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Time Course Production of Urolithins from Ellagic Acid by Human Gut Microbiota

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ABSTRACT: Ellagic acid (EA) is converted to urolithins by gut microbiota. Urolithins have beneficial biological effects in humans, but differences in urolithin production capacity among individuals have been shown. Therefore, the identification of the urolithin production pathways and the microorganisms implicated is of high interest. EA was incubated with gut microbiota from two volunteers able to produce urolithins but with different in vivo urolithin profiles (urolithin A and isourolithin A producers). The metabolic capabilities observed in vivo were retained in vitro. Both individuals showed a much higher abundance of Clostridium leptum group of Firmicutes phylum than Bacteroides/Prevotella. EA was either dissolved in DMSO or suspended in water. DMSO increased EA solubility but decreased urolithin production rate due to a delay in growth of some microbial groups, principally, Clostridium coccoides. This allowed the detection of catabolic intermediates [urolithins M-5, M-6, M-7, C, and 2,3,8,10-tetrahydroxy urolithin (urolithin E)]. Bacteria from C. coccoides group (or genera co-occurring in vivo with this group) seem to be involved in production of different urolithins.

KEYWORDS: ellagitannins, colon microbiota, metabolism, dibenzopyranones, polyphenols, human health

17 INTRODUCTION

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18 Ellagitannins and ellagic acid (EA) are plant secondary 19 metabolites that have relevant antioxidant activities in vitro, 20 potential cardiovascular protection, and anticarcinogenic and 21 anti-inflammatory effects. ¹ These phytochemicals are relevant 22 constituents in different foods including pomegranates, berries 23 (strawberry, raspberry, blackberry, camu-camu, etc.), nuts 24 (walnuts, acorns, chestnuts, etc.), muscadine grapes, oak-aged 25 wines, and medicinal plants and tisanes (geranium, oak leaves, 26 etc.). They are not absorbed in the gut and are metabolized in 27 vivo by the gut microbiota to produce a series of metabolites 28 known as urolithins. 4,5 Therefore, urolithins have been 29 suggested as biomarkers of intake for strawberries, raspberries, 30 pomegranates, walnuts, and oak-aged red wines. We have 31 previously suggested that urolithins, that are much better 32 absorbed than the original ellagitannins, could be responsible 33 for the systemic biological effects of ellagitannins. 2,5 Additional 34 evidence supports this suggestion as urolithins and their 35 glucuronide and sulfate metabolites have shown relevant effects 36 at the concentrations found in vivo.^{7–9} Urolithins share a 37 nucleus of a dibenzo-pyran-one, with different hydroxyl 38 substitutions. The first reported urolithins were urolithin A 39 and urolithin B that were found in kidney stones of sheep. 10 In 40 a recent study, it has been demonstrated that other mammals 41 also produce urolithins from ellagitannins, although they have 42 different structures and different hydroxylation patterns on the 43 urolithin nucleus. 11 Thus, isourolithin A has been found as the 44 main in vivo metabolite in beef cattle after oak leaf intake, ¹² and 45 a series of tetrahydroxy and trihydroxy urolithin metabolites 46 were produced by rats after Geranium extracts intake. 13

Recently, it has been reported that some humans are able to produce isourolithin A as the main in vivo metabolite instead of urolithin A after the intake of strawberries¹⁴ and raspberries.¹⁵ Furthermore, marked person-to-person differences have been observed in the level of in vivo urolithin production following 51 consumption of ellagitannin-containing food. In vitro anaerobic 52 metabolism of EA to urolithin A in human fecal suspensions 53 has also been reported. More recently, in vitro anaerobic 54 incubation of EA with human fecal suspensions demonstrated 55 conversion to some other urolithins, urolithins B, C, and 56 isourolithin A. However, neither comparison between the in 57 vitro and the in vivo urolithin patterns of these volunteers nor 58 their endogenous microbiota has been analyzed in any of these 59 studies.

In the present study, we evaluated the in vitro time course 61 production of urolithins from EA by human fecal microbiota 62 from two volunteers who have urolithin A and isourolithin A 63 urinary excretion patterns. Detailed in vitro and in vivo 64 experiments were performed to identify the intermediary and 65 final catabolites of EA conversion by the human gut microbiota 66 to elucidate interindividual urolithin production routes and to 67 determine if EA metabolism could be linked to the microbiota 68 composition of these individuals.

MATERIALS AND METHODS

Chemicals. EA and 6,7-dihydroxycoumarin were obtained from 71 Sigma-Aldrich (St. Louis, MO, USA). Urolithins A and B were 72 chemically synthesized by Villapharma SL (Parque Tecnológico de 73 Fuente Álamo, Murcia, Spain). Urolithins C (3,7,8-tetrahydroxy-6H- 74 dibenzo[b,d]pyran-6-one, Uro-C; > 95% purity) and D (2,3,7,8- 75 tetrahydroxy-6H-dibenzo[b,d]pyran-6-one, Uro-D; > 95% purity) were 76 purchased from Dalton Pharma Services (Toronto, Canada). 77 Methanol and acetonitrile were purchased from Romil (Barcelona, 78 Spain) and ethyl acetate and dimethyl sulfoxide from Labscan (Dublin, 79

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80 Ireland). Formic acid and hydrochloric acid was obtained from Panreac (Barcelona, Spain). Milli-Q system (Millipore Corp., Bedford, MA) ultrapure water was used throughout this experiment. Nutrient Broth (NB) was from Oxoid (Basingstoke, Hampshire, UK). L-84 Cysteine hydrocloride was from (Panreac Química, Barcelona, Spain). Dimethyl sulfoxide (DMSO) was from (Scharlab, Barcelona, Spain). All chemicals and reagents used in the preparation of buffers, macromineral, micromineral, and reducing solutions were obtained from Sigma-Aldrich, Panreac, and Scharlab.

Collection of Human Fecal Samples. These were donated by a healthy female (age 31) and a healthy male (age 36) in the Centro de Edafología y Biología Aplicada del Segura (CEBAS-CSIC, Murcia, Spain). These two volunteers were identified as urolithin A and isourolithin A producers, respectively, in a previous study in which urine samples were analyzed after the intake of strawberries and strawberry jam, although these metabolic groups were not reported. To conduct fermentation experiments in duplicate, each volunteer donated two fecal samples. They were stored at 4 °C and were further processed within 1 h of donation, as no differences in microbial concentrations were observed in fresh samples before and after processing samples for fermentation experiments. Anoxic conditions were maintained by using the AnaerobeGen compact system (Oxoid).

