#### 1 Microbiome Determinants and Physiological Effects of the Benzoate-Hippurate

#### 2 Microbial-Host Co-Metabolic Pathway

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44

# 45 ABSTRACT

46

47 **Objective.** Gut microbial products are involved in type 2 diabetes, obesity and insulin
48 resistance. In particular, hippurate, a hepatic phase 2 conjugation product of microbial
49 benzoate metabolism, has been associated with a healthy phenotype. This study aims to
50 identify metagenomic determinants and test protective effects of hippurate.

51

52 Design. We profiled the urine metabolome by <sup>1</sup>H Nuclear Magnetic Resonance (NMR) 53 spectroscopy to derive associations with metagenomic sequences in 271 middle-aged 54 Danish individuals to identify dietary patterns in which urine hippurate levels were associated 55 with health benefits. We follow up with benzoate and hippurate infusion in mice to 56 demonstrate causality on clinical phenotypes.

57

58 **Results.** In-depth analysis identifies that the urine hippurate concentration is associated with 59 microbial gene richness, microbial functional redundancy as well as functional modules for 60 microbial benzoate biosynthetic pathways across several enterotypes. Through dietary 61 stratification, we identify a subset of study participants consuming a diet rich in saturated fat 62 in which urine hippurate, independently of gene richness, accounts for links with metabolic 63 health that we previously associated with gene richness. We then demonstrate causality in 64 vivo through chronic subcutaneous infusions of hippurate or benzoate (20 nmol/day) 65 resulting in improved glycemic control in mice fed a high-fat diet. Hippurate improved insulin 66 secretion through increased  $\beta$ -cell mass and reduced liver inflammation and fibrosis, 67 whereas benzoate treatment resulted in liver inflammation. 68 69 **Conclusion.** Our translational study shows that the benzoate-hippurate pathway brings a

range of metabolic improvements in the context of high-fat diets, highlighting the potential of

71 hippurate as a mediator of metabolic health.

73 74

# 75 INTRODUCTION

76

77 The human obesity epidemic raises the risk of type 2 diabetes and cardiovascular disease. 78 Dysbiosis of the gut microbiome is now recognised as a key feature of these disorders.[1] 79 The microbiome collectively encodes up to 10 million different microbial genes.[2,3] In 80 particular, gene richness has been proposed as a marker of ecological diversity mirroring 81 improvements in metabolic health.[4.5] Although the microbiome directly impacts various biological processes of the host through production or degradation of a multitude of 82 83 compounds, the vast majority of molecules involved in this chemical crosstalk remains 84 elusive.[6-9] 85 There is growing evidence that hippurate, one of the most abundant microbial-host co-86 metabolites in human urine, is positively associated with metabolic health through inverse 87 associations with blood pressure, fatty liver disease, visceral fat mass and Crohn's 88 disease.[10-14] Its microbial precursor, benzoate is taken up by organic anion transporter MCT2[15] and conjugated with glycine in the liver and kidney to form hippurate. We showed 89 90 in a genetic cross between diabetic and normoglycemic rat strains that serum benzoate 91 under host genome control.[16] Hippurate was recently shown to be associated both with 92 microbiota diversity based on 16S rRNA gene sequencing[17] and with reduced risk of 93 metabolic syndrome.[13] 94 However, there is a critical need for an in-depth characterization of the complex nutrition-95 microbiome-host interaction in the benzoate-hippurate pathway, in relation to: i) associations 96 with enterotype, gene richness and functional redundancy, ii) population stratification 97 according to nutritional patterns to identify patient sub-groups in which hippurate improves 98 metabolic health, and iii) translational elucidation of these effects on host phenotypes in vivo. 99 To address these specific points, we characterized the urinary metabolome and the fecal 100 microbiome of 271 middle-aged non-diabetic Danish individuals from the MetaHIT study.[4] 101 Through dietary stratification we delineate the complex interaction between dietary intake, 102 microbiome and metabolome its impact on body weight, immune and metabolic markers, 103 which we further confirm and characterise in a mouse model of diet-induced obesity and 104 diabetes.

105

# 106 METHODS

### 107

## 108 Human subjects

- 109 All analyses were done on non-diabetic Danish individuals from the MetaHIT study
- 110 (n=271),[4,18] including the subset of 193 individuals who completed a validated food
- 111 frequency questionnaire (FFQ).[19] The study was approved by the Ethical Committees of
- the Capital Region of Denmark (HC-2008-017 and H-15000306) and was in accordance with
- the principles of the Declaration of Helsinki. All individuals gave written informed consent
- before participating in the study. Sampling and clinical phenotyping were performed as
- described previously.[4,18] In short, all study participants were recruited from the population-
- based Inter99 study.[20]. The study program consisted of two visits with approximately 14
- 117 days apart. At the first visit all participants were examined in the morning after an overnight
- fast. Blood sampling was performed at the fasting state, and urine was collected upon arrival
- at the centre. At the second visit, a Dual Energy X-Ray Absorptiometry (DXA) scan was
- 120 performed. Serum glycine levels were previously assessed.[19] Estimated glomerular
- 121 filtration rate (eGFR) was calculated with the CKD-EPI formula with age, gender and
- 122 creatinine (µmol/L) and without ethnicity factor.[21]

# 123 Dietary assessment

- 124 A subset of the study participants (n=193) completed a validated FFQ in order to obtain
- information on their habitual dietary habits.[22] The FFQ gathered dietary information from
- all meals during a day and the intake frequencies within the past months were recorded. The
- dietary data were evaluated by determining the consumed quantity and multiplying the
- 128 portion size by the corresponding consumption frequency as reported in the FFQ. Standard
- 129 portion sizes for women and men, separately, were used in this calculation; all food items in
- the FFQ were linked to food items in the Danish Food Composition Databank as previously
- described.[19] Estimation of daily intake of macro-and micronutrients for each participant
- 132 was based on calculations in the software program FoodCalc version1.3
- 133 (http://www.ibt.ku.dk/jesper/FoodCalc/Default.htm).

# 134 Sample collection

- 135 Fecal sample collection and analysys was performed as previously described.[4] Urine was
- 136 collected at the first visit upon arrival at the study site and stored at -80°C until analysis.
- 137 Metabolic profiling
- 138 Urine samples were randomized, prepared and measured on a NMR spectrometer (Bruker)
- 139 operating at 600.22 MHz <sup>1</sup>H frequency using previously published experimental
- 140 parameters[23] The <sup>1</sup>H NMR spectra were then pre-processed and analyzed as previously

- 141 reported[10] using Statistical Recoupling of Variables-algorithm.[24] Structural assignment
- 142 was performed as reviewed in [25], using in-house and publicly available databases.

