca 7)
 206 to 5.1 (Maca 4). Mesquite presented the highest b^* yellow component (34.1), followed by cat's
 207 claw (on average 31.6), five maca samples (22.2-26.2) and graviola (on average, 25.6); maca 1 and
 208 maca 4 had values different from the other maca (28.3-30.2). Purple corn presented the lowest b^*
 209 value, hence the major blue component (2.8), followed by mashua (5.3) and tocosh (9.7). The
 210 differences observed between maca samples may be due either to the different treatments utilized
 211 for their preparation (ONWUDE *et al.*, 2017) or to cultivars with different chromatic characteristics.
 212 No information or comparisons for the color components are available in literature.
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214 3.2. Total polyphenol content

215 Supplementary Table 3 reports the results of TPC, performed on the EtOH:H₂O and
 216 MeOH:H₂O:acet extracts, as well as the results of the LSD test comparing the products. The great
 217 heterogeneity of the samples led to ANOVAs (not shown) always with significant differences. The
 218 average values, obtained by grouping the samples according to the species, are depicted in Fig. 2.



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220 **Figure 2.** Total polyphenol content (TPC) of the ethanol 80% (EtOH:H₂O) and
 221 methanol:H₂O:acetic acid (MeOH:H₂O:acet) extracts of powdered products from 11 species.
 222 Different letters indicate significant differences ($p \leq 0.05$) among species.
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224 EtOH:H₂O showed a lower TPC extraction capacity than MeOH:H₂O:acet, but the information
 225 provided was similar, as demonstrated by their very high linear coefficient of correlation ($r=0.98$).
 226 TPC was maximum for camu camu (38.3 and 76.4 g GAE/kg DM, respectively), followed by cat's
 227 claw 2 (24.2 and 53.6 g GAE/kg DM, respectively); the lowest TPCs were recorded in tocosh (2.5
 228 and 2.8 g GAE/kg DM), amaranth (on average, 3.1 and 2.8 g GAE/kg DM), lucuma (2.8 and 7.5 g
 229 GAE/kg DM) and six maca samples (on average, 3.6 and 6.7 g GAE/kg DM). The values generally
 230 fell within the range of variation reported in the literature for camu camu (FUJITA *et al.*, 2013),
 231 cat's claw (BERLOWSKI *et al.*, 2013; GALVEZ RANILLA *et al.*, 2010), amaranth (REPO-
 232 CARRASCO-VALENCIA *et al.*, 2010), lucuma (FUENTEALBA *et al.*, 2016), mesquite
 233 (CARDOZO *et al.*, 2010), maca (GALVEZ RANILLA *et al.*, 2010; CAMPOS *et al.*, 2013), mashua
 234 (CHIRINOS *et al.*, 2007; CHIRINOS *et al.*, 2013), yacon (CAMPOS *et al.*, 2013), but were lower
 235 than those described for graviola frozen pulp (ZIELINSKI *et al.*, 2014). For Peruvian purple maize
 236 the information available on TPC is reported in chlorogenic acid equivalent and is not directly
 237 comparable to our results, while for tocosh no similar information was found in literature.

238 **The Folin-Ciocalteu method sometimes overstates total phenolics content, because some reducing**
 239 **sugars (e.g. glucose and fructose) may interfere with the results; however, in this research the**
 240 **powders with the highest sugars content (yacon, mashua, lucuma and maca), generally have low**
 241 **TPC; conversely, the two highest TPC values were from camu camu and cat's claw, which showed**
 242 **very low sugars content.**