Fermentation Medium. This was prepared as described by 103 Jaganath et al. ¹⁹ with some modifications. Briefly, 2 g of tryptone, 2 g 104 of glucose, 1 g of maltose and 2 g of yeast extract were mixed in 400 105 mL of distilled water and 100 μL of micromineral solutions (consisting 106 of 13.2 g of CaCl₂·2H₂O, 10 g of MnCl₂·4H₂O, 1 g of CoCl₂·6H₂O, 8 107 g of FeCl₃·6H₂O, and distilled water up to 100 mL), 200 mL of buffer 108 solution (2 g of NH₃CO₃, 17.5 g Na₂CO₃, and distilled water up 500 mL), 200 mL of macromineral solution (2.85 g of NA₂HPO₄, 3.1 g 110 KH₂PO₄, 0.3 g MgSO₄·7H₂O, and distilled water up to 500 mL), 1 mL 111 of 1% (w/v) of resazurin solution (a redox indicator) and 0.5 mg/L of 112 Vit K₁, and 5 mg/L and 625 mg/L hemein. The medium was adjusted 113 to pH 7 using HCl, dispensed into the fermentation vessels, and 114 autoclaved at 121 °C for 15 min and allowed to cool under oxygen-115 free nitrogen (OFN) to remove oxygen.

Conversion Experiments of EA into Urolithins with Human 117 Fecal Cultures. Preparation of fecal suspensions and subsequent 118 culturing experiments were conducted under anoxic conditions in an 119 anaerobic chamber (Don Whitley Scientific Limited, Shipley, UK) 120 with an atmosphere consisting of N₂/H₂/CO₂ (80:10:10) at 37 °C. 121 Aliquots of fecal samples (10 g) were diluted 1/10 (w/v) in NB 122 supplemented with 0.05% L-cysteine hydrochloride and homogenized 123 by stomacher in filter bags. Aliquots of filtered fecal suspensions (2 124 mL) were inoculated into 200 mL of prereduced fermentation 125 medium containing EA at 30 μ M with and without 1% DMSO. DMSO 126 was used in order to increase EA solubility, and EA dissolved in water 127 instead of DMSO was used as control. Duplicate cultures were 128 prepared in parallel from each fecal suspension. In addition, controls 129 were used; ones without fecal suspension and others without EA. 130 Samples (6 mL) were collected at appropriate time intervals during 8 131 day incubation at 37 °C and stored at -20 °C until further HPLC 132 analysis. Similarly, 3 mL were collected every 2 h during 2 day 133 incubation at 37 °C and stored at −20 °C until further qPCR analysis. Sample Cleanup for LC Analyses. Feces and urine of the two 135 volunteers were extracted and analyzed. Urine samples were defrosted, 136 vortexed, and centrifuged at 14000g for 10 min at 4 °C. The 137 supernatant was filtered through a 0.45 μ m PVDF filter and analyzed 138 by HPLC-DAD-IT MS. When the intensity was not enough to 139 distinguish the metabolites, urine samples were concentrated using a 140 Sep-Pak reverse phase C-18 extraction cartridge (Waters Millipore). 141 The cartridges were previously activated with 10 mL of MeOH and 10 142 mL of water. Then 25 mL of urine acidified with 250 μ L of formic acid 143 (1%) were passed through the cartridge that was then dried with air. 144 The metabolites remaining in the cartridge were eluted with 3 mL of 145 MeOH/H2O (50:50, v/v). Samples were analyzed after filtration 146 through 0.45 μ m PVDF filter. Feces samples (1 g) were defrosted and 147 homogenized with 10 mL of MeOH/DMSO/H2O (40:40:20) with 148 0.1% HCl using an Ultra-Turrax for 1 min at 24000 rpm. The mixture

was centrifuged at 5000g for 10 min at room temperature and the 149 supernatant filtered through a 0.45 μ m PVDF filter before analysis.

Samples (5 mL) obtained in fermentation experiments with human 151 faecal suspensions were extracted with 5 mL of ethyl acetate acidified 152 with 1.5% formic acid. The mixture was vortexed for 2 min and 153 centrifuged at 3500g for 10 min. The organic phase was separated and 154 evaporated under reduced pressure until dryness. The dry samples 155 were then redissolved in 250 μ L of methanol and filtered through a 156 0.45 μ m PVDF filter. Then 5 μ L of 100 μ g/mL of internal standard 157 (6,7-dihydroxycoumarin) was added to 50 μ L of sample prior to the 158 injection onto a column for LC-UV/vis and LC-MS analysis under the 159 conditions described below.

LC-UV/Vis and LC-MS/MS Analyses. The analyses were 161 performed using an Agilent 1100 HPLC system equipped with a 162 photodiode array detector and an ion-trap mass spectrometer detector 163 in series (Agilent Technologies, Waldbronn, Germany). Chromato- 164 graphic separation was carried out on a reverse phase LiChroCART C- 165 18 column (Merck, Darmstadt, Germany) (250 mm \times 4 mm, 4.5 μ m 166 particle size) using water with 1% formic acid (A) and acetonitrile (B) 167 as the mobile phases. The gradient profile was: 0–20 min, 5–30% B, 168 20–30 min, 30–55% B, 30–38 min, and 55–90% B, and this 169 percentage was maintained for 2 min and then came back to the initial 170 conditions. A volume of 10 μ L of sample was injected onto the column 171 operating at room temperature and a flow rate of 1 mL/min. UV 172 chromatograms were recorded at 280, 360, and 305 nm.

The HPLC system was coupled in series to an ion trap mass 174 spectrometer (IT) equipped with an electrospray interface (ESI). 175 Nitrogen was used as drying gas with flow of 11 L/min and 176 temperature of 350 $^{\circ}$ C and nebulizing gas at pressure of 65 psi. The 177 capillary voltage was set at 4 kV. Mass scan (MS) and daughter (MS- 178 MS) spectra were recorded in negative mode in the range of m/z 179 100–700 with target mass of 300. Maximum accumulation time of ion 180 trap and the number of MS repetitions to obtain the MS average 181 spectra were set at 200 ms and 3, respectively. Compound stability was 182 set at 75%.

