Bioactivity of food melanoidins is mediated by gut microbiota

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21 ABSTRACT

22 Melanoidins are an important component of the human diet (average consumption 10g/day), which escape gastrointestinal digestion and are fermented by the gut microbiota. 23 In this paper melanoidins from different food sources (coffee, bread, beer, balsamic 24 vinegar, sweet wine, biscuit, chocolate, and breakfast cereals) were submitted to an in vitro 25 26 digestion and fermentation process, and their bioactivity was assessed. Some melanoidins were extensively used by gut microbes, increasing production of short chain fatty acids 27 (mainly acetate and lactate) and favoring growth of the beneficial genera Bifidobacterium 28 (bread crust, pilsner and black beers, chocolate and sweet wine melanoidins) and 29 Faecalibacterium (biscuit melanoidins). Quantification of individual phenolic compounds 30 after in vitro fermentation allowed their identification as microbial metabolites or phenolics 31 melanoidins backbone released from the (specially pyrogallol, 2-(3,4-32 dihydroxyphenyl)acetic and 3-(3,4-dihydroxyphenyl)propionic acids). Our results also 33 34 showed that antioxidant capacity of melanoidins is affected by gut microbiota fermentation.

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36 KEYWORDS: melanoidins, gut microbiota, short chain fatty acids, polyphenols,
37 antioxidant capacity.

38 **1. Introduction**

Melanoidins are end products of Maillard reaction (MR), which occurs among the 39 amino group of an amino acid, protein, or vitamin and the carbonyl group of a reductive 40 sugar or oxidized lipid (Rufián-Henares & Pastoriza, 2016). The MR is very common in 41 foods since it happens during heating but also during storage at room temperature (Rufián-42 Henares, Guerra-Hernández & García-Villanova, 2006; Tagliazucchi & Verzelloni, 2014). 43 Melanoidins are therefore high molecular weight products responsible for the brown color 44 of thermally treated foods like bread, coffee, cocoa, etc. The detailed structure of 45 melanoidins remains unknown due to the varied nature of the foods in which melanoidins 46 are formed. Thus, in biscuits and bread, melanoidins are mainly composed of 47 48 polysaccharides and proteins (melanoproteins), while in coffee and cocoa melanoidins are usually smaller and incorporate phenolic compounds such as chlorogenic acids or catechins 49 (Morales, Somoza, & Fogliano, 2012). 50

Melanoidins make an important contribution to our diet: on average, around 10 g of 51 52 melanoidins per day could be ingested (Fogliano & Morales, 2011; Pastoriza and Rufián-Henares, 2014). Therefore, knowing the possible benefits or risks of melanoidins to human 53 health is important. Accordingly, many different biological effects have been attributed to 54 melanoidins. They can behave as prebiotic (Helou et al., 2015; Jiménez-Zamora, Pastoriza 55 & Rufián-Henares, 2015), they have an important antioxidant capacity (Delgado-Andrade 56 & Morales, 2005; Carvalho, Correia, Lopes, & Guido, 2014; Pastoriza& Rufián-Henares, 57 2014; de la Cueva, Seiguer, Mesías, Rufián-Henares, & Delgado-Andrade, 2017) and 58 antimicrobial activity (Rufián-Henares & de la Cueva, 2009), and they can possess 59

60 inhibitory activity against angiotensin I-converting enzyme (Rufián-Henares & Morales,
61 2007).

In the gut, dietary melanoidins escape gastrointestinal digestion (similarly to fiber) 62 and reach the colon where they become substrates for the gut microbiota (Wang, Qian, & 63 Yao, 2011). However, the potential bioactivity of melanoidins after being exposed to the 64 gut microbiota has not been extensively studied. It has been proposed that the colon is the 65 main region of the gut where melanoidins exert their effects since they escape digestion 66 (Rufián-Henares & Pastoriza, 2015). However, fermentation of melanoidins by gut 67 microbes and resulting production of short chain fatty acids (SCFAs) have not been yet 68 deeply studied. SCFAs are the main microbial metabolites and have been attributed to 69 70 several health effects. For example, decreasing the luminal pH is one of the most obvious effects of SCFAs release in the gut, which can hinder the growth of pathogenic bacteria. 71 Butyrate is used as a substrate by the epithelial cells, and the three main SCFAs (acetate, 72 73 propionate and butyrate) are important for the maintenance of the gut barrier (Ríos-Covián et al., 2016). In addition, all these SCFAs have a protective role in diet-induced obesity 74 (Lin et al., 2012) since butyrate and propionate have been related to the production of gut 75 hormones and therefore reduction of food intake (Ríos-Covián et al., 2016). SCFAs (mostly 76 butyrate) could also have an important role in colorectal cancer protection via reducing 77 78 inflammation and increasing cell apoptosis (Donohoe et al., 2014).

Taking all this information into account, the objective of this research was to investigate the fate of melanoidins during simulated gastrointestinal digestion and subsequent fermentation by human gut microbiota. We assessed how melanoidins shaped gut microbial community structure and its functionality (SCFAs production), and we

measured the release of phenolic compounds and antioxidants as consequence of microbial
metabolism.

- 85
- 86 **2.** Materials and methods
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2.1. Reagents.