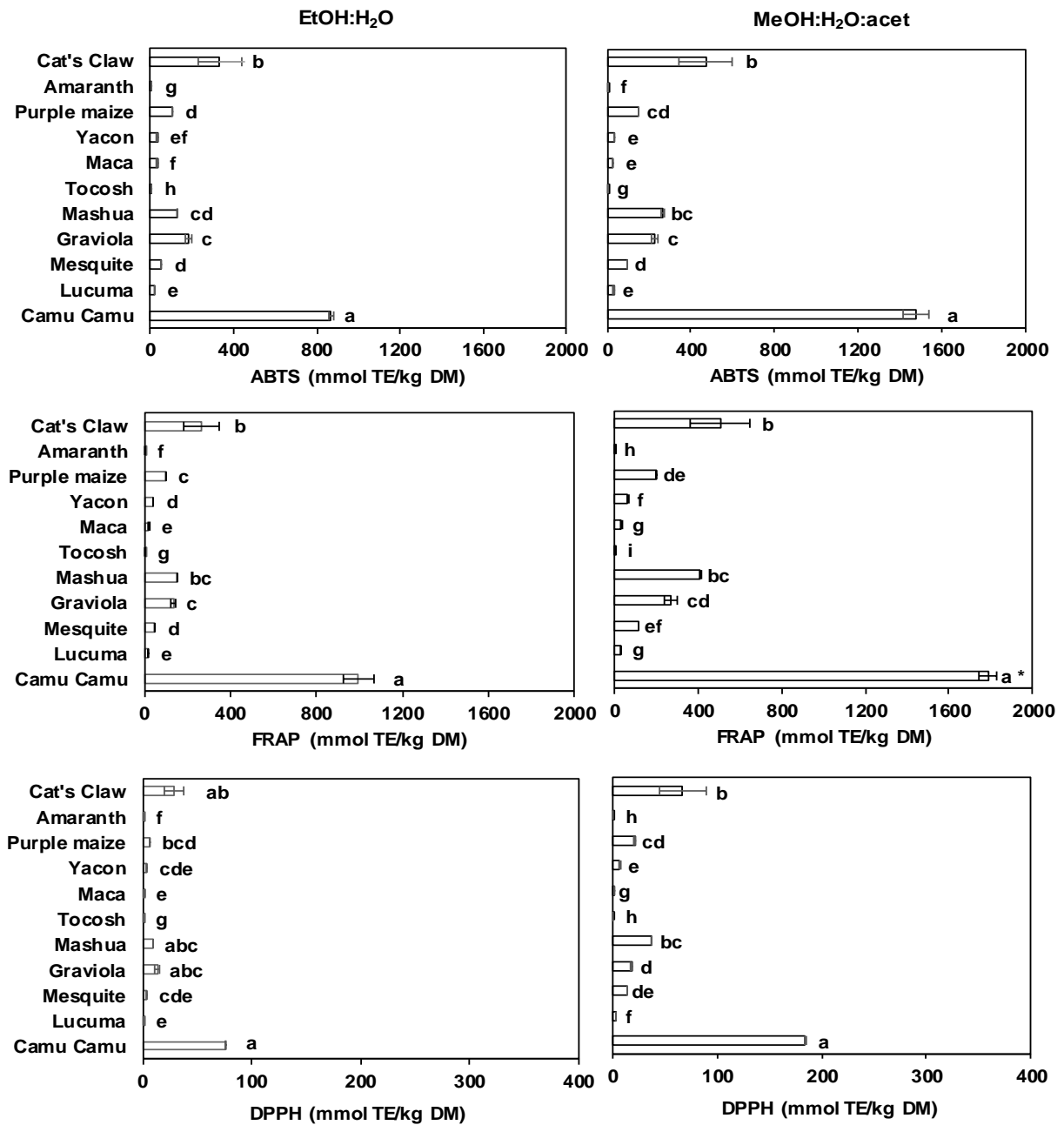
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245 3.3. Antioxidant capacity

246 The antioxidant capacity of the samples, assessed by the ABTS, FRAP and DPPH tests carried out
 247 on the EtOH:H₂O and MeOH:H₂O:acet extracts are shown in Supplementary Table 3, along with
 248 the results of the LSD test. The great heterogeneity among samples led to ANOVAs (not presented)

249 always indicating significant differences, as previously remarked for color and total polyphenols
 250 content. The average antioxidant capacities obtained by grouping the samples according to the type
 251 of product are presented in Fig. 3.



