

1 **Determinants of post-prandial plasma bile acid kinetics in human volunteers**

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32 **Abstract**

33 Bile acids (BA) are signaling molecules with a wide range of biological effects, also
34 identified amongst the most responsive plasma metabolites in the post-prandial state. We
35 here describe this response to different dietary challenges and report on key determinants
36 linked to its inter-individual variability.

37 Healthy men and women (N=72, 62 ± 8 years) were enrolled into a 12-week weight loss
38 intervention. All subjects underwent an oral glucose tolerance test (OGTT) and a mixed
39 meal tolerance test (MMTT) before and after the intervention. BA were quantified in plasma
40 by LC-MS/MS combined with whole genome exome sequencing and fecal microbiota
41 profiling.

42 Considering the average response of all 72 subjects, no effect of the successful weight loss
43 intervention was found on plasma BA profiles. Fasting and post-prandial BA profiles
44 revealed high inter-individual variability and 3 main patterns in post-prandial BA response
45 were identified using multivariate analysis. Although the women enrolled were
46 postmenopausal, gender effects in BA response were evident. Exome data revealed the
47 contribution of preselected genes to the observed inter-individual variability. In particular, a
48 variant in the *SLCO1A2* gene, encoding the small intestinal BA transporter OATP1A2 was
49 associated with delayed post-prandial BA increases. Fecal microbiota analysis did not
50 reveal evidence for a significant influence of bacterial diversity and/or composition on
51 plasma BA profiles.

52 The analysis of plasma BA profiles in response to two different dietary challenges revealed
53 a high inter-individual variability, which was mainly determined by genetics and gender of
54 host with minimal effects of the microbiota.

55

56 **New and Noteworthy:**

57 Considering the average response of all 72 subjects, no effect of the successful weight loss
58 intervention was found on plasma BA profiles.

59 Despite high inter-individual variability, 3 main patterns in post-prandial BA response were
60 identified using multivariate analysis.

61 A variant in the *SLCO1A2* gene, encoding the small intestinal BA transporter OATP1A2
62 was associated with delayed post-prandial BA increases to both the OGTT and MMTT.

63

64 **Keywords:** bile acids, *SLCO1A2*, post-prandial, OGTT, MMTT

65

66 **Introduction**

67 In addition to their role in solubilizing lipids in the intestine, bile acids (BA) are now
68 recognized as important signaling molecules. They serve as ligands of the farnesoid X
69 receptor (FXR) expressed in intestine and liver and thus control synthesis and transport of
70 BA but also modulate expression of other genes(24, 28, 39, 43, 57, 64). After the discovery
71 of a G protein-coupled receptor (TGR5) for BA(28, 39), acute metabolic effects of BA
72 received considerable interest. With the expression of TGR5 and FXR in a wide range of
73 cell types and tissues, BA effects cover immunomodulation, metabolism of adipose tissue
74 and muscle as well as gastrointestinal hormone secretion (e.g. GLP-1)(27, 28). TGR5
75 activation by BA was shown to activate deiodinase, increasing the level of active thyroid
76 hormone which in turn elevates the expression of enzymes involved in fatty acid oxidation
77 in brown adipose tissue and skeletal muscle(24, 28, 39, 43, 57, 60, 64, 66). TGR5 was also
78 identified as a target for treatment of obesity and type-2 diabetes. BA can (via farnesoid X
79 receptor (FXR)-induced activation of short heterodimer partner (SHP) and its effect on
80 hepatocyte nuclear factor 4 (HNF-4) and/or forkhead box O1 (Foxo1)) decrease the
81 expression of gluconeogenic enzymes with a beneficial effect on glucose homeostasis in
82 insulin resistance(28, 35, 38, 39). FXR also participates in the regulation of insulin synthesis
83 and secretion as well as in the protection of pancreatic islets from lipotoxicity(27, 28, 44,
84 45). Via SREBP-1c activation, BA were also shown to modulate triacylglycerol synthesis in
85 rodent liver(61).