For the antioxidant assays, 2,4,6-Tri(2-pyridyl)-s-triazine (TPTZ), potassium
persulfate, iron (III) chloride hexahydrate, 2,2-Diphenyl-1-picrylhydrazyl, 2,2'-Azinobis(3-ethylbenzothiazoline-6-sulfonic acid) diammonium salt, trolox ((±)-6-Hydroxy2,5,7,8-tetramethylchromane-2-carboxylic acid), and methanol were purchased from
Sigma-Aldrich (Germany). Short chain fatty acids and phenolic standards were from
Sigma-Aldrich (Germany).

For the *in vitro* digestion and fermentation, the following reagents were used: 94 potassium di-hydrogen phosphate, potassium chloride, magnesium chloride hexahydrate, 95 sodium chloride, calcium chloride dihydrate, sodium mono-hydrogen carbonate, 96 97 ammonium carbonate, hydrochloric acid, all obtained from Sigma-Aldrich (Germany). The enzymes - salivary alpha-amylase, pepsin from porcine, and bile acids (bile extract 98 porcine) - were purchased from Sigma-Aldrich, and porcine pancreatin was fromAlfa 99 Aesar (United Kingdom). The fermentation reagents (sodium di-hydrogen phosphate, 100 sodium sulfide, tryptone, cysteine, and resazurin) were obtained from Sigma-Aldrich 101 (Germany). 102

For individual phenolic quantification the following standards were used: hydroxymethyl-furan-carbaldehyde, 3,4-Dihydroxybenzoic acid, 3,5-Dihydroxybenzoic acid, 3,4-Dihydroxyphenilacetic acid, 3,4-Dihydroxybenzaldehyde, p-Hydroxyphenilacetic

vanillina, acid, cafeic acid. chlorogenic acid, 4-O-Cafeoylquinic 106 acid, 3.4-Hydroxyphenilpropionic acid, trans-hydroxycinnamic acid, p-Cumaric acid, epicatechin, 107 epicatechin gallate, ferulic acid, 3,4-Dimethoxybenzaldehide, o-Cumaric acid, pyrogallol, 108 resveratrol, daidzein, glicitein, genistein and formonotein, were purchased from Sigma-109 Aldrich (Germany). Moreover, diethyl ether for phenolic compounds extraction was 110 purchased from Sigma-Aldrich (Germany). 111

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2.2. Melanoidins preparation

Melanoidins were isolated from ground coffee "Cafés Cumbal", pilsner beer 114 115 "Alhambra" and Irish black beer "Guinness", corn breakfast cereals "Corn Flakes", bread crust separated from French bread by grinding, Marie biscuits, 85% black chocolate 116 "Lindt", balsamic vinegar "Borges", and sweet wine "Málaga Vírgen". All products were 117 purchased in Carrefour in Granada (Spain). Spent coffee grounds were obtained after 118 brewing the same coffee used for melanoidins extraction in an espresso coffee machine. In 119 the case of melanoproteins (biscuit and bread crust) they were obtained after in vitro 120 digestion with Pronase E and subsequent diafiltration (Pastoriza, Roncero-Ramos, Rufián-121 Henares, & Delgado-Andrade, 2014). In vitro digestion of bread crust and biscuits were 122 carried out in sodium tetraborate buffer 0.1M and pH 8.2 at a ratio of 150 g of food/L of 123 buffer during 72h at 37°C. Pronase E concentration was 100 mg/L of buffer. The digested 124 solution was submitted to diafiltration for melanoidin isolation using a 5 kDa membrane 125 from Sartorius (United Kingdom) and a peristaltic pump from Cole Parmer (USA). 126 Melanosaccharides (obtained from coffee, beers, chocolate, balsamic vinegar and sweet 127 wine) were directly isolated by diafiltration using the same membrane and pump as before 128

as described in Rufián-Henares & de la Cueva, (2009). In the case of chocolate, it was first
dissolved in Milli-Q water. Melanoidins were stored at -80 °C until analysis.

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2.3. Spent coffee grounds preparation

Spent coffee grounds were included in the experiments since they still retain melanoidins from coffee and other bioactive compounds (Jiménez-Zamora et al., 2015). Spent coffee grounds were obtained after brewing coffee in an expresso coffee maker as described in Pérez-Burillo et al. (2019). Spent coffee grounds were afterwards freeze-dried and stored at -80 °C until further analysis.

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139 2.4. In vitro gastrointestinal digestion and fermentation

All samples were subjected to an *in vitro* digestion and fermentation to mimic 140 physiological processes in the human gut. The *in vitro* digestion method was carried out 141 with alpha-amylase, pepsin, pancreatin and bile salts according to the protocol described by 142 Pérez-Burillo, Rufián-Henares & Pastoriza (2018). The in vitro fermentation was carried 143 out according to the protocol described by Pérez-Burillo et al. (2018). Briefly, the solid 144 residue obtained after in vitro digestion plus a 10% of the digestion supernatant was 145 146 fermented by gut microbiota from faecal samples obtained from three healthy adult donors (mean age range 27.3 years, not taking antibiotics, mean Body Mass Index = 21.3). 147

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2.5. High-throughput amplicon sequencing