Identification of all metabolites was carried out by direct 184 comparison (UV spectra and MS) with pure standards and confirmed 185 by their spectral properties and molecular mass. Calibration curves 186 were obtained for EA, urolithin A, urolithin B, and urolithin C with 187 good linearity ($R^2 > 0.998$). EA was quantified at 360 nm and 188 urolithins at 305 nm. The limits of detection (LODs) were determined 189 based on a signal-to-noise ratio (S/N) of 3 and of 10 for the limit of 190 quantification (LOQ). EA, urolithin A, and urolithin B showed LODs 191 of 0.5 μ M and LOQs of 1.67 μ M and urolithin C an LOD of 0.2 μ M 192 and an LOQ of 0.67 μ M. Repeatability was evaluated by injecting 20 193 μM of a mixture of standards four times in the same day (intraday 194 repeatability) and in four different days (interday repeatability). The 195 results expressed as the relative standard deviation (RSD) of peak area 196 were ≤5% for intraday repeatability and ≤8% for interday repeatability. 197 The recovery of the compounds was calculated, spiking the medium in 198 the presence of inactivated bacteria with a standard solution of EA, 199 urolithin A, and urolithin B in DMSO at a final concentration of 20 200 μ M. Recoveries of 75%, 90%, and 83% were obtained for EA, urolithin 201 A, and urolithin B, respectively. Isourolithin A was quantified at 305 202 nm with the urolithin A calibration curve and urolithins M-5, M-6, and 203 M-7 at 360 nm with the EA calibration curve.

Fecal Dry Weight Determination and DNA Extraction. To 205 determine fecal moisture content, approximately 0.5 g (wet weight) of 206 each fecal specimen was placed in a vacuum dryer for 3 d and 207 reweighed. Percent fecal dry weight was calculated. Total DNA was 208 extracted from human fecal samples using a commercial DNA 209 extraction kit (QIAampR DNA Stool Mini Kit, Qiagen Inc., Valencia, 210 CA). Although this kit did not contain beads, an additional step of 211 vigorous shaking using the FastPrep Instrument was carried out. 212 Briefly, 10 mg of fresh fecal samples (25.3 ± 8.9% dry matter) were 213 added to 2 mL tubes containing specialized beads (MP Biomedicals, 214 LLC, Ohio, USA) and homogenized in the lysing matrix using the 215 FastPrep Instrument for 30 s at a speed setting of 5.5. The following 216 steps were carried out attending protocol supplied with the kit: briefly, 217 incubation temperatures (95 °C, 5 min and 70 °C, 10 min), cell lysis 218

Table 1. Primer and Probe Sequences Used for Detection of Main Bacterial Groups in Human Feces by Quantitative Real-Time Polymerase Chain Reaction

target organism groups and species ²⁴	primers and probes	sequence 5'-3'	refs
Clostridium leptum group	F_Clept 09	CCT TCC GTG CCG SAG TTA	24
Clostridium leptum	R_Clept 08	GAA TTA AAC CAC ATA CTC CAC TGC TT	
Faecalibacterium prausnitzii	P_Clep 01 ^a	6FAM-CAC AATAAG TAA TCC ACC	
Ruminococus albus			
ifidobacterium genus	F_Bifid 09c	CGG GTG AGT AAT GCG TGA CC	24
Bifidobacterium adolescentis	R_Bifid 06	TGA TAG GAC GCG ACC CCA	
Bifidobacterium breve	P_Bifid ^a	6FAM-CTC CTG GAA ACG GGT G	
Bifidobacterium infantis			
Clostridium coccoides group	F_Ccoc 07	GAC GCC GCG TGA AGG A	24
Clostridium coccoides	R_Ccoc 14	AGC CCC AGC CTT TCA CAT C	
Ruminococcus gnavus	P_Erec482 ^a	VIC-CGG TAC CTG ACT AAG AAG	25
Ruminococcus hansenii			
Eubacterium rectale			
Bacteroides/Prevotella group	F_Bacter 11	CCT WCG ATG GAT AGG GGT T	24
Bacteroides fragilis	R_Bacter 08	CAC GCT ACT TGG CTG GTT CAG	
Bacteroides ovatus	P_Bac303 ^a	VIC-AAG GTC CCC CAC ATT G	26
Bacteroides thetaiotaomicron			
Bacteroides uniformis			
Bacteroides vulgatus			
Bacteroides caccae			
Bacteroides eggerthii			
Prevotella oralis			
Prevotella buccae			
Prevotella albensis			
Escherichia coli species	E.coli F	CAT GCC GCG TGT ATG AAG AA	27
	E.coli R	CGG GTA ACG TCA ATG AGC AAA	
Lactobacillus/Leuconostoc/Pediococcus group	F_Lacto 05	AGC AGT AGG GAA TCT TCC A	24
Lactobacillus acidophilus	R_Lacto 04	CGC CAC TGG TGT TCY TCC ATA TA	
Lactobacillus casei			
Lactobacillus paracasei			
Lactobacillus delbrueckii			
Lactobacillus fermentum			
Lactobacillus johnsonii			
Lactobacillus plantarum			
Lactobacillus rhamnosus			
Lactobacillus helvetic			
Pediococcus inopinatus Pediococcus parvulus			
Pediococcus parvuius Pediococcus cellicola			
Pediococcus cententi Pediococcus acidilactici			
Leuconostoc mesenteroides			
Leuconostoc pseudomesenteroides			
-			
Probe sequences.			

 $_{219}$ and homogenization (centrifugation at 14,000 rpm, 1 min), adsorption $_{220}$ of inhibitors (InhibitEx tablet), and approximate time to completion $_{221}$ (60–80 min).