86 Taken the relevance of BA as receptor ligands and their increase in circulation in the post-
87 prandial state, it is remarkable that impressive differences in plasma levels of BA between
88 individuals are not explained as yet. Within the NutriTech study, we recorded changes in
89 BA plasma concentrations in 72 healthy male and female (postmenopausal) volunteers in
90 response to an oral glucose tolerance (OGTT) and a mixed meal tolerance test (MMTT)
91 combined with exome sequencing and fecal microbiota analysis. We here report on a large
92 inter-individual variability of plasma BA profiles with pronounced gender effects and the
93 identification of a number of gene variants underlying hepatic BA synthesis and
94 enterohepatic recirculation.

95 **Methods**

96 **1. Study design**

97 The research project NutriTech was funded by the European Union 7 Framework program
98 (clinicaltrials.gov record: NCT01684917). It aimed at better phenotyping human volunteers
99 in response to standardized challenge tests. 72 volunteers (37 women and 35 men) in
100 average 59.2±4.2 years-old, BMI 29.7±2.7, healthy at the screening considering fasting
101 glucose and insulin concentrations and blood pressure values were recruited. The subjects
102 underwent comprehensive phenotyping, including MRI scanning, food intake recording, and
103 blood profiling for metabolites, hormones and chemokines. All subjects underwent an
104 OGTT and a MMTT before and after a 12-week period, in which 40 participants followed a
105 20% caloric restriction diet while subjects in the control group consumed an average
106 european diet which was matched for their energy expenditure to maintain body weight.
107 Challenge tests always started at 09:00 am after 12 hours of fasting. The OGTT drink
108 consisted of 75g glucose in 250 mL of water consumed within 5 minutes. Blood was
109 collected in heparin-coated tubes at times 0, 15, 30, 60, 90, 120 and 240 minutes. The
110 MMTT (also known as the “PhenFlex drink”) was a high-fat, high-glucose, high-caloric drink
111 (400 ml) consisting of 320 ml tap water, 75g glucose, 60g palm olein, 20g profitar (protein
112 supplement, Nutricia, Netherlands) and 0,5g artificial vanilla aroma resulting in a shake with
113 33, 59 and 8% of energy from carbohydrates, lipids and proteins, respectively. The drink
114 was ingested within 5 minutes and blood was collected at 0, 30, 60, 120, 240, 360 and 480
115 minutes. Plasma was separated and stored at -80°C for later analyses. Clinical chemistry
116 parameters were assayed using enzymatic colorimetric kits and GLP-1 was measured
117 using an in-house radioimmunoassay as previously described(30).

118 **2. Bile acid analysis**

119 Plasma bile acids were analyzed by a modified method originally described by Tagliacozzi
120 et al (53). Briefly, 10 µL of plasma were mixed with 10 µL internal standard solution and 500
121 µL ice-cold methanol was added for deproteinization. Samples were vortexed and
122 centrifuged at 15.000 g for 10 minutes. The supernatant was transferred to a 96 deep-well
123 plate and evaporated under a stream of N₂. The solid remnants were re-suspended in
124 methanol:water (1:1). Analysis was performed by LC-MS/MS using a triple quadrupole
125 mass spectrometer (HPLC Agilent - CA, USA; QTrap 5500 - ABSciex MA, USA). BA were
126 separated using a gradient with 0.2% formic acid in water and acetonitrile (acetonitrile going
127 from 30 % at the start of the run to 100 % at 20 minutes) with a flow rate of 0.6 ml/min and