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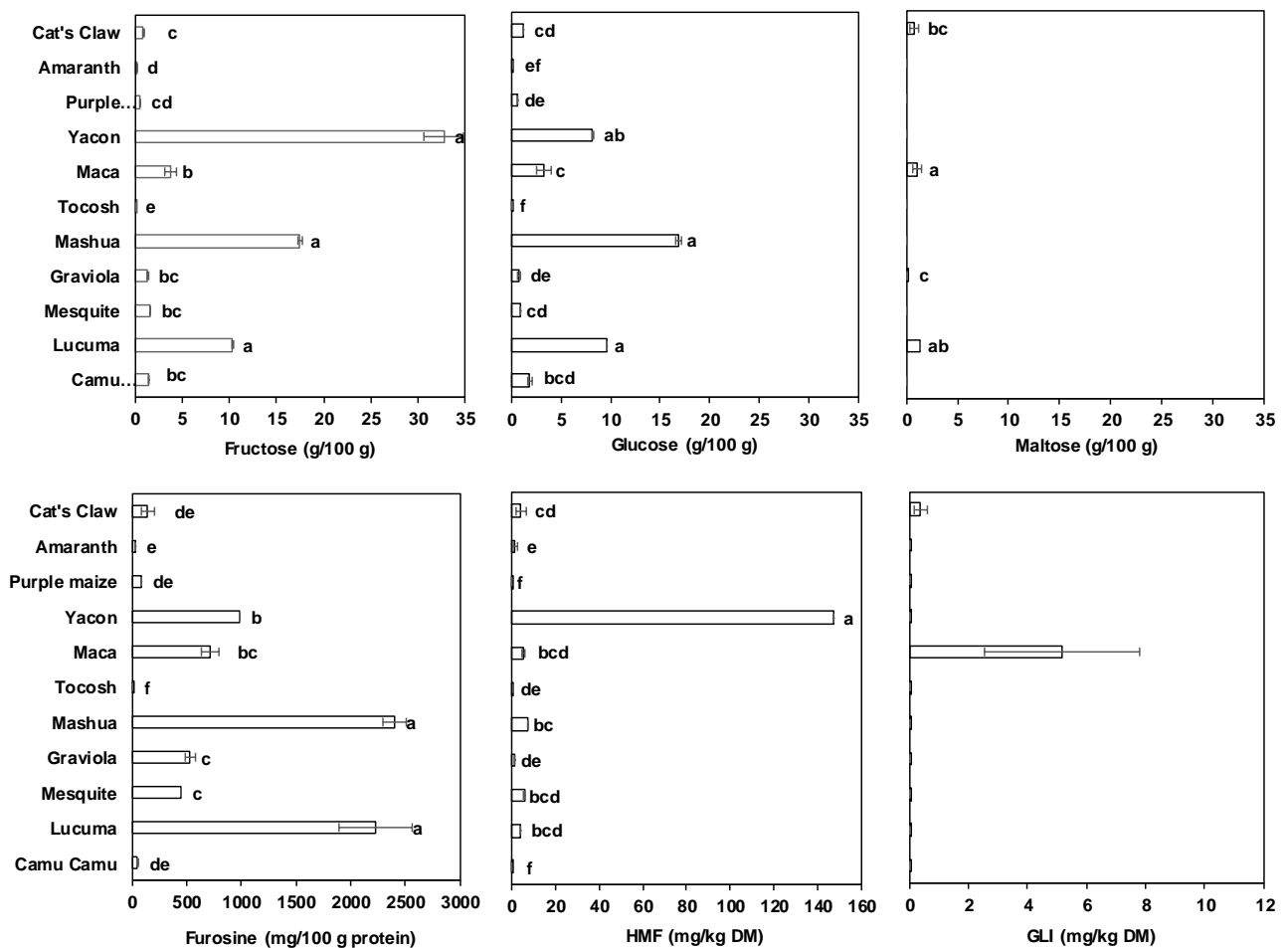
253 **Figure 3.** Antioxidant capacity (ABTS, FRAP and DPPH tests,)of the ethanol 80% (EtOH:H₂O)
 254 and methanol:H₂O:acetic acid (MeOH:H₂O:acet) extracts of powdered products from 11 species.
 255 Different letters indicate significant differences ($p \leq 0.05$) among species.
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257 The ABTS, FRAP and DPPH tests give similar and highly correlated results (r between 0.98 and
 258 1.00 for both EtOH:H₂O and MeOH:H₂O:acet extracts). A higher antioxidant capacity was
 259 observed in the MeOH extracts, the exceptions being amaranth (for all three methods), maca and
 260 tocosh (ABTS), graviola tea and maca 5 (DPPH). Camu camu, which had the highest TPC
 261 concentration (Fig. 2) but also an outstanding vitamin C content (FUJITA *et al.*, 2013), showed the
 262 highest antioxidant capacity (Fig. 3), followed by cat's claw, graviola, mashua, purple maize and