Real-Time qPCR. DNA from dominant groups of fecal bacteria was quantified with real-time qPCR and primers as summarized in Table 1. Real-time qPCR was performed using an ABI 7500 sequence detection system. Amplification and detection were carried out in 96-226 well plates with TaqMan Universal PCR 2× master mix (Applied Biosystems) or with SYBR-Green PCR 2× master mix (Applied Biosystems). Each reaction was run in duplicate in a final volume of 25 μ L with 0.2 mM final concentration of each primer, 0.25 mM final

concentration of each probe, and 10 mL of appropriate dilutions of 230 DNA samples. Probes were designed with molecular-groove binding 231 nonfluorescence quencher (MGBNFQ). Amplifications were carried 232 out using the following ramping profile: 1 cycle at 95 °C for 10 min, 233 followed by 40 cycles of 95 °C for 30 s and 60 °C for 1 min. For 234 SYBR-Green amplifications, the following dissociation protocol (a 235 hold at 95 °C for 15 s, a hold at 60 °C for 20 s, and a slow ramp for 20 236 min from 60 to 95 °C) was followed after the last cycle in order to 237 check the expected amplification products.

When PCR was performed on unknown fecal samples, the cycle 239 threshold of each sample was compared with a standard curve 240

241 (performed in triplicate) made by diluting genomic DNA (6-fold serial 242 dilutions of *Bifidobacterium longum* DSM 20088T for *Bifidobacterium* 243 genus, *Escherichia coli* CECT 515T for *E. coli* species, *Clostridium* 244 leptum DSM 753T and *Clostridium coccoides* DSM 935T for *C. leptum* 245 and *C. coccoides* groups, *Bacteroides ovatus* DSM 1896T for *Bacteroides/* 246 *Prevotella* group and *Lactobacillus plantarum* CECT 748T for 247 *Lactobacillus/Leuconostoc/Pediococcus* group). Selected bacteria strains 248 were cultured anaerobically on selective broth as recommended by 249 DSMZ. For each culture, the total number of bacteria, in terms of 250 CFU, was determined by plating. Aliquots of 1 mL of culture were 251 used for DNA extraction by DNeasy blood and tissue kit (Quiagen) 252 following the manufacturer's instructions.

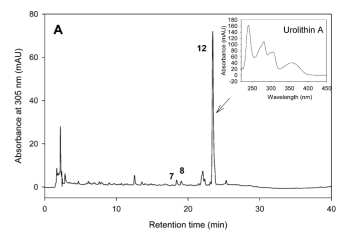
Statistical Analyses and Data Modeling. Fecal samples were analyzed in triplicate, and the quantitative estimates represent mean values ± standard error. Data were subjected to statistical analysis of variance (ANOVA). Normality of variances was tested by the Shapiro—Wilk test before determining the ANOVA. Statistical analyses were performed using SPSS V.14 for Windows.

Microbial growth curves were fitted using the function of Baranyi et 260 al. 20 to estimate the main growth parameters (maximum specific 261 growth rate, lag time of microorganisms before the onset of growth, 262 and estimated correlation coefficient that indicates the goodness of fit 263 of the parameters derived from experimental data). Only growth 264 curves with at least 10 data points were used for modeling, as 265 suggested by the authors.

266 RESULTS

In Vivo Metabolism of EA and Ellagitanins by Human 267 Gut Microbiota. Two volunteers who had different routes of 268 269 EA metabolism in vivo were selected for the present study. Volunteer 1 produced urolithin A, and volunteer 2 produced 271 isourolithin A. We had previously identified these two urolithin 272 metabolic profile groups in urine samples from different volunteers after the intake of walnuts, strawberries, and 274 raspberries (unpublished data). The analyses of fecal samples 275 from both volunteers after the intake of walnuts (0.6 g/kg body 276 weight per day for 5 days) showed that volunteer 1 had a chromatographic profile characterized by urolithin A that was identified by its MS spectrum, its characteristic UV spectrum, 11 and by chromatographic comparisons with an authentic standard, while volunteer 2 produced isourolithin A as a main metabolite (Figure 1). The UV spectra of urolithin A and isourolithin A allow a clear and fast identification of both metabolites. 11 In addition, other minor metabolites were also detected in fecal samples of both volunteers [urolithin M-6 (7) and urolithin C (8)] (Table 2). When the urine of the same volunteers was analyzed, their metabolic profiles were consistent with those found in feces, and volunteer 1 excreted mainly urolithin A glucuronide (1), while volunteer 2 excreted two isomers of isourolithin A glucuronide (2, 3) instead (Figure 2). Trace amounts of urolithin B-glucuronide (10) were also detected in the urine of volunteer 2 (Figure 2B).

Bacterial Populations of Human Fecal Samples Used in This Study. Fecal bacteria densities of the two volunteers, who donated feces for the investigation of the metabolic fate of EA in vivo, were also analyzed. Interindividual differences in fecal microbiota were analyzed by qRT-PCR using primers and probes (Table 1) that target the main bacterial groups: Bifidobacterium genus; C. leptum group (C. leptum, Faecalibactery rium prausnitzii, and some species of Ruminococcus genus); C. soo coccoides group (C. coccoides, Eubacterium rectale spp., and some species of Ruminococcus genus); E. coli species; Bacteroides/Prevotella group; Lactobacillus/Leuconostoc/Pediococcus group (Table 1). All bacteria results were presented as the mean of the log CFU per g feces. Fecal dry weight (%) was 31.7 and



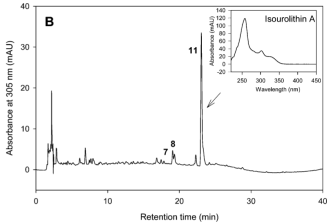


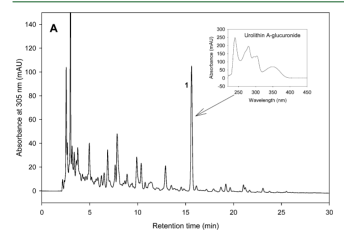
Figure 1. HPLC-UV analysis at 305 nm of fecal samples of volunteer 1 (A) and volunteer 2 (B) after the intake of walnuts (0.6 g/kg body weight per day for 5 days). (7) urolithin M-6; (8) urolithin C; (11) isourolithin A; (12) urolithin A.

19.0% in volunteers 1 and 2. To overcome the fact that fecal $_{305}$ samples contained different water, a correction factor ($_{19/31.7}$ $_{306}$ = 0.6) was applied to microbial data of volunteer 1.