## 143 Metagenomics

- 144 Shotgun sequencing of microbial DNA and metagenomics processing workflow for gene
- richness was performed as previously published.[4] Sequences were mapped onto the
- 146 previously released integrated gene catalog.[2] Following the previously published
- strategy,[26] we built a novel set of manually curated gut-specific metabolic modules
- 148 (GMMs) to describe and map microbial phenylpropanoid metabolism from shotgun
- 149 metagenomic data. The set of 20 modules, following KEGG syntax, is provided in
- 150 supplement, including references to the original publications where pathways were
- 151 discovered and described (*Supplementary List*).

# 152 Univariate statistical analysis

- 153 A ROUT test was performed to identify potential outliers (Q=1%). For statistical comparisons
- between study groups, normality was tested using D'Agostino-Pearson omnibus normality
- 155 test, then one-way ANOVA was used, followed by Tukey's HSD post hoc testing when data
- 156 were normally distributed, otherwise groups were compared using the two-tailed Mann-
- 157 Whitney test. Data are displayed as mean ± s.e.m in all figures. Multiple testing corrections
- 158 were performed using Storey's procedure.[27]

# 159 Multivariate statistics

- 160 Probabilistic principal component analysis (PCA) was performed using MATLAB R2014a
- 161 function 'ppca' to handle missing values. Unconstrained ordination was performed using
- 162 principal coordinates analysis (PCoA) to visualize inter-individual variation in microbiota
- 163 composition using Bray-Curtis dissimilarity on the genus-level abundance matrix using the R
- 164 package *vegan*.[28] Distance based Redundancy Analysis (dbRDA) was used to determine
- the cumulative contributions of a matrix of explanatory variables on the response data,
- 166 hippurate excretion inter-individual variation (Euclidian distance on log-transformed urine
- 167 hippurate concentration matrix) in R package *vegan*.[28]. Orthogonal partial least squares
- discriminant analysis (O-PLS-DA) was performed in MATLAB R2014a for supervised
- 169 multivariate analysis as previously described.[29] The predictive capability of O-PLS-DA
- 170 models was evaluated through 7-fold cross-validation[29] to compute Q<sup>2</sup>Yhat goodness-of-
- 171 prediction parameters. The empirical significance of the  $Q^2_{Yhat}$  parameter was evaluated by
- 172 random permutation testing (10,000 iterations).[30]

# 173 Animal experiments

- 174 Six-week-old C57BL/6J male mice (Janvier Labs, Courtaboeuf, France) were maintained in
- 175 specific pathogen free condition on a 12h light/dark cycle and fed a standard chow diet

- 176 (R04-40, Safe, Augy, France) for a week. Groups of 10 randomly selected mice were then
- 177 fed either control chow diet (CHD) (D 12450Ki, Research diets, NJ) or high fat (60% fat and
- 178 sucrose) diet (HFD) (D12492i, Research diets, NJ). One week later, osmotic minipumps
- 179 (Alzet® model 2006, Charles River Lab France, l'Arbresle, France) filled with a solution of
- 180 either hippuric acid or benzoic acid (5.55mM in 0.9% NaCl) (Sigma Aldrich, St Quentin,
- 181 France) were inserted subcutaneously under isoflurane anesthesia. The metabolites were
- delivered at a rate of 0.15  $\mu$ L/hour over a 42-day period (20 nmol/day). 182
- 183 Glycemia and body weight were recorded weekly. After 3 weeks of metabolite treatment,
- 184 mice underwent an intra-peritoneal glucose tolerance test (IPGTT, 2g/kg). Blood was
- collected from the tail vein before glucose injection and 15, 30, 60 and 120 minutes 185
- 186 afterwards to determine glycemia using an Accu-Check® Performa (Roche Diagnostics,
- 187 Meylan, France). Cumulative glycemia (AUC) was calculated as the sum of plasma glucose
- 188 values during the IPGTT and cumulative glucose increase ( $\Delta G$ ) parameter was calculated
- 189 as AUC above the fasting glycemia baseline integrated over the 120 min of the IPGTT.
- 190 Insulinemia was determined at baseline and at 30 minutes using insulin ELISA kits
- 191 (Mercodia, Uppsala, Sweden). After 6 weeks of metabolite infusion, mice were killed by
- 192 decapitation and organs were dissected and weighed. Triglycerides were quantified using a
- 193 colorimetric assay (ab65336, Abcam, Paris, France) of liver homogenates. All procedures
- 194 were authorized following review by the institutional ethic committee and carried out under
- 195 national licence condition (Ref 00486.02).

#### 196 Histology and immunohistochemistry of animal tissues

- 197 Liver fibrogenesis and immunohistochemistry were determined as previously described [31].
- An epitope-specific antibody (C27C9) was used for immunohistochemistry detection of 198
- 199 insulin on pancreas sections (8508S Ozyme, Saint Quentin en Yvelines, France).

#### 200 **RNA** isolation and quantitative **RT-PCR**

- 201 Liver RNA preparation and reverse transcription were performed as previously reported [31].
- 202 Quantitative RT-PCRs were performed using oligonucleotides designed for the genes Col1
- 203 (Forward: CACCCCAGCGAAGAACTCATA; Reverse:
- 204 GCCACCATTGATAGTCTCTCCTAAC) and Col3 (Forward: GCACAGCAGTCCAACGTAGA;
- 205 Reverse: TCTCCAAATGGGATCTCTGG) and using the cyclophilin A housekeeping
- 206 gene.[31]

### 208 **RESULTS**

209

# Hippurate is the urine metabolite most strongly associated with fecal microbial generichness.

212 To identify microbial and host compounds mediating beneficial effects in metabolic health, 213 we profiled the urinary metabolome of the MetaHIT population[4] using <sup>1</sup>H nuclear magnetic 214 resonance (NMR) spectroscopy to perform a Metabolome-Wide Association Study 215 (MWAS)[11] for microbial gene richness, a proposed criterion of metabolic and immune 216 health[4] (Figure 1). We first built an orthogonal partial least squares (O-PLS) regression 217 model based on the <sup>1</sup>H NMR spectra to stratify the population by gene richness quartiles 218 computed using our previously published 10-million integrated gene catalog (IGC)[2] (Figure 219 1A, P=5.8x10<sup>-21</sup>). The cross-validated model significantly predicted variance associated with 220 gene richness through a permutation test (Figure 1B, P=9.7x10<sup>-5</sup>, 10.000 randomizations). 221 Model coefficients for this discrimination revealed hippurate as having the strongest 222 association with microbial gene count (Figure 1C): individuals with low microbial richness 223 present significantly lower urinary hippurate levels than individuals with high microbial 224 richness (Figure 1D, P=1.99x10<sup>-9</sup>, r<sup>2</sup>=0.173). These data support reports of association 225 between hippurate levels and microbial functional redundancy[26] and Shannon's diversity 226 index[17] (Figure1E, P=0.024 and Supplementary Figure 1A, P=0.0058).