263 mesquite. On the other hand, tocosh, amaranth, yacon, maca and lucuma had low antioxidant
 264 activities, like that of wheat (YILMAZ *et al.*, 2015). Comparable results were reported for camu
 265 camu (DPPH: 153-185 $\mu\text{mol TE/g FW}$; CHIRINOS *et al.*, 2010), cat's claw (ABTS: 513 mmol
 266 TE/kg DM; FRAP: 507 mmol Fe/kg DM; BERLOWSKI *et al.*, 2013), graviola (ABTS: about 200
 267 mmol TE/kg DM; BERLOWSKI *et al.*, 2013), mashua (ABTS: 24.3-247.7 $\mu\text{mol TE/g DM}$; DPPH:
 268 23.2-157.1 $\mu\text{mol TE/g DM}$; CHIRINOS *et al.*, 2013), mesquite (ABTS: 57.0-61.6 $\mu\text{mol TE/g DM}$;
 269 CARDOZO *et al.*, 2010), yacon (ABTS 23-136 $\mu\text{mol TE/g DM}$; CAMPOS *et al.*, 2012), purple
 270 maize (DPPH: 23.1 $\mu\text{mol TE/g DM}$; CEVALLOS-CASALS and CISNEROS-CEVALLOS, 2003),
 271 lucuma (ABTS: 5.6-304.6 $\mu\text{mol TE/g DM}$; DPPH: 0.7-132.9 $\mu\text{mol TE/g DM}$; FUENTEALBA *et*
 272 *al.*, 2016) and amaranth (ABTS: 3.7 $\mu\text{mol TE/g DM}$; DPPH: 1.2 $\mu\text{mol TE/g DM}$; CHIRINOS *et al.*,
 273 2013). On the other hand, those of maca were slightly lower than the levels (ABTS: 67 $\mu\text{mol TE/g}$
 274 DM; FRAP: 11 mmol Fe/kg DM) observed by FUENTEALBA *et al.* (2016).
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276 3.4. Sugars content

277 The ANOVA (not presented) showed the existence of significant differences for sugars content
 278 among samples. The average values and the results of the LSD test for the different sugars are
 279 reported in Supplementary Table 2. The reducing sugars results obtained grouping the samples by
 280 species are presented in Fig. 4.



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282 **Figure 4.** Reducing sugars (fructose, glucose and maltose) and heat damage indices (furosine;
 283 hydroxymethylfurfural, HMF; glycosylisomaltol, GLI) of powder products from 11 species.
 284 Different letters indicate significant differences ($p \leq 0.05$) among species.

286 Fructose and glucose were detected in all samples, and were particularly abundant in yacon, mashua
287 and lucuma; maltose was detected only in cat's claw, most maca samples, lucuma and graviola.
288 Overall, yacon (40.9 g/100 g DM), mashua (34.4 g/100 g DM) and lucuma (21.2 g/100 g DM)
289 showed the highest content of reducing sugars which, on the other hand, were almost absent in
290 tocosh (0.11 g/100 g DM) and amaranth (0.30 g/100 g DM). Sucrose (a non-reducing sugar, but a
291 possible source of monosaccharides) was present in moderate quantities in mesquite, maca, mashua,
292 lucuma and yacon (42.5, 25.9, 16.4, 8.30, 8.20 g/100 g DM, respectively) and was very scarce in all
293 the other products. The presence of reducing sugars is important, because they are one of the basic
294 reactants involved in the formation of Amadori products during the Maillard reaction, when
295 exposed to high temperatures (e.g. oven drying, cooking, baking): therefore, higher reducing sugars
296 concentrations forebode higher heat damage during products manufacturing. Among the plants
297 tested, yacon is a well-known source of fructo-oligo-saccharides (CAMPOS *et al.*, 2012) and our
298 results are confirmed by the observations (49.2 g/100 g) of SCHER *et al.* (2009). The reducing
299 sugars content found in mashua is analogous to the quantity (6.4-45.3 g/100 g DM, average 28.4
300 g/100 g DM) reported by GUEVARA-FREIRE *et al.* (2018), while those of maca are slightly lower
301 than the value (13.10±0.17 g/100 g DM) described by RONDÁN-SANABRIA and FINARDI-
302 FILHO (2009), and that of mesquite is slightly inferior to the data (3.17-3.74 g/100 g DM) reported
303 by Cardozo *et al.* (2010) for different *Prosopis* spp. For fructose and glucose content our lucuma
304 results are within the broad range of variation (1.28-12.71 and 2.48-17.37 g/100 g DM) reported by
305 FUENTEALBA *et al.* (2016), and the amaranth ones are very similar to those (0.12 and 0.34 g/100
306 g DM) presented by GAMEL *et al.* (2006).