High-throughput sequencing of microbial 16S rRNA genes was carried out as
previously described (Pérez-Burillo et al., 2019). Bacterial genomic DNA was isolated
from each sample using ZR Fungal/Bacterial DNA MiniPrep kit (Zymo Research). DNA

was amplified using two pairs of primers, one targeting 16S rDNA V1-V2 region [forward 153 primer 16S gene complementary sequence AGRGTTYGATYMTGGCTCAG and reverse 154 primer 16S gene complementary sequence GCWGCCWCCCGTAGGWGT], and another 155 156 targeting V4 region [forward] GCCAGCMGCCGCGG and reverse GGACTACHVGGGTWTCTAAT complementary sequences, respectively]. 157 PCR amplification was performed with 25ng of starting DNA material and included 10 cycles of 158 159 linear elongation with only the forward primers used, followed by 25 cycles of traditional exponential PCR (Paliy & Foy, 2011). High throughput sequencing was performed on Ion 160 161 Torrent Personal Genome Machine. We obtained an average of 17,490 sequence reads per 162 sample. Sequence reads were processed in QIIME (Caporaso et al., 2010). Sequence read counts for each OTU were adjusted by dividing them by known or predicted number of 16S 163 rRNA gene copies in that organism's genome following a previously described approach 164 (Rigsbee, Agans, Foy, & Paliy, 2011), and the resulting cell counts were sub-sampled to the 165 lowest value among samples. The cell counts obtained independently for each sample based 166 on the sequencing of V1-V2 and V4 16S rRNA gene regions were merged together into a 167 single taxon abundance estimate via $A_{CUM} = \sqrt{(A_{V1V2}^2 + A_{V4}^2)/2}$ calculation, where A is an 168 abundance value for each taxon. 169

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2.6. *Analysis of short chain fatty acids*

SCFAs determination was carried out by UV-HPLC according to the procedure
 described in Panzella et al. (2017). After the fermentation process, 1 mL of melanoidin
 fermentation supernatant was centrifuged to remove solid particles, filtered through a 0.22
 µm nylon filter, and finally transferred to a vial for HPLC analysis.

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2.7. Analysis of phenolic compounds

Phenolic compounds were analyzed through UV-HPLC as described in Moreno-178 Montoro et al. (2015). In brief, one mL of melanoidin fermentation supernatant was mixed 179 with 1 mL of diethyl ether and kept in the dark at 4°C for 24 hours. The organic phase was 180 then collected and another two extractions with diethyl ether were performed. These 3 mL 181 182 of diethyl ether were dried in a rotary evaporator set at 30°C and the solid residue was resuspended in 1 mL of methanol:water (50:50 v/v) mix. The mixture was then ready to be 183 injected into HPLC system. Identification and quantification were carried out by comparing 184 185 retention times obtained from pure standards (listed in *reagents* section).

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2.8. Antioxidant assays

188 The antioxidant capacity of melanoidins before and after *in vitro* digestion-189 fermentation was measured using three different approaches.

190 $TEAC_{ABTS}$ assay. The radical scavenging activity of samples was performed 191 following the method described by Re et al. (1999) adapted to a microplate reader 192 (Jiménez-Zamora, Delgado-Andrade & Rufián-Henares, 2016). The results were expressed 193 as µmol Trolox equivalents per g of sample.

194 $TEAC_{FRAP}$ assay. The reducing capacity of iron was analyzed by the method 195 described by Benzie & Strain (1996) adapted to a microplate reader (Rufián-Henares, 196 García-Villanova & Guerra-Hernández, 2006). The results obtained were expressed as 197 µmol Trolox equivalents per g of sample.

198 $TEAC_{DPPH}$ assay. This method was carried out according to the procedure described 199 by Yen & Chen (1995). Results were expressed as µmol equivalents of Trolox per g of 200 sample.

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2.9. Multivariate statistical analyses

Unconstrained principal coordinates analysis (PCoA) utilizing phylogenetic weighted UniFrac distance as a measure of sample dissimilarity was performed on the genus-level microbial abundance dataset. Principal component analysis was carried out to assess sample similarity based on the levels of short chain fatty acids and phenolic compounds. Multivariate analyses were run in R and Matlab (Paliy & Shankar, 2016).

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3. Results and discussion

Melanoidins are bioactive compounds generated during the thermal processing of 210 foods (Rufián-Henares, Delgado-Andrade & Morales, 2006) with bioactive effects 211 212 potentially similar to those of fiber-enriched foods (Delgado-Andrade, Rufián-Henares & Morales, 2007). Thus, in order to assess the bioactivity of food melanoidins and the effect 213 of the gut microbiota over such bioactivity, ten melanoidins were isolated from different 214 foods and subjected to an *in vitro* digestion-fermentation process designed to mimic natural 215 digestion in the human oral, gastric, and intestinal chambers. Melanoidins bioactivity was 216 measured as their ability to (i) modify the gut microbial community (gut microbiota 217 218 composition) and activity (production of SCFAs), (ii) release phenolic compounds, and (iii) 219 provide antioxidant capacity.