227

228 Consistent with associations previously reported for microbial gene richness in the MetaHIT 229 population[4] and associations between hippurate and reduced cardiometabolic disease risk[11,12,14,17], urinary hippurate significantly correlated with low values for body mass 230 231 index (BMI), bodyweight, the homeostasis model assessment of insulin resistance (HOMA-232 IR) and fasting circulating levels of IL6, insulin, and C-peptide, whilst adjusting for age and 233 gender (partial Spearman's rank-based correlations, g<0.1, Supplementary Figure 1B). 234 Moreover, stratification on urinary hippurate concentrations in lean (BMI <25 kg/m<sup>2</sup>), and 235 overweight or obese (BMI >25 kg/m<sup>2</sup>) individuals showed improved glucose homeostasis in 236 participants excreting higher levels of hippurate (Supplementary Figure 1C, median 237 threshold). These observations depict hippurate, one of the main microbial-mammalian co-238 metabolites found in human urine, as a key molecular marker associated with gene richness 239 which may underlie some of its health benefits. These results however raise questions 240 related to the entanglement of gene richness and hippurate as potential markers of 241 health[13,17]. Adjusting for hippurate weakens associations between gene richness and 242 bioclinical variables (Supplementary Figure 1B), consistent with the idea that hippurate could

243 mediate some of the observed benefits for subjects with higher gene richness. However, 244 hippurate associations with bioclinical variables adjusted for gene richness are no longer 245 significant, suggesting that the gene richness signal overrides hippurate associations in the presence of confounding variation affecting urinary concentrations, such as diet, microbial 246 247 synthesis, host conjugation and clearance. Hippurate did not correlate either to glycine 248 bioavailability, which is required for hippurate synthesis through conjugation with gut 249 microbial benzoate[32] or kidney function (eGFR) which could limit hippurate synthesis and 250 clearance (Supplementary Figure 1D-E).

251

254

# 252 Microbiome determinants of hippurate production in the phenylpropanoid pathway

253 To characterize the microbial determinants of benzoate production, we next focussed on

annotated functions of the IGC to KEGG Orthology (KO) groups and found 2,733 KEGG and

high-throughput shotgun sequencing fecal metagenomics data (n=271). We functionally

- 256 6,931 EggNOG modules positively associated with urine hippurate levels (pFDR<0.05,
- 257 Supplementary Tables S1-2). Of specifically curated enzymatic modules[26] involved in

258 microbial benzoate metabolism (4 aerobic and 15 anaerobic; Supplementary List), only three

- 259 modules significantly correlated with urine hippurate levels: MC0004 (detected in 271
- samples) corresponding to a 2-enoate reductase converting cinnamic acid into 3-(3-

261 hydroxyphenyl)-propionic acid, MC0005 (observed in 201 samples) converting cinnamate

into benzoate and MC0014, a benzoate 4-monooxygenase only observed in fewer than 15%

of individuals (Figure 2A, Supplementary Table S3). Abundance of these modules also

correlated with gene richness (Supplementary Table S4), thereby providing a functional

265 basis for the association between gene richness and urine hippurate levels observed in this

population (Figure 1). Genes involved in MC0004 and MC0005 were predominantly found in

267 genomes from unclassified Firmicutes and Clostridiales (Figure 2B, Supplementary Tables

268 S5-6). Among classified Firmicutes, the genera *Lachnoclostridium*, *Eubacterium* and *Blautia* 

harbored MC0004. Conversely, MC0005 was encoded by Proteobacteria, i.e. *Klebsiella*,

270 Enterobacter, Suterella and Comamonas (Figure 2B). We then mapped these modules into

the enteroscape (as observed on the principal coordinates plot derived from normalized

272 genus abundances using Bray-Curtis distances,[26] Figure 2C), revealing that the

273 conversion of cinnamic acid into 3-hydroxy-3-phenylpropionic acid is linked to the

- 274 *Ruminococcus* enterotype, while capacity to convert cinnamate to benzoate is more
- 275 ubiquitously distributed across gut community types. Phenylpropanoid pathway potential is

276 increased in the *Ruminococcaceae*-enterotype compared to the *Bacteroides*- or *Prevotella*-

277 enterotypes, the former being confirmed by analyzing gradients of key taxa instead of

enterotypes (Supplementary Table S7). The results altogether suggest a wide range of
substrates, taxa and species are involved in benzoate production, consistent with its
association with gene richness.