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308 3.5. Heat damage

309 The ANOVA (not presented) showed the existence of significant differences for heat damage
310 among the samples. The average values and the results of the LSD test for heat damage indices, i.e.
311 furosine, GLI and HMF, are reported in Supplementary Table 2. The results obtained grouping the
312 samples by species are reported in Fig. 4.

313 Non-enzymatic browning in dried products may be influenced by water activity, drying
314 temperature, pH and chemical composition of foods (Sagar Suresh Kumar, 2010). Furosine is an
315 index of the first steps of Maillard reaction, while GLI and HMF are markers of intermediate
316 phases; GLI is formed by the heating of maltose and aminoacids (especially glutamine), while HMF
317 is created not only by degradation of Amadori compounds but also of sugars.

318 Furosine content was very high in mashua and lucuma (>2000 mg/100 g protein), high in yacon and
319 most maca samples and low in camu camu, amaranth, purple maize as well as in two cat's claw
320 samples. Maca 6 and maca 7 had significantly lower furosine content than the other maca samples.
321 HMF was high only in yacon, but was detected, at lower levels, in several other samples, while GLI
322 was found only in maca 3, maca 6 and cat's claw 1. No Maillard reactions developed in tocosh (a
323 characteristic food, obtained by natural bacterial fermentation of straw-wrapped potatoes kept in
324 running water for several months) as furosine and GLI were lower than the detection limit and HMF
325 was very low, while camu camu, purple maize, amaranths, and cat's claw tea had limited heat
326 damage (low furosine levels and generally below-detection GLI and HMF). Furfural, an indicator of
327 more advanced Maillard reaction stages mainly produced by pentose degradation or thermal
328 degradation of HMF during caramelization, was absent in all samples, even if the method used for
329 GLI and HMF analysis is able to determine its presence.

330 Since water activity values for all the samples were very similar, ranging between 0.492 and 0.585,
331 and no correlation between heat damage indices and protein content (Table 1) exists, the
332 development of the Maillard reaction seems completely attributable to processing conditions and
333 reducing sugar concentration. The lofty heat damage of yacon (981.3 mg/100 g protein of furosine
334 and 146.6 mg/kg DM of HMF) is justified by its high fructose and glucose content and by the