Using high-throughput sequencing of 16S rRNA gene, microbiota community

Fermentation of melanoidins from different foods by human fecal 221 3.1. 222 *microbiota promotes different community structures*

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structure was determined in all fermented samples. As depicted in Figure 1A, samples 224 225 could be separated into 4 distinct groups based on microbiota composition. Digested melanoidins from bread crust, cereals, and pilsner and black beers promoted a remarkable 226 expansion of members of genus Bifidobacterium, which represented on average 92% of all 227 microbial cells in these samples. Microbiota that was fed with digested chocolate and sweet 228 wine melanoidins also contained significant amounts of bifidobacteria, albeit at a lower 229 230 level (68% on average). In contrast, biscuit melanoidins promoted the expansion of Faecalibacterium (30% relative abundance in this community vs 5% average abundance in 231 all other samples). The remaining three melanoidin samples maintained microbial 232 communities more similar to that of the baseline inocula (represented by the blank sample 233 that only received inocula and buffer, but no additional nutrient sources). This baseline 234 community was relatively abundant in members of genera Blautia, Roseburia, Citrobacter 235 and Enterobacter. 236

The separation of communities into 4 distinct groups was confirmed by the 237 phylogenetic PCoA ordination analysis (Figure 1B). Abundances of Bifidobacterium and 238 Faecalibacterium were the two main drivers of sample distribution in the PCoA space (see 239 Figures 1C and 1D). Based on the community structure analysis, it appears that 240 241 melanoidins from spent coffee grounds were largely not fermented by human fecal microbiota, because the community structure was very similar to that maintained in the 242 buffer medium. Abundances of Bifidobacterium and Faecalibacterium displayed a 243 reciprocal relationship (Spearman rank correlation R_s= -0.97), likely indicating different 244

enzymatic capacities of these bacteria to break down different melanoidin structures. 245 Bifidobacterium expansion resulted in a remarkable reduction in community diversity 246 (Shannon H' index of 0.5 for group 1 samples vs 2.3 for the baseline group communities), 247 possibly indicating that only members of this genus were able to utilize melanoidins from 248 bread crust, cereals, and pilsner and black beers for growth. Note that both Bifidobacterium 249 and Faecalibacterium are considered beneficial to human health (Murri et al., 2013; Flint, 250 251 Duncan, Scott, & Louis, 2015), thus their expansion can be viewed as a positive effect and these melanoidins could be used to formulate functional foods. 252

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254 3.2. SCFAs production from melanoidins fermentation

Short chain fatty acids are the main end products of microbial fermentation in the 255 gut. Thus, the concentrations of four major SCFAs (acetate, butyrate, lactate, and 256 propionate) were measured in all fermented samples (Flint et al., 2015). Overall, acetate 257 and lactate were the main end products of fermentation of melanoidins, with levels of 258 butyrate and propionate being significantly lower (Figure 2A). Acetate and lactate are the 259 main end products of Bifidobacterium fermentation (Flint et al., 2015), and their levels 260 correlated with this genus abundance (Spearman rank correlation $R_s = 0.75$ and 0.78, 261 respectively). Thus, the SCFAs measurements fit well with the gut microbiota analysis 262 described above. As anticipated, blank sample had very low levels of all four SCFAs, and 263 other samples in group 4 also generated lower SCFAs levels, indicating poor melanoidin 264 fermentation. In contrast, melanoidins from bread crust, cereals, pilsner and black beers 265 (group 1) were highly fermented and produced the largest SCFAs amounts (see Figure 2A). 266 Since melanoidins from these foods promoted *Bifidobacterium*, they can be considered 267 possible prebiotics for improvement of gut health. 268

The output of the principal component analysis of SCFAs abundance dataset also
corroborated the separation of samples into four groups (Figure 3A). Samples from group 4
where SCFAs production was low were well split from other samples, which were more
tightly clustered. Co-inertia analysis (Shankar et al., 2017) was used to assess if sample

(ure 3A). Samples from group 4 270 her samples, which were more 271 272) was used to assess if sample distribution was similar between SCFAs-based PCA and genus-based PCoA ordinations. 273 As shown in Figure 3C, samples were indeed distributed similarly in both ordination spaces 274 (p = 0.03), indicating that microbiota and short chain fatty acids measurements 275 corroborated each other. 276

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3.3. 278 Release of phenolic compounds during melanoidins fermentation

Melanoidins can contain different phenolics depending on the food source 279 (Tagliazucchi & Bellesia, 2015). Therefore, the metabolization of melanoidins by gut 280 microbes could result in a release of phenolic compounds, which could elicit anti-oxidant 281 properties. Moreover, these phenolics could be further metabolized by the microbes, 282 usually yielding smaller compounds (Selma, Espín, & Tomás-Barberán, 2009). UV-HPLC 283 was used to measure the concentrations of 26 phenolic compounds in all profiled samples 284 (Figure 2B and Table 1). In contrast to the SCFAs production, PCA analysis of phenolic 285 286 compounds dataset failed to distribute samples according to the four groups defined above. Instead, most samples clustered together, with an exception of chocolate and coffee 287 samples that were significant outliers (Figure 3B). Most samples differed significantly in 288 their metabolite profiles, and the detected compounds in each sample are described below. 289