281

# Urine hippurate levels associate with improved metabolic health in patients with diets rich in meat and saturated fats

284 We next assessed individual nutritional intake through validated FFQs available in 193 285 MetaHIT individuals.[19] Associations with metabolic health were stratified according to 286 multivariate dietary patterns (Figure 3). A PCA of 133 dietary intake descriptors summarises 287 dietary patterns and loadings highlight four archetypal diets within our population: higher 288 consumption of fruits and vegetables (n=96) vs. high consumption of meat containing 289 saturated fats (n=97) on the first principal component (PC1) and carbohydrate-rich foods vs. 290 fish containing unsaturated fats on PC2 (Figure 3A), a trend which was observed at the food 291 ingredient and nutrient level (Supplementary Figure 2A-B). It is therefore possible to stratify 292 the population according to the median of dietary PC1 highlighting contrasts between 293 healthy (higher consumption of fruit and vegetables) and at-risk (higher consumption of 294 carbohydrates and meat) diets. Although hippurate was not correlated with the first two 295 dietary principal components, its dynamic range was similar whilst stratifiying according to 296 the first two principal components (Supplementary Figure 2C-E). To summarise the main 297 factors influencing inter-individual variation in urine hippurate excretion, we calculated the 298 cumulative contribution of several covariates using a dbRDA ordination approach (Figure 299 3B). Gene richness accounted for 12% (P=0.002), followed by MC0020 encoding a 300 hippurate dehydrolase (4%, P=0.002, observed in 271 subjects) catalysing the 301 retroconversion of hippurate into benzoate, and HOMA-IR (1.5%, P=0.008; n=265). When 302 taking diet into account (i.e., PC1 fruits and vegetables vs. meat; n=193) in the dbRDA, gene 303 richness contributes to 15% (P=0.002), diet adding another 4% (P=0.002) and hippurate 304 retroconversion 3% (P=0.004), suggesting that the pattern of hippurate associations could 305 be diet-dependent and requiring further analysis. We therefore stratified the data according 306 to diet (n=193) using a median threshold for the first dietary principal component, 307 highlighting a healthy diet associated with vegetables and fruit intake (low PC1 values, n=96) 308 and an at-risk diet rich in saturated fats derived from meat intake (n=97). For this latter 309 subset of individuals consuming a diet rich in saturated fats on the first dietary principal 310 component, urine hippurate levels significantly associated with decreased HOMA-IR, 311 circulating fasting levels of insulin, fasting associated adipocyte factor (FIAF, also known as 312 Angiopoietin-like 4, a peroxisome proliferator-activated receptor target gene environmentally

313 modulated by the gut microbiota inhibiting lipoprotein lipase in adipose tissue [33]) and TNF-

 $\alpha$ , whilst displaying increased plasma levels of fasting adiponectin (Figure 3C-E, 314

Supplementary Table S8). Urine hippurate was not associated with any health benefits in the 315

- 316 subsets of participants consuming mostly a fruit and vegetable diet, a pescetarian diet or a
- 317 carbohydrate-rich diet (Supplementary Table S8).
- 318

319 To disentangle contributions arising from hippurate and gene richness to bioclinical variables 320 in subjects consuming a diet rich in fats, we compared unadjusted and adjusted Spearman's 321 rank-based correlations (Figure 3H). In the population consuming higher amounts of meat 322 and saturated fats, elevated urine hippurate levels significantly associated with an increase 323 in fasting plasma adiponectin and a reduction in adiposity, BMI, HOMA-IR and fasting 324 plasma insulin, which is consistent with gene richness being significantly associated with an 325 increase in adiponectin and a decrease in HOMA-IR and fasting plasma insulin. However, 326 the associations between gene richness and bioclinical variables were no longer significant 327 when adjusting for urine hippurate levels. Conversely, hippurate associations with insulin 328 and HOMA-IR were still significant after gene richness adjustment. We exemplified this 329 through a correlation graph taking into account the correlation between hippurate and gene 330 richness (r=0.44): this unadjusted correlation between gene richness and HOMA-IR 331 collapses when adjusting for gene richness (rho=0.143, n.s.) and is in fact contributed for by 332 a partial correlation between urine hippurate and HOMA-IR (Figure 3I). The latter finding 333 suggests a possible preventive role for hippurate in metabolic disease driven by diets high in

334 meat and saturated fats, independently of gene richness.

335

#### Hippurate and benzoate improve glucose tolerance in HFD-fed mice 336

337 To further study the impact of benzoate and hippurate on host physiology, we treated mice

338 with subcutaneous infusion of hippurate (0.14 mg/kg/day) and benzoate (0.1 mg/kg/day) in

339 CHD and HFD (Figure 4). Under control diet, metabolite treatments had no effect on body

340 weight, BMI and fasting glycemia (Supplementary Figure 4). Benzoate caused a significant

- 341 elevation of the adiposity index and a reduction of the normalised heart weight
- 342 (Supplementary Figure 5A). During an IPGTT, both metabolites induced a slight increase in
- 343 glycemia (Figure 4A-B) and  $\Delta G$  parameter (Figure 4C), respectively. Also, hippurate induced
- 344 a stronger glucose-stimulated insulin secretion than benzoate, compared to the saline-
- 345 treated mice (Figure 4D). Whilst HFD-feeding increased body weight and adiposity,
- 346 hyperglycemia and glucose intolerance (Figure 4E-H, Supplementary Figs 4D, E and 5),
- 347 mice treated with hippurate or benzoate showed a parallel improvement in glucose tolerance

348 compared to saline-treated controls (Figure 4E). This effect was illustrated by the highly 349 significant reduction of both the cumulative glycemia during the test (Hippurate vs. control -350 23.90% P=0,001, benzoate vs. ctl -31.52%, P=0.001) and the  $\Delta G$  parameter (Hippurate vs. ctrl -37.22% P=0.001, benzoate vs ctrl -33.35%, P=0.001) (Figure 4F,G). Hippurate and 351 352 benzoate treatments also significantly increased glucose-induced insulin secretion (Figure 353 4H). These data suggest that both metabolites have the capacity to improve glucose 354 tolerance and stimulate glucose-induced insulin secretion in vivo specifically in diet-induced 355 obesity and diabetes.

356

# 357 Effects of hippurate and benzoate on the morphology of pancreatic islets

358 To investigate the possible cause of stimulated insulin secretion by hippurate and benzoate,

- 359 we performed out a histological analysis of pancreatic islet structure. We confirmed that
- insulin positive area was increased by hippurate (+194%, P=0.04, one-tailed) or benzoate
- 361 (+437%, p=0.02, one-tailed) respectively in mice fed control diet (Figure 5); hippurate
- 362 treatment Insulin positive area was also increased in HFD-fed mice treated with hippurate
- 363 (+168%, P=0.04, one-tailed). These data suggest that increased  $\beta$ -cell mass may explain
- 364 enhanced insulin production and secretion induced by hippurate and benzoate treatment.
- 365

### 366 Effects of hippurate and benzoate on liver histopathology and function

Liver triglyceride accumulation, fibrosis and inflammation are hallmarks of structural and
 biochemical adaptations to HFD feeding. Liver triglycerides were more elevated in HFD fed