Fecal sample analyses from both urolithin A (volunteer 1) 308 and isourolithin A (volunteer 2) producers were analyzed 309 (Figure 3). This study showed that one of the most highly 310 f3 represented bacterial groups, in human stools of both 311 volunteers, was the C. leptum group of the Firmicutes phylum. 312 No significant differences in levels of C. leptum group $(7-34 \times 313)$ 10^7 CFU/g) and C. coccoides group $((5-8) \times 10^6$ CFU/g) of 314 the Firmicutes phylum as well as the Bacteroides/Prevotella group 315 $(5-12 \times 10^{3} \text{ CFU/g})$ of the Bacteroidetes phylum were 316 observed between fecal samples from volunteers 1 and 2. 317 However, concentrations of other bacterial groups in the 318 volunteer 1 fecal sample were noticeably different from those of 319 the volunteer 2 fecal sample ($P \le 0.01$) (Figure 3). Thus, the 320 microbiota from volunteer 1 showed a higher concentration of 321 Bifidobacterium genus (6 \times 10⁸ CFU/g) of the Actinobacteria 322 phylum when compared with that of volunteer 2 (1 \times 10⁶ log 323 CFU/g) (Figure 3A). Furthermore, in volunteer 1, the fecal 324 sample contained a higher concentration of E. coli species (1 \times 325 10⁶ CFU/g) of the *Proteobacteria* phylum than fecal sample of 326 volunteer 2 (8 \times 10⁴ CFU/g). In contrast, a lower abundance 327 of Lactobacillus/Leuconostoc/Pediococcus group of the Firmicutes 328 phylum was found in volunteer 1 (2 \times 10⁴ CFU/g) which were 329 2.7 log units lower than counts for fecal sample of volunteer 2 330 $(1 \times 10^7 \text{ CFU/g})$ (Figure 3). These studies showed that the 331

Table 2. Metabolites identified in urine (U), Feces (F), Fermentation Medium Incubated with Human Fecal Suspensions in the Presence of Ellagic Acid (FM) or in the Presence or Ellagic Acid Dissolved in 1% DMSO (FM-DMSO) (sh, Shoulder)^a

no.	compds	volunteers	origins	$R_{\rm t}$ (min)	$[M - H]^{-}$	$\lambda_{ m max}$
1	urolithin A glucuronide	1	U	15.75	403	246, 278, 303, 35
2	isourolithin A glucuronide	2	U	15.52	403	255, 299sh, 325
3	isourolithin A glucuronide	2	U	16.14	403	255, 299sh, 325
4	urolithin M-5	1	FM and FM-DMSO	14.65	275	261, 292sh,352
		2	FM-DMSO			
5	urolithin E	1	FM-DMSO	16.55	259	252, 276sh,365
6	ellagic acid	1 and 2	FM and FM-DMSO	17.62	301	254, 300sh, 366
7	urolithin M-6	1 and 2	F and FM and FM-DMSO	17.94	259	259, 291sh, 350
8	urolithin C	1	F and FM	18.83	243	255, 304, 349
		2	F and FM and FM-DMSO			
9	urolithin M-7	1	FM and FM-DMSO	20.49	243	250, 280sh, 367
		2	FM-DMSO			
10	urolithin B glucuronide	2	U	20.56	387	249, 273, 297, 320
11	isourolithin A	2	F and FM and FM-DMSO	23.06	227	256, 302, 329
12	urolithin A	1	F and FM and FM-DMSO	24.11	227	246, 280, 305, 356
		2	FM and FM-DMSO			
13	urolithin B	2	FM	28.21	211	249, 276, 301, 333



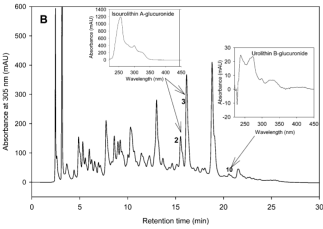
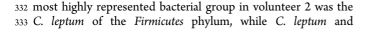


Figure 2. HPLC-UV analysis at 305 nm of urine samples of volunteer 1 (A) and volunteer 2 (B) after the intake of walnuts (0.6 g/kg body weight per day for 5 days). (2) urolithin A glucuronide; (3) isourolithin A glucuronide; (10) urolithin B glucuronide.



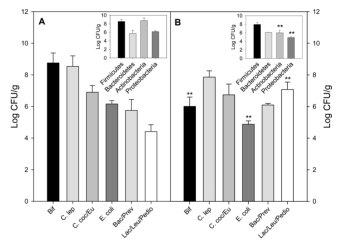


Figure 3. Quantities of dominant groups of bacteria in fecal samples of volunteer 1 (A) and volunteer 2 (B) as determined by real-time polymerase chain reaction. The mean (±SD) counts from triplicate determinations (three DNA samples isolated from the same feces) were used to quantify the bacterial groups. (Bif) Bifidobacterium genus; (C. lep) Clostridium leptum group; (C. coc/Eu) Clostridium coccoides group; (E. coli) E. coli species; (Bac/Prev) Bacteroides/Prevotella group; (Lac/Leu/Pedio) Lactobacillus/Leuconostoc/Pediococcus group. **Significant differences ($P \le 0.01$) in bacteria levels between volunteer 1 (A) and volunteer 2 (B). An estimation of the total amount of Firmicutes was obtained by adding bacterial values obtained from C. coccoides, C. leptum, and Lactobacillus/Leuconostoc/Pediococcus.

Bifidobacterium genus of the Firmicutes and Actinobacteria phyla 334 were the major representative in volunteer 1.