Bakery products melanoidins: Biscuit, bread crust, and breakfast cereal melanoidins 290 (melanoproteins) released similar amounts of polyphenols during fermentation (2.08-2.44 291 $\mu g/g$). Most of them (60-82%) have been identified as microbial metabolites from other 292

phenolic compounds (Rowland et al., 2018). These melanoidins showed high amounts of 293 pyrogallol (Table 1), which could indicate the presence of epicatechin or gallic acid in the 294 melanoidin backbone (Rowland et al., 2018). Biscuit melanoidins also released noticeable 295 amounts of vanillin and daidzein, probably coming from the food source. Bread crust 296 melanoidins showed high amounts of 2-(3,4-dihydroxyphenyl)acetic acid and 3-(3,4-297 dihydroxyphenyl)propionic acid (Table 1), microbial metabolites of quercetin, myricetin, 298 epicatechin, or hydroxycinnamic acids (Selma et al., 2009; Marín, Miguélez, Villar, 299 &Lombó, 2015; Rowland et al., 2018). Breakfast cereals melanoidins showed high 300 amounts of p-hydroxyphenylacetic acid (Table 1), a potential microbial metabolite of 301 302 hydroxycinnamic acids (Rowland et al., 2018). Relatively high amounts of ferulic acid were also found, probably from the food sources. 303

Beer melanoidins: Black and pilsner beer melanoidins released similar amounts of 304 polyphenols during in vitro fermentation, 7.38 and 7.42 µg/g respectively. However, the 305 306 percentage of possible microbial metabolites was higher (52%) in black beer than in pilsner (35%). Both contained notable amounts of the microbial metabolites 2-(3,4-307 dihydroxyphenyl)acetic acid, 3-(3,4-dihydroxyphenyl)propionic acid and pyrogallol. Both 308 309 of them also showed high amounts of 3,5-dihydroxybenzoic acid, which could come from the beer itself (Wannenmacher, Gastl & Becker, 2018), since it has not been related to 310 311 microbial metabolism. Black beer was characterized by high amounts of 4-O-312 caffeoylquinic acid and epigallocatechin gallate and the microbial metabolite 3,4-313 dihidroxybenzoic acid, metabolite from hydroxycinnamic acids and cyanidin (Rowland et al., 2018) (Table 1). Pilsner beer, on the other hand, was characterized by high amounts of 314

ferulic acid and the microbial metabolite 3,4-dihydroxybenzaldehyde, which, according to
(Selma et al., 2009), results from microbial degradation of quercetin (Table 1).

Balsamic vinegar melanoidins: Released the lowest amount of polyphenols (1.09
 μg/g), almost exclusively 3,5-dihydroxybenzoic acid and the microbial metabolite 2-(3,4 Dihydroxyphenyl)acetic acid.

Sweet wine melanoidins: The amount of polyphenols released from these 320 melanoidins was similar to that of bakery products, 1.93 µg/g. Most of them (74%) were 321 possible microbial metabolites: 2-(3,4-dihydroxyphenyl)acetic acid, 322 3,4dihydroxybenzaldehyde, p-hydroxyphenylacetic acid, 3-(3,4-dihydroxyphenyl)propionic 323 acid, and pyrogallol. Regarding non-microbial metabolites, the higher amounts were 324 showed by orto- and meta-coumaric acids. 325

Chocolate, and coffee melanoidins, and spent coffee grounds: Released the highest 326 amounts of polyphenols, 60.15, 48.76, and 58.45 µg/g, respectively. Phenolic profile of 327 chocolate melanoidins and spent coffee grounds were similar, with most phenols being 328 non-microbial metabolites. Especially large amounts of 3,5-dihydroxybenzoic acid were 329 found in both samples. Chocolate melanoidins showed also noticeable amounts of the 330 331 microbial metabolites 3,4-dihidroxybenzoic acid, 2-(3,4-dihydroxyphenyl)acetic acid, 3-(3,4-dihydroxyphenyl)propionic acid, and pyrogallol. Spent coffee grounds were largely 332 not metabolized, since 90% of the phenolics were represented by 3,5-dihydroxybenzoic 333 acid, which is not described as microbial metabolite. On the other hand, 51% of coffee 334 melanoidins phenolics were represented by microbial metabolites, especially p-335 hydroxyphenylacetic acid (probably coming from hydroxycinnamates metabolism), and 336

pyrogallol (from epicatechin or gallic acid metabolism) (Table 1). A large concentration of
chlorogenic acid was also released from coffee melanoidins.

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3.4. Melanoidins possess significant antioxidant capacity

According to Tagliazucchi & Verzelloni (2014) there is a strong correlation between 341 the phenolic content of melanoidins and their antioxidant capacity. Thus, three different 342 343 methods were used to estimate the antioxidant capacity of fermented and digested samples: ABTS (measures most antioxidants including phenolics and thiols), FRAP (measures Fe³⁺ 344 reduction) and DPPH (measures antiradical activity). A strong correlation among the three 345 antioxidant methods was found ($R_s \ge 0.95$). All three measurements also showed a 346 moderate correlation with the total amounts of phenolics detected in each sample ($0.35 \le R_s$) 347 \leq 0.44), though the relationships were not statistically significant (p \geq 0.2). In all three 348 assays, coffee and spent coffee melanoidins showed the highest antioxidant capacity after 349 fermentation, followed by sweet wine melanoidins and balsamic vinegar melanoidins 350 (Figure 2C). On the other hand, pilsner beer, breakfast cereals, and bread crust melanoidins 351 were always those with the lowest values. Our results are in agreement with Verzelloni, 352 Tagliazucchi & Conte (2010) who reported that coffee melanoidins had more antioxidants 353 than black beer and balsamic vinegar melanoidins. Even though the structure of 354 melanoidins varies significantly, coffee melanoidins as well as sweet wine and chocolate 355 melanoidins contain phenolic moieties that could make them more antioxidant than others 356 (Morales et al., 2012). 357