128 a reverse phase column (Phenomenex Luna C18(2) 150 x 4,6mm; 5µm particle size), kept
129 at 40 °C. The mass spectrometer was operated in negative ion mode and mass spectra
130 were obtained using the multiple reaction-monitoring mode (MRM). Integration of the peaks
131 was done using Analyst Software (ABSciex MA, USA). Analyte concentration was
132 calculated using deuterated internal standards (d4-Deoxycholic acid, d4-
133 Glycoursodeoxycholic acid, d4-Glycodeoxycholic acid, d4-Glycocholic acid and d5-
134 Taurocholic acid). Samples were randomized so that every batch contained samples from
135 men and women, and from OGTT and MMTT performed before and after the intervention to
136 avoid batch effects. Reference fasting plasma samples (Recipe Clinical Diagnostics
137 Munich, Germany) were included in each batch and a dilution row of BA standards was
138 processed in a similar manner in every batch for quantitation purposes. In total, 13 BA were
139 quantified: ursodeoxycholic acid (UDCA), cholic acid (CA), chenodeoxycholic acid (CDCA),
140 deoxycholic acid (DCA), glycoursodeoxycholic acid (GUDCA), glycocholic acid (GCA),
141 glycochenodeoxycholic acid (GCDCA), glycodeoxycholic acid (GDCA),
142 tauroursodeoxycholic acid (TUDCA), taurocholic acid (TCA), taurochenodeoxycholic acid
143 (TCDCA), taurodeoxycholic acid (TDCA) and taurolithocholic acid (TLCA). Due to the low
144 abundance and values below the lower limit of quantitation, UDCA TUDCA and TLCA were
145 not taken into consideration in the statistical analysis. In our analyses, BA were grouped as
146 primary (sum of CA, CDCA, and their taurine and glycine conjugates) or secondary +
147 tertiary (DCA, LCA, UDCA including taurine and glycine conjugates thereof). The terms
148 taurine-conjugated, glycine-conjugated, or unconjugated BA are used regardless of whether
149 BA are primary or secondary + tertiary. In the text, total BA refers to the sum of all individual
150 BA.

151

152 **3. Exome DNA sequencing**

153 Genomic DNA was extracted from saliva, collected using the Oragene® DNA sample
154 collection kit (DNA Genotek, Ottawa, ON, Canada), following manufacturer's protocol.
155 Whole-exome sequencing libraries were prepared from DNA using SureSelectXT Human
156 All Exon V4+UTRs (71Mb) kit (Agilent Technologies, Santa Clara, CA). Sequencing was
157 performed on a HiSeq25000 platform generating 100 base-pair end reads. After quality
158 control using FastQC(29) version 0.10.0, sequencing reads were mapped to the GRCh37
159 (hg19) reference assembly of the human genome using BWA-MEM(36) version 0.7.2.(36)
160 Variant calling was performed using GATK version 2.6(41) and quality filtered variants were
161 annotated in ANNOVAR(58). Annotated exonic variants were filtered based on *in-silico*

162 predictive functional effects on the protein by two algorithms incorporated in the ANNOVAR
163 suite: i) SIFT(31), which relies on the degree of conservation of amino acid residues in
164 highly conserved regions in sequence alignments derived from closely related sequences
165 and ii) Polyphen-2(2) which predicts the possible impact of amino acid substitutions on the
166 stability and function of human proteins using structural and comparative evolutionary
167 considerations. These approaches indicate a high probability that an amino acid change is
168 “deleterious” based on a combination of these attributes. The effect of the identified variants
169 on post-prandial BA metabolism was analyzed by creating allelic gene scores, which
170 represented the sum of deleterious variants per gene involved in BA synthesis, metabolism
171 and transport. Where necessary individual variants were analyzed separately.