In Vitro Catabolism of EA by Human Gut Microbiota. 336 The production of urolithins by the gut microbiota from the 337 two volunteers was followed in vitro in order to study the time 338 course of the production of the two microbial metabolite 339 patterns found in vivo and identify potential intermediate 340 catabolites in the route from EA to urolithin A or isourolithin A 341 (Figure 4). The first metabolites observed when EA was 342 f4 incubated with the volunteer 1 (urolithin A producer) fecal 343 microbiota were urolithin M-5 (pentahydroxy-urolithin) and 344 urolithin M-6 (tetrahydroxy-urolithin), which peaked at 12 h 345

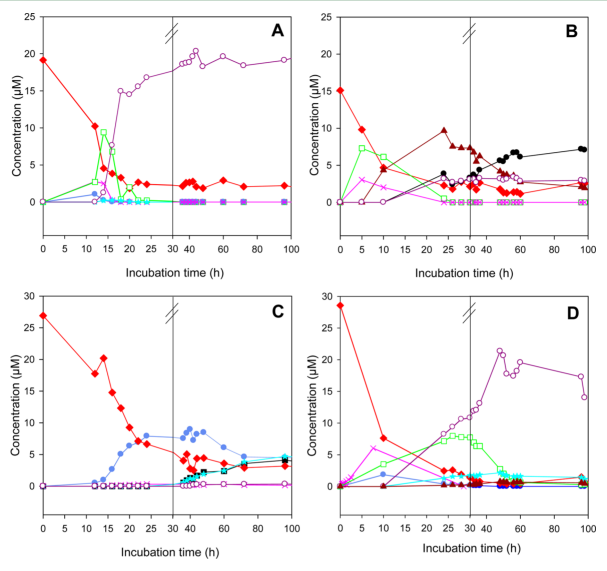


Figure 4. In vitro conversion of EA during batch fermentation with human feces from two volunteers. Human fecal cultures consisted of (A) volunteer 1 fecal suspension in fermentation medium with 30 μ M EA; (B) volunteer 2 fecal suspension in fermentation medium with 30 μ M EA dissolved with 1% DMSO; (D) volunteer 2 fecal suspension in fermentation medium with 30 μ M EA dissolved with 1% DMSO. Ellagic acid (red line with red solid diamonds); urolithin M-5 (blue line with blue solid circles); urolithin E (black lines with black solid squares); urolithin M-6 (magenta lines with magenta Xs); urolithin M-7 (cyan lines with cyan solid stars); urolithin C (green lines with green open squares); urolithin A (purple lines with purple open circles); isourolithin A (brown lines with brown solid triangles); urolithin B (black lines with black solid circles).

346 with a maximum concentration of 1 and 2.5 μ M, respectively 347 (Figure 4A). Then, urolithin C (trihydroxy-urolithin) started to 348 be accumulated reaching 9.4 μ M at 14 h. At that moment, a 349 small amount of another trihydroxy-urolithin (urolithin M-7) 350 started to be produced with a maximum of 0.29 μ M at 16 h. 351 Urolithin A was detected after 14 h incubation, and a maximum 352 of 19.6 μ M was reached at 42 h, then a plateau was maintained 353 (Figure 4A). A large proportion of the EA supplied (30 μ M) 354 was finally converted into urolithin A (21 μ M), although some 355 EA remained unmetabolized in the medium (1.3 μ M).

In the case of volunteer 2 (isourolithin A producer), no urolithin M-5 was detected and both urolithin C and urolithin M-6 appeared with a maximum after 5 h (7.5 and 3.5 μ M, so prespectively) (Figure 4B). Then isourolithin A started to be produced, reaching a maximum of 10 μ M after 24 h. Isourolithin A then decreased, and urolithin B increased to reach 6.5 μ M after 60 h and then reached a plateau. At the same

time as urolithin B was produced, some urolithin A was also $_{363}$ observed, although it reached concentrations much lower than $_{364}$ those found for volunteer 1 (3 μM). The maximum EA $_{365}$ conversion to urolithins was lower in the case of volunteer 2 $_{366}$ fecal fermentation (15 μM) (Figure 4B) than in the case of $_{367}$ volunteer 1 fecal fermentation (21 μM) (Figure 4A) and some $_{368}$ EA remained unmetabolized in the medium (2 μM).

The urolithin production from EA when this was dissolved in 370 DMSO and added to the fermentation medium (1%) to 371 increase its solubility, and potentially its bacterial accessibility, 372 was also evaluated. The EA catabolism was very different from 373 that observed when EA was added to the fermentation medium 374 without adding any organic solvent, and significant changes 375 were observed in growth parameters of some bacterial groups 376 analyzed as described in the section below. In the case of 377 volunteer 1, the main in vitro metabolite produced in the 378 presence of DMSO was urolithin M-5 (pentahydroxy-urolithin) 379

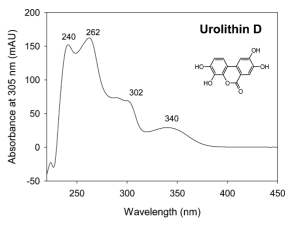
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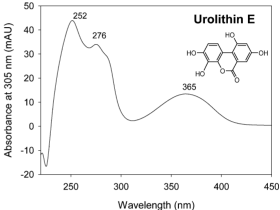
380 instead of urolithin A, which started to be detected after 12 h, 381 with a maximum concentration of 9 μ M at 40 h incubation. 382 Urolithin M-6, urolithin M-7, and urolithin A were also 383 observed but not before 16, 36, and 42 h incubation, 384 respectively, and in much smaller maximal concentrations 385 (0.3, 1.3, and 0.5 μ M, respectively) (Figure 4C) than in the absence of DMSO (2.5, 0.3, and 19.6 μ M, respectively) (Figure 387 4A). Urolithin C was not produced in the presence of DMSO 388 (Figure 4C). Despite the smaller and slower urolithin 389 production in presence of the DMSO, a large proportion of 390 EA supplied was metabolized in the medium (24 μ M) (Figure 391 4C). A relevant unidentified tetrahydroxy urolithin was started 392 to be produced after 36 h of incubation, and this was named as 393 urolithin E (Figure 4C). This had the same mass spectrum as urolithin M-6. To identify the structure of urolithin E, this was compared with the other two available tetrahydroxy-urolithins, 396 urolithin D and urolithin M-6. All of them had the same MS spectrum but differed in the UV spectrum (Figure 5). Urolithin 398 M-6 had a characteristic UV spectrum, and urolithin D had a 399 similar spectrum to urolithin A. Urolithin E had a different UV spectrum, with a significant band I at 360 nm, a characteristic of those urolithins without a hydroxyl at 9-position (Figure 5).¹¹ Therefore, urolithin E was tentatively identified as 3,4,8,10-403 tetrahydroxy-urolithin, a new urolithin metabolite. NMR analysis of this urolithin was not possible due to the small amount produced. 405

In the case of volunteer 2, when EA added was dissolved in DMSO, this started to be converted to urolithin M-5, urolithin 408 M-6, and urolithin C with a maximum concentration of 2, 6, 409 and 7.5 μ M after 7.5, 10, and 25 h, respectively. Then urolithin 410 A started to be produced, reaching a maximum (22 μ M) after 411 40 h (Figure 4D). In this case, only a small concentration of 412 isourolithin A was detected (1 μ M). Urolithin M-7 was also 413 observed and reached maximal concentrations of 3 μ M after 50 414 h. Most EA was metabolized, and only low concentrations (1 415 μ M) remained unmetabolized in the medium (Figure 4D).