Measurements of antioxidant capacity of digested but not fermented melanoidins similarly revealed that coffee melanoidins had the highest capacity, whereas the bread crust melanoidins had the lowest (**Table 2**). For biscuit, breakfast cereals, and balsamic vinegar

melanoidins the antioxidant capacity increased after fermentation; for other melanoidins it 361 was opposite. According to Wang et al. (2011) melanoidins antioxidant capacity is thought 362 to be partly related to their metal chelating ability because of their anionic nature and partly 363 due to their scavenging activity. After fermentation, melanoidins are degraded to some 364 extent by the gut microbiota, thus decreasing their chelating capacity. Moreover, 365 scavenging capacity is directly related to the presence of phenolic moieties in melanodin 366 367 structure (Wang et al., 2011), and the biotransformation of some of them by gut microbes could result in a decreased antioxidant capacity. Accordingly, a statistically significant 368 positive correlation was found between the ratio of antioxidant capacities prior to/after 369 fermentation and the total of identified phenolic compounds ($R_s = 0.70$, p < 0.05). These 370 data potentially indicate that transformation of melanoidins by gut microbiota can reduce 371 their antioxidant capacity. This was true for black and pilsner beer, chocolate, coffee and 372 sweet wine melanoidins (Table 2). This hypothesis was supported by a negative correlation 373 found between antioxidant capacity of the fermented samples and the total SCFAs 374 produced in those samples ($R_s = -0.47$, p < 0.05). 375

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4. Conclusions

Melanoidins are an important part of our diet, reaching average amounts of 10 g/day. Here we have demonstrated that melanoidins can be used by gut microbes resulting in SCFAs production and also shaping their communities. Many melanoidins favored the growth of beneficial genera such as *Bifidobacterium* and *Faecalibacterium*. Gut microbes, by fermenting melanoidins, were able to release some phenolics initially linked to the melanoidin backbone, which in turn could increase phenolics absorption. Such analysis of polyphenols could be used to investigate melanoidins structure and also to investigate

385 microbial pathways. Therefore, these results suggest that melanoidins should be considered386 as potential prebiotic agents.

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394 Conflict of interest

The authors declare that there is no conflict of interest regarding the publication of this paper.

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532 Figure legends

533

Figure 1. Fermentation of melanoidin preparations by human fecal microbiota 534 promotes different community structures. Panel A shows relative abundances of the top 535 nine most abundant microbial genera across all samples. Each column represents a 536 community derived from an independent microbiota-based fermentation of particular 537 538 melanoidin preparation. Abundances of other genera were summed and are represented cumulatively as "other genera". Sample names were abbreviated as follows: BRCR - bread 539 crust; CERL - cereal; BLBR - black beer; PLBR - pilsner beer; BISC - biscuits; CHOC -540 chocolate; SWWN - sweet wine; BLVN - balsamic vinegar; COFF - coffee; SPCF - spent 541 coffee; BLNK - blank. Four groups of samples with similar genus abundance profiles 542 within group are separated by dotted lines. Panel **B** displays the output of the unconstrained 543 PCoA ordination analysis of microbial genus abundance dataset among all profiled 544 samples. Phylogenetic weighted UniFrac distance was used to calculate the sample 545 dissimilarity matrix. The percent of dataset variability explained by each principal 546 coordinate is shown in parentheses in axis titles. Samples are colored based on groups 547 defined in Panel A. Panels C and D show the same PCoA output as in panel B, but with 548 samples colored according to the abundance of Bifidobacterium (panel C) or 549 Faecalibacterium (panel D) genera. 550

Figure 2. Metabolite measurements. Concentrations of short chain fatty acids (in mM),
phenolic compounds (in parts per million), and antioxidants (defined as µmol of Trolox
equivalent/g of sample) and are displayed in panels A, B, and C, respectively.
Concentrations are visualized in log-scale and are represented by color gradient as shown

next to each heatmap. Four groups of samples defined in Figure 1A are separated by
horizontal black lines. Sample abbreviations are defined in Figure 1 legend.

557 Figure 3. Ordination analysis of metabolite data. Panels A and B display the output of Euclidean-distance based principal component analysis of short chain fatty acids and 558 phenolic compounds, respectively. Due to vast differences in concentrations among 559 different compounds, all values for each compound were standardized across samples. 560 Sample abbreviations are defined in Figure 1 legend. Co-inertia analysis depicted in panel 561 C reveals congruency of sample dispersal in ordination space based on SCFAs and genus 562 abundance profiles. The distance between each sample position on two ordination plots is 563 indicated by a connecting line. Shorter lines represent similar sample positioning in each 564 plot. The statistical significance and the relative 'fit' of the ordinations were assessed by p-565 value and RV coefficient, respectively. 566

Table 1. Microbial phenolic metabolites, parent phenolic compounds and microbial types (Data obtained from Selma et al., 2009; Marín et al., 2015; Rowland et al., 2018).