- 369 mice (29.30±4.15 mg/g) than in mice fed control diet (8.63±2.19 mg/g, P=0.002)
- 370 (Supplementary Figure 6). Hepatic triglycerides were not significantly affected by hippurate
- 371 or benzoate treatment in either diets. Benzoate induced a singifnicant decrease in liver
- triglycerides compared to hippurate in HFD (-57.35%, P= 0.02) (Supplementary Figure 6).
- 373 We next analysed hepatic fibrosis through quantitative analysis of collagen detected by red
- 374 picrosirius staining of histological sections (Figure 6A). Hippurate treatment resulted in a
- 375 marked reduction of liver collagen in mice fed control diet (-53.2%) or HFD (-55.7%),
- 376 whereas benzoate had no effects on collagen levels in these mice (Figure 6B,C). These
- 377 results were confirmed by liver expression of the genes encoding collagen 1 (*Col1*) and
- 378 collagen 3 (*Col3*) (Figure 6B,C): hippurate treatment induced a significant reduction of the
- 379 expression of *Col1* (-79.89%, P=0.02) and *Col3* (-29.37%, P=0.01) under control diet (Figure
- 6B). but the effect of the metaolites on *Col1* and *Col3* I HFD was marginal (Figure 6C).
- 381 Finally, we assessed the effects of hippurate and benzoate on liver inflammation through  $\alpha$ -
- 382 SMA (alpha Smooth Muscle Actin) staining[34,35] (Figure 7A), which is increased by HFD-

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feeding (+273.86%) (Figure 7B,C). Hippurate induced a marked reduction in  $\alpha$ -SMA staining 383 384 in mice fed control diet (-82.71%) or HFD (-94.87%) (Figure 7B,C), suggesting reduced 385 presence of stellar cells and reduced liver inflammation when compared to saline-treated 386 controls. In contrast, benzoate treatment in chow diet induced a strongly significant increase 387 in stellar cells when compared to mice treated with saline (+564%, P=10-7) or hippurate (+3,741%, P=10<sup>-7</sup>), thereby indicating liver inflammation in these mice (Figure 7B). 388 389 Collectively, these data show that hippurate decreases fibrosis and inflammation regardless 390 of diet, whereas benzoate reduces triglycerides and collagen accumulation in obese mice 391 fed HFD whilst stimulating inflammation in lean mice fed control diet.

392

#### 393 DISCUSSION

394

Integrated analysis of metabolome profiling and deep metagenome sequencing data of 271 395 396 middle-aged non-diabetic Danish subjects from the MetaHIT study sample [4] identified 397 urinary hippurate as the metabolite most significantly associated with microbial gene 398 richness based on the microbial IGC.[2] This observation largely confirms previous reports 399 associating hippurate with gene richness in steatosis and bariatric surgery contexts[14,36] 400 as well as with increased gut microbial diversity as assessed by sequencing the 16S rRNA 401 gene amplicon.[17] Hippurate having previously been inversely corrected with blood 402 pressure, obesity and steatosis,[11-14] this study highlights diet-dependent relationships 403 between microbiota-host co-metabolism of benzoate and hippurate and health benefits 404 associated with gene richness. Our in-depth dissection of the metagenomic determinants of 405 urinary hippurate highlighted a series of richness-responsive gene modules functionally 406 related to benzoate synthesis. These modules are shared across several enterotypes and 407 taxonomic gradients. Population stratification analyses demonstrated that hippurate only 408 benefits individuals consuming a diet rich in saturated fats. This hypothesis of a diet-409 dependent beneficial health effect of benzoate metabolism was confirmed in vivo in a 410 preclinical model of HFD-induced obesity.

411

#### 412 Hippurate brings diet-dependent metabolic improvements in pancreas and liver

413 Our study shows that chronic hippurate treatment in a model of obesity induced by HFD-414 feeding reduces glucose intolerance, stimulates insulin secretion, enhances β-cell mass and 415 reduces hepatic inflammation and fibrosis. Metabolomic studies have consistently shown 416 inverse associations variations between hippurate levels and pathophysiological elements of the metabolic syndrome. Urinary hippurate is reduced in mouse models of insulin 417

- 418 resistance[10] and in rat models of spontaneously occurring hypertension (SHR), type 2
- 419 diabetes (Goto Kakizaki, GK) and obesity (Zucker) or the WKY rat.[37-39] This is consistent
- 420 with our previous reports showing an inverse association among hippurate, insulin
- 421 resistance, hypertension, obesity or liver steatosis[10-12,14] and observations that hippurate
- 422 exerts protective effects in  $\beta$ -cells.[40] We also showed in HFD-fed isogenic mice that
- 423 urinary hippurate measured before a 3-week HFD induction predicts the future development
- 424 of obesity, suggesting that functional variations in microbiome predicts disease risk
- independently of genetics.[41] Whilst both hippurate and benzoate have similar *in vivo*
- 426 effects, including greatly improved glucose tolerance and stimulation of insulin secretion,
- 427 only hippurate results in beneficial effects on increased  $\beta$ -cell mass or reduced liver fibrosis.
- 428

# 429 The phenylpropanoid-benzoate-hippurate pathway in metabolic health

430 The range of responses observed in the animal model treated with hippurate and benzoate 431 depict subtle and context-dependent microbiome-host interactions. Benzoate and its co-432 metabolite hippurate are the endproducts of several converging microbial biosynthetic 433 pathways.[15] The phenylpropanoid pathway is a broad network of reactions connecting a 434 wide range of dietary substrates such as phenylalanine, guinic acid, shikimic acid or 435 chlorogenic acid for instance to 4-coumaryl-coA. These pathways lead to much simpler 436 molecules, benzoate being their common endpoint. Dietary and microbial intermediates 437 (including cinnamic acids, coumarins, stillbenoids, flavonoids and isoflavonoids) in the 438 phenylpropanoid and connected pathways are associated with beneficial health

- 439 outcomes.[15,42]
- 440

441 **Conclusion.** Our study depicts hippurate as a pivotal microbial-host co-metabolite mediating 442 part of the beneficial metabolic improvements associated with high microbial gene richness 443 in the context of Western-style diets. This work unifies previous reports in which hippurate was associated improvements in insulin resistance, steatosis, hypertension and obesity[10-444 445 12,14] and microbial ecological diversity.[14,17] Beyond the diversity of microbial 446 ecosystems and functions associated with hippurate, we uncover beneficial bioactivities in 447 the liver and pancreas resulting in health benefits in terms of metabolic control, liver 448 inflammation and fibrosis. Our observations support the existence of diet-dependent 449 microbial-host metabolic axis in which hippurate partly offsets unhealthy diets, further 450 exemplifying that the microbiome determines key components of human biochemical 451 individuality[43] and provides critical diagnostic and therapeutic potential in personalized 452 nutrition and stratified medicine.[44,45]