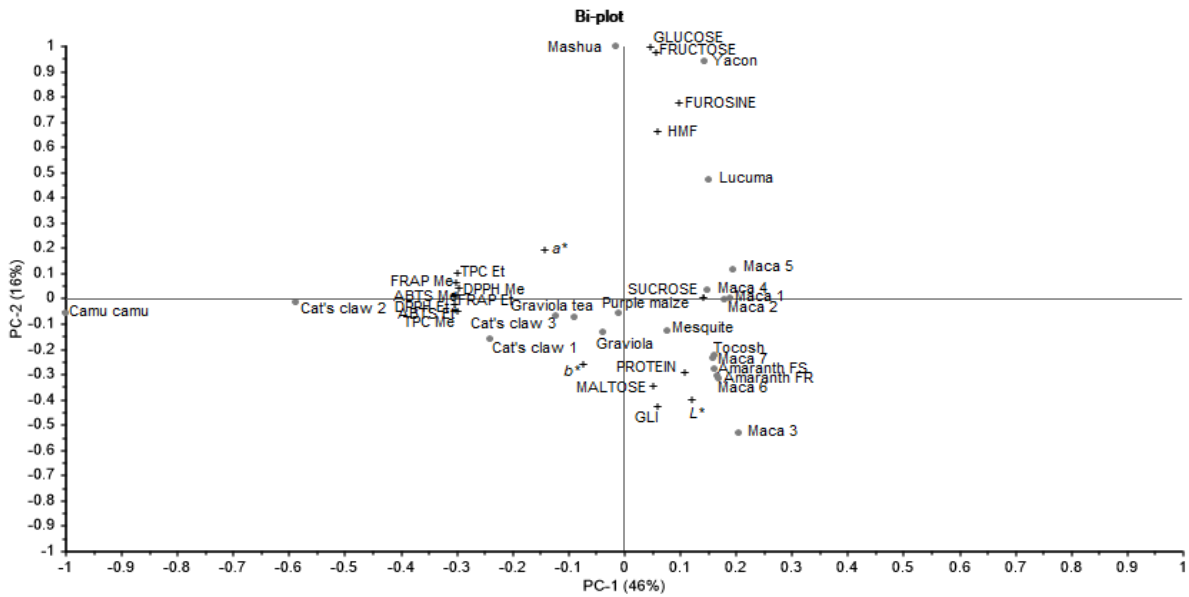
335 strong thermal treatment needed to inactivate its highly thermostable polyphenol oxidase enzyme
336 (NEVES DA SILVA, 2007) for a luminous color (Figure 1). These conditions lead to the
337 degradation of the Amadori compounds and to the formation of intermediate compounds (i.e.
338 HMF). The furosine levels of the other samples correlate well to the concentration of reducing
339 sugars ($r = 0.93$). Thus, mashua, lucuma and most of maca powders with relevant content of
340 reducing sugars have high furosine levels while samples with low reducing sugars content present
341 limited furosine. However, furosine alone is not suitable to completely describe the heat damage of
342 all powder products. The presence of HMF (0.9-11.8 mg/kg DM) in most samples points to an
343 intermediate development of Maillard reaction. The samples with detectable GLI (maca 3, maca 6
344 and cat's claw 1) showed the highest maltose levels but relatively low reducing sugar
345 concentrations (4.0-7.7 g/100 g DM). The simultaneous formation of HMF suggests higher-than-
346 the-average processing temperatures for these samples. A similar hypothesis can be made for some
347 samples with very low reducing sugar concentrations (0.2-2.5 g/100 g DM) and significant HMF
348 content (1.3-6.0 mg/kg DM).

349 To the best of our knowledge, no information on heat damage in this type of powder products is
350 available. With reference to other vegetables, furosine contents of 14-262 mg/100 g protein
351 (RUFÍAN-HENARES *et al.*, 2013) and of 457-1172 mg/100 g protein (BIGNARDI *et al.*, 2016) are
352 reported in sweet pepper and in dried red chili pepper, respectively; similarly, RÍOS-RÍOS *et al.*
353 (2018) describe furosine concentrations of 46.6-110.1 mg/100 g protein in black garlic powder,
354 while HMF levels of 1.3-9.5 mg/kg DM are recorded by SORIA *et al.* (2009) in carrots dried under
355 different conditions.