Changes in Human Gut Microbiota in Vitro. Changes in 417 the fecal microbiota of volunteers 1 and 2 were followed during 418 the in vitro metabolism of EA in order to elucidate which 419 microbial groups could be implicated in the production of 420 urolithins. The growth characteristics of the main bacterial 421 groups, determined by q-PCR, during the conversion of EA in 422 vitro batch fermentation are shown in Table 3. In the absence 423 of DMSO, the main differences between fecal microbiota 424 development of volunteers 1 and 2 were detected in the 425 Bifidobacterium genus. In fact, the bifidobacteria of volunteer 2 426 were not able to grow in the in vitro conditions despite 427 urolithins being produced. Lactobacillus/Leuconostoc/Pediococ-428 cus group also grew slower in volunteer 2 than in volunteer 1, 429 but a higher level of bacteria was obtained at the end of the 430 growth (N_{max}) in volunteer 2 (5.0 log CFU/mL) than in 431 volunteer 1 culture (4.4 log CFU/mL). In contrast, C. coccoides 432 group reached a higher level in volunteer 1 (N_{max} : 6.1 log CFU/ 433 mL) than in volunteer 2 culture (N_{max} : 5.0 log CFU/mL) 434 (Table 3). In the same way, urolithin concentrations were 435 higher in volunteer 1 than in volunteer 2 culture.

In the presence of DMSO, a delay in the growth of *C.* 437 coccoides group was observed in both volunteer fecal 438 fermentations. Thus, the lag period was extended in 9.7 and 439 22 h in the case of volunteers 1 and 2, respectively (Table 3). 440 For human fecal suspensions of volunteer 1, the extension of 441 the lag phase in 9.7 h of *C. coccoides* group in the presence of 442 DMSO was accompanied by a delay of 10 h in the beginning of





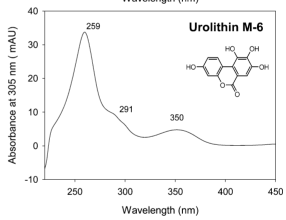


Figure 5. UV spectra of the three tetrahydroxy urolithins ($[M - H]^-$ m/z 259): urolithin D, urolithin E, and urolithin M-6.

urolithin production. A similar tendency was observed in fecal 443 suspension of volunteer 2, where the presence of DMSO 444 lengthens both lag period (24.6 h) and time needed to achieve 445 the maximum urolithin concentration. In contrast, the 446 Bacteroides/Prevotella group achieved higher levels in the 447 absence of DMSO ($N_{\rm max}$: 7.2 log CFU/mL) than when 448 DMSO was present ($N_{\rm max}$: 6.8 log CFU/mL) in the case of 449 volunteer 2 (Table 3).

DISCUSSION

The studies carried out so far with human biological fluids 452 (urine and plasma) have allowed the identification of urolithin 453 A, urolithin B, urolithin C, and isourolithin A.⁵ Other 454 metabolites such as urolithins M-5, M-6, and M-7 had been 455 reported in rat fecal samples and other animal materials.^{11–13} 456

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Table 3. Growth Characteristics of Main Bacterial Groups Determined by q-PCR during Conversion of Ellagic Acid in Vitro Batch Fermentation with Human Fecal Suspensions^a

		-	1	0		1
microbial groups	volunteer	fermentation medium + EA	lag time (h)		$(\log \frac{N_{ m max}}{ m CFU/mL})$	correlation coefficient (R^2)
Clostridium coccoides group	1	without DMSO	0	0.12	6.08	0.99
		with DMSO	9.66** ^a	0.05** ^a	5.87	0.99
	2	without DMSO	2.59	0.10	5.04	0.99
		with DMSO	24.60*** ^a	0.14	5.08	0.99
Bifidobacterium genus	1	without DMSO	0	0.15	7.91	0.92
		with DMSO	0	0.10	7.93	0.83
	2	without DMSO	no growth	no growth	no growth	
		with DMSO	no growth	no growth	no growth	
Clostridium leptum group	1	without DMSO	0	0.07	6.89	0.95
		with DMSO	0	0.04** ^a	6.60* ^a	0.87
	2	without DMSO	0	0.11	7.08	0.99
		with DMSO	0	0.09	7.12	0.97
Lactobacillus/Leuconostoc/Pediococcus	1	without DMSO	0	0.16	4.40	0.99
		with DMSO	0	0.18	4.34	0.99
	2	without DMSO	0	0.02	5.00	0.99
		with DMSO	0	0.05** ^a	4.96	0.99
Bacteroides/Prevotella group	1	without DMSO	0	0.12	6.40	0.96
		with DMSO	0	0.24** ^a	6.41	0.98
	2	without DMSO	0	0.21	7.17	0.99
		with DMSO	0	0.21	6.81* ^a	0.99
E. coli species	1	without DMSO	0	0.45	8.22	0.98
		with DMSO	0	0.40	8.30	0.99
	2	without DMSO	0	0.31	7.37	0.99
		with DMSO	0	0.27	$7.83*^{a}$	0.99

"Fecal cultures consisted of volunteer 1 and 2 fecal suspensions in fermentation medium with 30 μ M ellagic acid (EA) dissolved or not with 1% DMSO. Significant differences in growth without and with DMSO addition. * ($P \le 0.05$), ** ($P \le 0.05$).