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- 462

# 463 Contributions

- 464 F.B. J.C., T.N. and S.V.S. and contributed equally to this work. F.B. J.C. and G.I.M. acquired
- 465 data, J.C., T.N., S.V.S. performed analyses, G.F. and S.V.S. curated the gut-specific
- 466 metabolic modules, L.H., A.L.N., A.R.M., N.P., S.F., E.L.C., and A.M.L.L. participated in data
- 467 collection and processing, J.C., T.N., S.V.S., F.B. and G.F. performed statistical analyses.
- 468 M.-E.D., D.G., J.R. and O.P. designed the study. T.H., J.K.N. K.C., P.B., S.D.E., participated
- in the study design and interpretation of the results, M.E.D. wrote the manuscript with
- 470 contributions from F.B., J.C., T.N., S.V.S, G.F., J.R., O.P and D.G.
- 471

## 472 Competing interests

473 The authors declare no competing financial interests.

#### 474 **Figure Legends**



# 475

476 Figure 1. Hippurate is the main metabolite correlated with gene richness and 477 functional redundancy of the microbiome. (A) Scores plot (predictive component 1) 478 obtained for an O-PLS-DA model fitted usingurinary<sup>1</sup>H NMR-spectra to predict microbial 479 gene richness, showing a significant association between gene richness quartiles and <sup>1</sup>H NMR spectra (p=5.84x10<sup>-21</sup> for a significantly non-zero slope using F-test, n=271). (B) 480 Empirical assessment of the significance of O-PLS goodness-of-fit parameter Q2<sub>Yhat</sub> by 481 generating a null distribution with 10,000 random permutations (p=9.68x10<sup>-5</sup>). (C) Manhattan 482 483 plot highlighting associations between <sup>1</sup>H NMR variables and gene count displayed in a 484 pseudo-spectrum layout. A negative value (blue circles) means a negative correlation while 485 a positive value (red circles) means a positive correlation. Grey circles are clusters with a pvalue>0.01. Size of circles represents the covariance of the cluster with the gene count.( D) 486 487 Association between urinary hippurate intensity and gene count quartiles (p=1.99x10<sup>-9</sup> for a 488 significantly non-zero slope using F-test). (E) Association between urinary hippurate intensity 489 and microbial functional redundancy [26] quartiles (p=0.0239 for a significantly non-zero 490 slope using F-test, n=271).





492 Figure 2. Detection of microbial phenylpropanoid metabolism-related modules in fecal 493 metagenomes of healthy volunteers and their associations with urine hippurate

494 concentrations. (A) Visualisation of gut-specific metabolic modules (GMMs) encoding

495 anaerobic phenylpropanoid metabolism-related pathways detected in more than 20% of

- individuals; MC0004 (orange; Spearman rho=0.19, q-value=0.005) and MC0005 (blue;
  Spearman rho=0.21, q-value=0.005) correlate positively to urine hippurate concentrations
  (n=271). All metabolites are connected to benzoate but for clarity the non-significant
  reactions were omitted. (B) Metagenomic species encoding modules MC0004 and MC0005.
  (C) [top panel] Fecal microbiomes dissimilarity visualised on the first plane of the genus-level
  principal coordinates analysis (PCoA, Bray-Curtis dissimilarity), with individual samples
- 502 colored according to enterotypes (R, Ruminococcaceae; B, Bacteroides; P, Prevotella).
- 503 [middle and bottom panels] Same genus-level PCoA overlaid with a mesh colored according
- to the median abundances of GMMs MC0004 (red) and MC0005 (blue) in samples falling
- 505 within each cell of the mesh (n=271). See Supplementary Table S3 for correlation between
- 506 hippurate and GMMs.



507

508 Figure 3. Hippurate associates with improved glucose homeostasis only in 509 participants consuming a diet rich in saturated fats. (A) Biplot of the principal

510 component analysis (PCA) of dietary intakes highlights opposite diets along first two

- 511 principal components (PCs). The main drivers of each principal components are named and
- 512 represented by blue arrows. (B) Cumulative contributions of explanatory variables to inter-
- individual variation in hippurate excretion, estimated by redundancy analysis (dbRDA). 513
- 514 Explanatory variables included microbiota gene count, microbiota phenylpropanoid
- 515 metabolism modules, host dietary principal components and clinical parameters (age,
- 516 gender, BMI, HOMA-IR, CRP, serum glycine levels, and glomerular filtration rate (eGFR)
- 517 estimation with CKD-EPI). (C-F) Evaluation of hippurate stratification (high hippurate, n=49

- 518 vs low hippurate, n=48) on bioclinical variables (q<0.1, Supplementary Table S8) for
- 519 individuals on high PC1 (i.e. high meat / high saturated fat diet). For full name description of
- 520 physiological data see Supplementary Table S8.

522

523





526 (A) Heatmap summarising Spearman's partial correlation between gene richness.

527 hippurate,gene richness adjusted for hippurateand hippurate adjusted for gene richness and bioclinical variables, all correlations adjusted for age and gender. Stars represent significant 528 529 pFDR corrected using Benjamini Hochberg procedure \* pFDR<0.1, \*\* pFDR<0.05, \*\*\* 530 p<0.01. (B) Representation of the Spearman's correlations and partial correlations between 531 gene count and hippurate, hippurate and HOMA-IR adjusted for gene richness and between 532 gene richness and HOMA-IR adjusted for hippurate. (C) Plasma glucose during a glucose 533 tolerance test (GTT). (D) Area under the curve for glucose during the GTT. (E) Body weight 534 of mice during the 6 weeks of hippurate treatment. (F) Body mass index at sacrifice (G) 535 Adipose tissue weight normalised to body weight. (H) Plasma adiponectin concentration. For 536 chow diet and chow diet + hippurate, (n=10) and for HFD and HFD + hippurate (n=6). Data 537 shown are mean±SEM. Statistical analysis was performed using two-way ANOVA with 538 Tukey's posthoc test. \*\* p<0.01, \*\*\*\* p<0.0001. For panel (D), (F), (G) and (H), groups with 539 different superscript letters are significantly different (P<0.05), according to Tukey's posthoc 540 test.