5	69
0	05

Phenolic compound group	Phenolic compound sub- group	Parent compound	Principal metabolites	Microbial types
Phenolic acids	Benzoic acids	Gallic acid	Urolithins A & B, isourolithins A & B	Gordonibacterurolithinfaciens, Gordonibacterpamelaeae
		Ellagitannins	Pyrogallol	
	Hydroxycinnamic acids	Chlorogenic acid	3-(3,4-Dihydroxyphenyl)- propionic acid	Escherichia coli, Bifidobacterium lactis, Lactobacillus gasseri
		Hydroxycinnamates	3-(3-hydroxyphenyl)- propionic acid	
			3-(4-hydroxyphenyl)- propionic acid	
			Hydroxyphenyl-ethanol	
			Vanillin	
			Phenylacetic acids and Benzoic acids	
Flavonoids	Flavonols	Kaempferol	2-(4-	Clostridium orbiscidens,
			Hydroxyphenyl)propionic acid	Enterococcus casseliflavus
		Kaempferol	2-(3,4- Dihydroxyphenyl)acetic acid	C. orbiscidens, Eubacterium
		Kaempferol	2-(3-hydroxyphenyl)acetic acid	Clostridium strains
	0	Quercetin	2-(3,4- dihydroxyphenyl)acetic acid	Clostridium orbiscindens, Eubacteriumoxidoreducens, Butyrivibrio spp.

		Isoxanthohumol (from hops)	8-Prenyl-naringenin	Eubacteriumlimosum
			3- phenylpropionic acid	
			acid	
1 14 1 41		Traingonn	Hydroxyphenyl)propionic	Eubacteriumramulus
Flava	nones	Naringenin	3-(4-	Clostridium strains
			2-(3-Hydroxyphenyl)acetic	
		<u> </u>	Dinydroxyphenyl)acetic acid	
			2-(3,5-	
			acid	
			Hydroxyphenyl)propionic	oxidoreducens
			3-(3-	Clostridium orbiscidens, E.
			acid	
			Dihydroxyphenyl)propionic	
			3-(3,4-	Eubacteriumramulus
		wiyneetiii	acid	
		Muricetin	2 (3 Hydroxynhenyl)scotic	Clostridium strains
		Quereetiii	acetic acid	Eubacterium oxidoreducens
		Quercetin	3 (3 4 Dihydroxymhenyl)	Clostridium orbiscidans
		Quereetiii	acid (vanillic acid)	
			3-methovy-A-bydrovybenzoic	
		Quercetin	3.4 dihydroxybenzoic acid	
		Quercetin	<u>acia</u> <u>3 4-dihydroxybenzaldehyde</u>	
			dihydroxyphenyl)propionic	
		Quercetin	3-(3,4-	
			acid	
		Quercetin	2-(3-nydroxypnenyl)acetic	

Flavan-3-ols	Catechin	3-(3-	Clostridium coccoides,
		Hydroxyphenyl)propionic	Bifidobacterium infantis
		acid	
		5-(3',4'-Dihydroxyphenyl)	
		valero-lactone	
	Epicatechin	5-(3,4-	
		Dihydroxyphenyl)valeric	
		acid	
		3-(3,4-	
		Dihydroxyphenyl)propionic	
		acid	
		Pyrogallol	
	Epigallocatechin	5-(3',4'-Dihydroxyphenyl)-γ-	
		valero-lactone	
		5-(3',5'-Dihydroxyphenyl)-γ-	
		valero-lactone	
Flavones	Luteolin	3-(3,4-Dihydroxyphenyl)-	Clostridium. orbiscindens,
		propionic acid	Enterococcus avium
	Apigenin	3-(4-hydroxyphenyl)-	Eubacteriumramulus,
		propionic acid	Bacteroidesdistasonis
		3-(3-hydroxyphenyl)-	
		propionic acid	
		4-hydroxycinnamic acid,	
		phloretin	
Isoflavones	Daidzein	Equol	Bacteroidesovatus, Streptococcus
			intermedius,
			Ruminococcusproductus
	7	O-desmethylangolensin	

			Eggerthellasp.Julong 732, Slakiaequolifaciens, Adlercreutziaisoflavoniconvertens, Slakiaequolifaciens
		0	Consortium of Lactobacillus, mucosae Enterococcus faecium, Finegoldia magna, Veillonella spp.
			Clostridium spp. HGHA136, Eubacteriumramulus
	Formononetin	Daidzein	
	Genistein	6'-ОН-О-	Eubacteriumramulus
		desmethylangolensin	
		2-(4-	
		hydroxyphenyl)propionic	
A with a swamiding	Cronidia	acid	
	Cyanidin	5,4-Dinydroxybenzoic acid	Clostridium saccharogumia, _ Eubacteriumramulus,
	Peonidin	3-Methoxy4-hydroxybenzoic acid	Lactobacillus plantarum, Lactobacillus
	Pelargonidin	3-hydroxycinnamic acid	casei, Lactobacillus acidophilus
	Malvidin	4-Hydroxybenzoic acid	- LA-5,
		3,4-Dimethoxybenzoic acid	Bifidobacterium lactis BB-12
30			