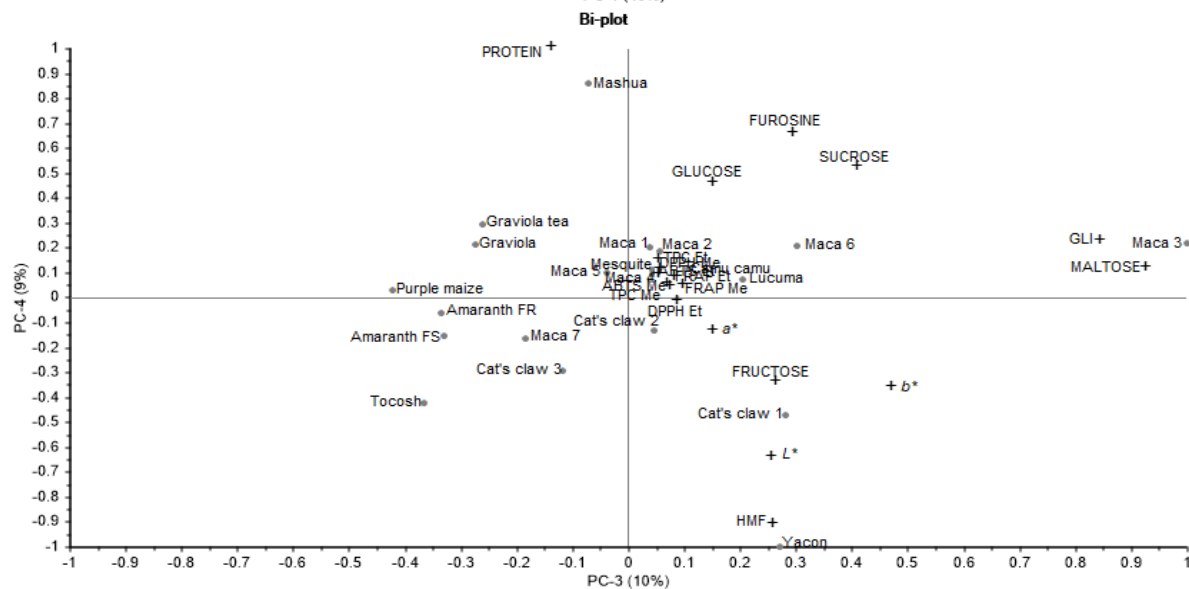
356

357 **3.6. Principal components analysis**

358 Fig. 5 depicts the biplots of scores and loadings obtained by the principal components analysis
359 (PCA) performed considering all samples and parameters. PC1 and PC2 describes 46% and 16% of
360 variation (Fig. 5A), while PC3 and PC4 10% and 9% (Fig. 5B); therefore, the initial four PC
361 explain 81% of total variation. The PCA unmistakably separates the different species. PC1,
362 characterized by antioxidant properties, differentiates camu camu and cat's claw along the left side,
363 while PC2, mainly related to heat damage, positions mashua, yacon and lucuma in the upper side
364 (high furosine, HMF, glucose and fructose contents), and maca 3, maca 6, amaranth and tocosh in
365 the bottom side (high L^* , GLI, maltose and protein). PC3 further splits purple maize, graviola,
366 amaranth and tocosh, from mesquite and maca samples, while PC4 divides mashua (top of the plot,
367 characterized by high protein and furosine contents) from yacon (bottom, high HMF and L^*); maca
368 3 sits alone in a spot defined by high GLI content.



369



370

371 **Figure 5.** Bi-plot of scores and loads for the first four principal components (PC1 vs. PC2, A; PC3
 372 vs. PC4, B) of the principal component analysis carried out on colour coordinates (L^* , a^* , b^*),
 373 protein content, total polyphenol content (TPC) and antioxidant capacity (ABTS, FRAP and DPPH
 374 tests) of the ethanol:H₂O (Et) and methanol:H₂O:acetic acid (Me) extracts, reducing sugars,
 375 furosine, hydroxymethylfurfural (HMF), and glycosylisomaltol (GLI) of 21 powder products.
 376

377 **4. CONCLUSIONS**

378 Our results show that, for the traits analysed, camu camu powder and cat's claw are excellent
 379 products, because they have high levels of total polyphenols and antioxidant capacity together with
 380 low heat damage. Other interesting products are the powders of graviola, purple maize and
 381 mesquite, while the high antioxidant properties of mashua are coupled to severe heat damage.
 382 For an effective use in the food industry, it is necessary to evaluate the stability of the antioxidant
 383 capacity of the powders during the manufacturing process and the digestion of innovative high-
 384 nutritional-value foods, as well as to assess the sensorial quality of the end products.
 385

386 **ACKNOWLEDGEMENTS**

387 We thank Giulia Malizia and Mattia Magistrelli for their assistance in the lab analyses.

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