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- 542



543

Figure 5. Effect of chronic administration of hippurate and benzoate on pancreatic 544

545 islets in C57BL6/J mice. The effect of chronic subcutaneous administration of the 546 metabolites (5.55 mM) for 42 days on islet density was tested in mice fed chow diet (CHD)

547 or high fat diet (HFD) for 56 days. Control mice were treated with saline. Each biological

548 replicate represents one slide per animal mounted with at least 3 tissue sections,

549 representing 3 technical replicates, the mean and variance of which is presented as the

550 result per biological replicate. Results are expressed as percentage of positive pixels.

551 ‡P<0.05, significantly different between mice treated with benzoate and hippurate.



552

553 Figure 6. Effect of chronic administration of hippurate and benzoate on liver fibrosis in C57BL6/J mice. (A) The effect of chronic subcutaneous administration of the metabolites 554 555 (5.55 mM) for 42 days on liver collagen was tested in mice fed control chow diet (CHD) or 556 high fat diet (HFD) for 56 days. Control mice were treated with saline. Red Picrosirius 557 staining of histological sections was used to visualise and guantify fibrosis in mice fed CHD 558 (B) or HFD (C). Each biological replicate represents one slide per animal mounted with at 559 least 3 tissue sections, representing 3 technical replicates, the mean and variance of which 560 is presented as the result per biological replicate (B,C). Liver expression of the genes 561 encoding collagen 1 (Col1) and collagen 3 (Col3) was assessed in mice fed CHD (B) or HFD 562 (C) by quantitative RT PCR in 6 mice per group. Data were analyzed using the unpaired 563 Mann-Whitney test. Results are means ± SEM. \*P<0.05 significantly different between mice 564 treated with hippurate and controls. ‡P<0.05, significantly different between mice treated 565 with benzoate and hippurate.





200	0.00
567	Figure 7. Effect of chronic administration of hippurate and benzoate on liver
568	inflammation in C57BL6/J mice. $\alpha$ SMA staining of liver slides was used to assess
569	inflammation in mice fed chow diet (CHD) or high fat diet (HFD) for 56 days and chronically
570	treated with subcutaneous administration of the metabolites (5.55mM) for 42 days (A).
571	Control mice were treated with saline. Each biological replicate represents one slide per
572	animal mounted with at least 3 tissue sections, representing 3 technical replicates, the mean
573	and variance of which is presented as the result per biological replicate in mice fed CHD (B)
574	or HFD (C). Data were analyzed using the unpaired Mann-Whitney test. Results are means
575	± SEM.
576	++++P<0.0001, significantly different between mice treated with benzoate and saline treated
577	controls. ####P<0.0001, significantly different between mice treated with benzoate and
578	hippurate.
579	

#### 580 SUPPLEMENTARY SECTION





# 582

#### 583 Supplementary Figure 1. Relationship between gene richness, hippurate and

#### 584 bioclinical variables

- (A) Association between urinary hippurate intensity and Shannon's diversity index quartiles 585
- 586 (p=6.04x10-8 for a significantly non-zero slope using F-test, n=271). (B) Heatmap
- summarising significant (pFDR<0.1) Spearman's correlation FDR corrected using Storey's 587
- 588 procedure[27] betweengene count, hippurate and gene count adjusted for hippurate and
- physiological data, all adjusted for age and gender. For full physiological data names and 589
- 590 units see Supplementary Table 8. (C) Association between hippurate and insulin resistance
- 591 (HOMA-IR) stratified according to BMI (lean (BMI < 25, n=87), overweight and obese (BMI > 25
- 592 ,n=184) and hippurate excretion levels (Mann-Whitney U test, p=0.0058). (D)
- 593 Representation of the absence of significant correlation between urinary hippurate and
- 594 eGFR. (E) Representation of the absence of significant correlation between urinary
- 595 hippurate and circulating glycine.



596

#### 597 Supplementary Figure 2. Urinary hippurate does not correlate with main dietary

#### 598 trends summarized and presents a high variability within each subgroup

- 599 (A) Biplot of the dietary data generated using only food items. (B) Biplot of thmicrobial e
- 600 dietary data generated using only macronutrients items. (C) Representation of the absence
- 601 of correlation between hippurate concentration and principal component 1. (D)
- 602 Representation of the absence of correlation between hippurate concentration and principal
- 603 component 2. (E) Distribution of hippurate urinary concentration within each
- 604 individualsubgroup stratified in high and low hippurate using median.



605

#### 606 Supplementary Figure 3. Experimental design for chronic six-week administration of

- benzoate and hippurate 607
- 608 Experiment design showing groups and durations of each step for the chronic treatments
- 609 with benzoate and hippurate in mice.







Supplementary Figure 4. Effects of chronic administration of hippurate and benzoate 613

614 on body growth and fasting glycemia. C57BL6/J mice fed control chow diet (A-C) or high

fat diet (D-F). The effects of chronic subcutaneous administration of the metabolites (5.55 615

616 mM) in mice were tested on body weight (A,D), body mass index (BMI) (B,E), fasting

617 glycemia (C,F). Control mice were treated with saline. BMI was calculated as body weight

618 divided by the squared of anal-nasal length. Results are means ± SEM.





620 Supplementary Figure 5. Organ weight in C57BL6/J mice treated chronically with hippurate

621 or benzoate for 42 days. Control mice were treated with saline. Mice were fed chow diet

622 (CHD) or high fat diet (HFD) for 56 days. Data are expressed as the ratio between organ

623 weight and body weight. Data were analyzed using the unpaired Mann-Whitney test. Results

624 are means ±SEM.

625 \*\*P<0.01, significantly different between mice treated with hippurate and controls. †P< 0.05,

++P<0.01, significantly different between mice treated with benzoate and saline treated 626

- 627 controls.
- 628

629





631 Supplementary Figure 6. Effect of chronic administration of hippurate and benzoate

632 on liver triglycerides content in C57BL6/J mice. The effect of chronic subcutaneous

633 administration of the metabolites (5.55mM) for 42 days on liver triglycerides was tested in

634 mice fed control chow diet (CHD) or high fat diet (HFD) for 56 days. Assay was carried out in

635 6 mice per group. Data were analyzed using the unpaired Mann-Whitney test.

Results are means ± SEM 636

637 ‡P<0.05, significantly different between mice treated with benzoate and hippurate.