457 However, this is the first time that urolithin M-5, urolithin M-6, 458 urolithin M-7, urolithin C, and the new identified metabolite 459 urolithin E are reported to be produced by human gut 460 microbiota. Urolithin M-6 (tetrahydroxy-urolithin) was de-461 tected with a maximum concentration between those of 462 urolithin M-5 and two trihydroxy-urolithins (urolithin C and urolithin M-7) as could be expected for the catabolic sequence (Figure 6). This shows that these metabolites occur in vivo in 465 human gut, although they can be quickly metabolized into 466 urolithin A, which is the main metabolite found in plasma and 467 urine. Analysis of human feces after the intake of ellagitannins 468 and EA should be carried out in order to evaluate the occurrence of these urolithin metabolites in the gut conditions. Degradation of EA by human gut microbiota leads to similar urolithin metabolites in vitro (fermentation medium containing EA at 30 μ M) than in vivo (after consumption of 0.6 g walnuts/ 473 kg body weight per day for 5 days). Thus, both urolithin A and isourolithin A producers in vivo (volunteers 1 and 2) led to urolithin A and isourolithin A in vitro, respectively (Figure 4A,B). Despite the fact that urolithin metabolic profiles found 477 in vivo were consistent with those found in vitro, further studies should be done in order to study the time course of the 479 production of the two microbial metabolite patterns when EA

480 containing foods are used in place of EA. This would elucidate

481 the influence of other food constituents such as fiber on

482 urolithin production rate.

In the human fecal culture of volunteer 1 (in vivo urolithin A 483 producer), EA was 62% converted to urolithins at 36 h, while in 484 volunteer 2 (in vivo isourolithin A producer) the conversion 485 efficiency to urolithins was much lower (46% at 36 h). Taking 486 into account the higher EA conversion efficiency of volunteer 1 487 and the fact that of the *C. coccoides* group achieved 1 log higher 488 concentration in fecal fermentation sample of volunteer 1 than 489 in volunteer 2, this suggests that this microbial group or other 490 in vivo co-occurring bacteria could be involved in EA 491 catabolism. In contrast, the lack of growth of *Bifidobacterium* 492 in volunteer 2 could indicate that this microbial group is not 493 involved in urolithin production.

Because of solubility problems of EA, this was alternatively 495 dissolved in DMSO and incorporated into the medium at 1% 496 v:v. In human fecal cultures where DMSO was added, different 497 EA conversion was observed (28% and 74% in the case of 498 volunteer 1 and 2, respectively). In the case of volunteer 1, the 499 EA catabolism did not reach the whole transformation to 500 urolithin A, and different intermediates were obtained. 501 Urolithin M-5 was observed as the first step in urolithin 502 production from EA to continue with urolithin M-6 and then 503 urolithin M-7 to end with urolithin A but in much lower 504 concentration than in the absence of DMSO. Furthermore, the 505 addition of DMSO prevented the conversion of urolithin M-6 506 into urolithin C being one trihydroxy-urolithin (urolithin M-7), 507 the only precursor found for the final catabolite (urolithin A) 508

Figure 6. Structures of the main urolithin metabolites detected and suggested catabolic routes of formation by gut microbiota.

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509 (Figure 6). Therefore, the use of DMSO in fecal cultures 510 offered the possibility of studying other metabolic intermediates 511 between EA and urolithin A, such as urolithin E, another 512 tetrahydroxy-urolithin precursor of urolithin M-7 (Figure 6). 513 Despite the smaller urolithin production in presence of the 514 DMSO, most of the EA was metabolized in the medium (24 515 μ M). This indicates that DMSO delayed urolithin producing 516 bacteria development, allowing the action of other bacteria able 517 to metabolize EA to simple organic acids such as phenyl acetic 518 acid and phenylpropionic acid. However, further studies should 519 be done in order to elucidate if higher abundance of 520 Bacteroidetes and less abundance of Firmicutes such as C. 521 coccoides are correlated with a lower urolithin production in 522 vivo.

In the case of human fecal culture of volunteer 2, it seems 523 524 that the catabolic route of EA to urolithins was modified by the effect of DMSO. In fact, much lower concentration of 526 isourolithin A and much higher concentration of urolithin A were detected in the presence of DMSO (Figure 4D). This suggests that the microbial strains responsible for isourolithin A production in volunteer 2 grew with difficulty under these in 530 vitro conditions. Furthermore, urolithin B was not detected, 531 which suggests isourolithin A and not urolithin A is its only 532 precursor and confirms this theory that was first proposed in 533 cattle after the administration of oak leaves. 12 In addition, this is 534 also in agreement with a previous in vitro study where urolithin B was only detected in those human fermentation cultures 536 where isourolithin A were detected. 17 A negative effect of 537 DMSO on the growth of some bacterial groups was also 538 observed, particularly on C. coccoides bacteria. A similar delay on 539 growth of this bacterial group and the time needed to achieve 540 the maximum urolithin concentration was observed, suggesting 541 that this bacterial group could be involved in the EA 542 metabolism of both volunteers.

In the literature, nothing is found about bacterial species or 543 groups responsible for urolithin production from ellagitanins. In 544 the present study, C. coccoides group (C. coccoides spp., E. rectale 545 spp., and some species of Ruminococcus genus) from the 546 Firmicutes phylum, or genera co-occurring in vivo with this 547 group, are proposed as the microorganisms involved in 548 urolithins production. Clostridium and Eubacterium genera are 549 a common element of the metabolism of several phenolic 550 compounds such as flavonoids.²¹ Given the phylogenetic 551 association between these two genera, the association of them 552 in the phenolic transformations is not surprising. The fecal 553 samples corresponding to in vivo urolithin A and isourolithin A 554 producers showed a higher abundance of either C. coccoides or 555 C. leptum than that of the Bacteroides/Prevotella group. 556 Recently, three robust clusters, referred to as "enterotypes", 557 which are not nation or continent specific have been identified. 558 Assignment of an individual microbiome into a given 559 enterotype is based upon the relative enrichment of that 560 microbiome in one of three genera: Bacteroides (enterotype 1), 561 Prevotella (enterotype 2), or Ruminococcus (enterotype 3). 22 562 Further studies should be done in order to elucidate existent 563 relations between human urolithins production capacity and 564 relative abundance of bacteria either in the different phyla or in 565 the different enterotypes.

Over the past decade, there has been enormous public and 567 scientific interest in urolithins because of their proposed health- 568 promoting effects, which, nonetheless, are far from being fully 569 understood. Future studies should take the newly identified 570 urolithin metabolites into account when investigating EA 571 absorption and metabolism as well as their physiologic effects 572 in humans. Isolation and identification of bacteria strains 573 implicated in urolithins production also require further 574 investigation.

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587 Notes

588 The authors declare no competing financial interest.

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