	Lignans	Lignans	Secoisolaricinresinoldiglucoside	Enterodiol	Bact. distasonis, Bact. fragilis, Bact. ovatus, Clostridium cocleatum, Clostridium.sp SDG- MT85-3Db, Butyribacteriummethylotrophicum, Eubacteriumcallanderi, Eubacteriumlimosum, Peptostreptococcusproductus, Clostridium scindens
				Enterolactone	Eggerthellalenta, ED-Mt61/PY-s6
	Secoiridoids	Secoiridoids	Oleuropein	Tyrosol	
			Ligstroside	Hydroxytyrosol	
570 571					

Table 2. Antioxidant capacity of melanoidins prior and after *in vitro* fermentation. Sample names were abbreviated as follows: BRCR
 bread crust; CERL – cereal; BLBR - black beer; PLBR - pilsner beer; BISC – biscuits; CHOC – chocolate; SWWN - sweet wine;

- 574 BLVN balsamic vinegar; COFF coffee; SPCF spent coffee grounds.
- 575 576

Sample	FRAP (µmol Trolox equivalent/g of sample)			FRAPABTSDPPH(μmol Trolox(μmol Trolox(μmol Troloxequivalent/g ofequivalent/g ofequivalent/g ofsample)sample)sample)						F (µmo equiva sai	RAI I Tr ilen npl	P colox t/g of e)	Α (μmo equiv sa	TS Trolox nt/g of le)	DPPH (µmol Trolox equivalent/g of sample)			
	Digested melanoidins								Fermented melanoidins									
BRCR	17.4	±	0.5	65.4	±	2.0*	25.6	<u>±</u>	0.8	16.9	±	0.9	52.2	±	2.0	15.5	±	1.9
CERL	3.05	±	0.07*	12.8	±	0.3*	1.64	±	0.04*	29.1	±	1.0	65.4	±	5.0	4.8	±	0.3
BLBR	63.6	±	2.9*	347	±	14.9*	52.9	±	2.4	47.3	±	2.3	203	±	3.0	46.0	±	1.8
PLBR	36.6	±	1.8*	175	±	9.3*	42.0	±	2.1*	21.7	±	1.5	98.7	±	3.6	22.2	±	1.1
BISC	43.1	±	1.5*	154	±	5.8*	12.0	±	0.5	103	±	3.6	319	±	1.3	25.5	±	1.9
CHOC	164	±	11.2	511	±	32.0*	97.9	±	6.7	108	±	2.7	232	±	12.9	103	±	5.6
SWWN	464	±	17.1	946	±	35.1*	539	±	19.0*	433	±	5.3	688	±	9.9	263	±	5.2
BLVN	219	±	3.2**	541	±	7.8**	139	±	2.0**	339	±	3.9	628	±	8.2	225	±	3.5
COFF	1475	±	41.0**	2700	±	78.1*	1595	±	46.1*	674	±	21.1	1065	±	13.4	628	±	5.6
SPCF	126	±	2.8*	198	±	4.6*	168	±	3.8	1048	±	19.7	1511	±	71.3	735	±	29.0

⁵⁷⁷ *whithin the column means statistical significance between digested and fermented melanoidins.



595	Figure 2																				
590 597	А	SCFAs		B Phenolics										С	its						
597 598 599 600 601 602 603 604 605 606 607	Acetate Butyrate Lactate	BRCR CERL BLBR PLBR BISC CHOC SWWN BLVN COFF SPCF BLNK	0 6 115	Chlorogenic acid	2-(3,4-Dihydroxyphenyl)acetic acid 3-(3,4-Dihydroxyphenyl)propionic acid	3,4-Dihydroxybenzaldenyde 3,4-Dihydroxybenzoic acid 3,4-Dimethoxybenzaldehyde	3,5-Dihytoxybenzoic acid	3,4-UIIITEUTOXYUETIZOIC acid 4-O-Caffeoylquinic acid		Epigallocatechin galate	Genistein	Hydroxymethyl-furan-carbaldehyde	-Coumaric acid	p-Coumaric acid p-Hydroxyphenylacetic acid	Pyrogallol Resveratrol	Trans-hydroxycinnamic (d) acid	BRCR CERL BLBR PLBR BISC CHOC SWWN BLVN COFF SPCF BLNK	0 1 55	ABTS	BRCR CERL BLBR PLBR BISC CHOC SWWN BLVN COFF SPCF BLNK	0 10 1510



624 3. Melanoidins increase SCFAs production by gut microbes.

- 625 4. Melanoidins metabolization release phenolic compounds.
- 626
- 627
- 628 Declaration of interests
- 629

K The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence
 the work reported in this paper.

633 The authors declare the following financial interests/personal relationships which may be considered as potential competing interests:

Credit Authors statement

643 Sergio Pérez Burillo: investigation, methodology and paper drafting; Sumudu Rajakamura: investigation, methodology and formal
 644 analysis; Oleg Paliy: formal analysis, conceptualization, paper drafting and founding acquisition; Silvia Pastoriza: conceptualization,
 645 supervision and review; José Ángel Rufián-Henares: validation, formal analysis, writing and review, founding acquisition.

- 04