- 638
- 639

640 Supplementary Table S1 641 Association between microbiota functional potential mapped to KEGG Orthologs (KOs) 642 database and urine hippurate levels. 643 644 Supplementary Table S2 645 Association between microbiota functional potential mapped to the eggNOG database and 646 urine hippurate levels. 647 648 Supplementary Table S3 649 Association between microbiota functional potential mapped to gut-specific metabolic 650 modules (GMMs) describing phenylpropanoid metabolism and urine hippurate levels. 651 652 **Supplementary Table S4** 653 Association between abundance of gut-specific metabolic modules (GMMs) describing 654 phenylpropanoid metabolism and gene richness. 655 656 Supplementary Table S5 657 Phenylpropanoid metabolism potential in metagenomic species, assessed by mapping to 658 GMMs significantly associated to urine hippurate levels. 659 660 Supplementary Table S6 Association between the microbiota composition profiled as metagenomic OTUs (mOTUs) 661 662 and urine hippurate levels. 663 664 Supplementary Table S7 665 pFDR from Spearman's rank-based correlations between GMMs describing 666 phenylpropanoid metabolism and gene richnessbioclinical variables using Storey's FDR 667 correction. 668 669 Supplementary Table 8 670 pFDR from Mann-Whitney U test for hippurate stratification in each bioclinical variable 671 between using Storey's FDR correction. 672 673

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### Supplementary List: Module definitions

# MC0001 phenylalanine degradation (cinnamate production)

cpd [phenylalanine] [cinnamate, NH3]

### K10775

ref MetaCyc Pathway: trans-cinnamoyl-CoA biosynthesis

### MC0002 chlorogenate degradation

cpd [chlorogenate] [caffeate, quinate]

K06889 K09252

ref MetaCyc Pathway: chlorogenic acid degradation

1

### MC0003 caffeate respiration

cpd [caffeate] [hydrocaffeate]

bactNOG04579

ref<sup>2</sup>

### MC0004 cinnamate conversion

cpd [cinnamate] [3-(3-hydroxyphenyl)propanoate]

K10797

ref KEGG Pathway: phenylalanine degradation

### MC0005 coumarate degradation

cpd [coumarate] [hydroxybenzoate]

cpd [ferulate] [vanillate]

cpd [caffeate] [protocatechuate]

cpd [cinnamate] [benzoate]

K01904 bactNOG05057

K01692 K01715 K01782 K01825 K13767 K15016 bactNOG19280

### K00141

ref MetaCyc Pathway: 4-coumarate degradation (anaerobic) <sup>3</sup>

### MC0006 (hydroxy)benzoate degradation

cpd [(hydroxy)benzoate] [3-hydroxypimeloyl-CoA]

K04105,K04107,K04108,K04109 K04110 bactNOG00950

K04112,K04113,K04114,K04115 K19515,K19516

K07537

K07538

K07539

ref	MetaCyc Pathway: 4-	coumarate degradation (anaerobic)	MetaCyc Pathway: benzoyl-CoA
degrada	ation II (anaerobic)	Kegg module: M00541	

### MC0007 ferulate degradation

cpd [ferulate] [vanillin]

K12508

K18383

ref MetaCyc Pathway: ferulate degradation

### MC0008 vanillate conversion

cpd [vanillin, (O2)] [protocatechuate]

bactNOG00059

K03862,K03863 K15066

ref MetaCyc Pathway: superpathway of vanillin and vanillate degradation Kegg Pathway: aminobenzoate degradation

### MC0009 cinnamate degradation

cpd [cinnamate] [benzoyl-CoA]

bactNOG08521

actNOG07134

### COG0277

bactNOG05297

ref MetaCyc Pathway: trans-cinnamoyl-CoA biosynthesis MetaCyc Pathway: benzoyl-CoA biosynthesis

#### MC0010 quinate degradation

cpd [quinate] [protocatechuate]

cpd [shikimate] [protocatechuate]

K05358 COG0169

K03785 K03786 K13832

K09483 K15652

ref 4

MetaCyc Pathway: quinate degradation I MetaCyc Pathway: quinate degradation II

#### MC0011 4-hydroxybenzoate conversion

cpd [4-hydroxybenzoate, O2] [protocatechuate]

K00481

5 Kegg Pathway: benzoate degradation 4 ref

#### MC0012 catechin degradation

cpd [catechin] [protocatechuate, phloroglucinol carboxylic acid]

bactNOG14887

6 ref

#### MC0013 3-hydroxybenzoate conversion

cpd [3-hydroxybenzoate, O2] [protocatechuate]

K19065

ref Kegg Pathway: benzoate degradation

MC0014 benzoate conversion cpd [benzoate, O2] [4-hydroxybenzoate]

### K07824

ref Kegg Pathway: benzoate degradation

### MC0015 benzoate degradation (aerobic)

cpd [benzoate, O2] [catechol, CO2]

K05549,K05550,K05784

K05783

ref Kegg module: M00551 MetaCyc Pathway: benzoate degradation I (aerobic)

# MC0016 benzoate degradation (anaerobic)

cpd [benzoyl-CoA] [3-hydroxypimeloyl-CoA]

K04112,K04113,K04114,K04115 K19515,K19516

K19066

K07534

K07535

K07536

K04118

ref MetaCyc Pathway: benzoyl-CoA degradation III (anaerobic) Kegg module: M00540

### MC0017 phenylalanine degradation

cpd [phenylalanine, 2-oxoglutarate] [CO2, glutamate, benzoyl-CoA]

### K00832 K00812 K00813 K11358 K00817

K04103

K00146 K00129

K01912

K18361

K18355,K18356,K18357,K18358,K18359

ref MetaCyc Pathway: phenylacetate degradation II (anaerobic) MetaCyc Pathway: Lphenylalanine degradation II (anaerobic)

MC001	8 cellulose and hemicellu	lose degradation (cellulolosome)				
cpd	[cellulose, hemicellulose]	[ferulate, polysaccharide, oligosaccharide]				
K01181 K13465						
K09252						
bactNOG05428						
ref	MetaCyc Pathway: cellulose and hemicellulose degradation (cellulolosome)					
MC001	9 feruloyl esterase (cellu	lolosome)				
cpd	[cellulose, hemicellulose]	[ferulate]				
K09252	2					
ref	MetaCyc Pathway: cellulose and	d hemicellulose degradation (cellulolosome)	MC0018			

MC0020	hippurate hydrolase

cpd [hippurate] [glycine, benzoate]

K01451

ref <sup>7</sup>

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