172

173 **4. High-throughput microbiome 16S rRNA gene amplicon sequencing**

174 Fresh stool sample were collected by subjects before and after a 12-week intervention
175 period. Samples were immediately frozen at -20°C by the subjects after defecation and
176 transported in frozen state to the laboratory at TNO where samples were mechanically
177 homogenized, split into aliquots in sterile 2 ml cryovials and stored at -80°C . For fecal
178 genomic DNA isolation, approximately 100-150 mg of fecal material was directly transferred
179 to DNA isolation plates. Phenol pH8.0 was added and the samples were mechanically
180 disrupted by bead beating with a 96-well plate Beadbeater (Biospec Products, Bartlesville).
181 Isolated DNA was extracted and purified with the AGOWA mag Mini DNA Isolation Kit
182 (AGOWA, LGC genomics, Berlin, Germany) as described in Kelder et al (2014)(29).
183 Microbiota analysis was performed by high-throughput sequencing of bar-coded amplicons
184 spanning the archaeal and bacterial V4 hypervariable region. These amplicons, generated
185 from a standardized quantity of template DNA (100pg) using adapted primers 515F and
186 806R, were bidirectionally sequenced using the MiSeq system (Illumina, San Diego, CA) as
187 described previously(11).

188 Data were analysed as described in detail previously(33). Raw reads were processed using
189 an in-house developed pipeline (www.imngs.org)(32) based on the UPARSE approach(16).
190 Sequences were trimmed to the first base with a quality score <3 and then paired. Those
191 with less than 200 and more than 300 nucleotides and paired reads with an expected error
192 >3 were excluded from the analysis. Remaining reads were trimmed by five nucleotides on
193 each end to avoid GC bias and non-random base composition. The presence of chimeras
194 was tested using UCHIME(15). Operational taxonomic units (OTUs) were clustered at 97%
195 sequence similarity, and only those with a relative abundance $>0.5\%$ in at least one sample

196 were kept. Taxonomies were assigned at 80% confidence level using the RDP
197 classifier(59).

198

199 **5. Statistical analysis**

200 Depending on the nature of comparisons, T-test, 1-way ANOVA or 2-way ANOVA with post
201 hoc multiple comparison Bonferroni's test were used. Unless otherwise stated, differences
202 with $P < 0.05$ were considered as significant. Shapiro-Wilk test was used to assess
203 normality. Results from calculation of area under the curve (AUC), refers to incremental
204 AUC, which considers individual values at $t=0$ as baseline. In cases where outliers were
205 removed, their detection was done using ROUT method.

206 The contribution of individual predictors to the overall variation of plasma BA concentrations
207 was quantified by means of percentaged marginal coefficients of determination (R^2) which
208 were obtained by means of ANOVA using a Mixed Design Model taking into account
209 repeated measures per subject during the MMTT. Individual predictors included allelic gene
210 scores, microbiota families, gender and MMTT time points. For this analysis, family level
211 microbiota data were used with a relative abundance of $>0.5\%$ in at least 30 % of the
212 samples. Analysis at lower taxonomic ranks is not compatible with the classification
213 confidence of short reads and the taxonomic incongruence of some dominant and prevalent
214 bacterial genera in the human gut.

215 Statistical analysis of microbiota profiles was performed in the R programming environment
216 using Rhea (<https://lagkouvardos.github.io/Rhea/>)(32). OTU tables were normalized to
217 account for differences in sequence depth. β -diversity was computed based on generalized
218 UniFrac distances(12) while α -diversity was assessed on the basis of species richness and
219 Shannon effective diversity(26). For de novo clustering of microbiota profiles, Partitioning
220 Around Medoids was performed as described previously(7).

221 **Results**

222 **Effect of the intervention on plasma bile acids**

223 All subjects underwent the OGTT and MMTT challenges two times, separated by 12 weeks
224 of weight loss intervention. While a subgroup of volunteers (n=32) followed a supervised
225 diet to keep body weight constant, 40 volunteers followed a diet with a 20% energy
226 restriction that led to a mean weight loss of 5.6 kg. This weight loss however did not cause
227 any significant changes in fasting glucose or insulin concentrations (data not shown) nor did
228 it change the composition and concentrations of circulating BA in fasting state and during
229 the dietary challenges (Figure 1). Based on the lack of an effect of the weight-loss we
230 calculated and used the mean values of the BA concentrations during the OGTT and
231 MMTT. In this way we minimized variation and could overcome problems caused by
232 eventual missing samples.

233 **Fasting plasma BA levels**

234 Quantification of the most abundant BA species in plasma (collected between 08:00 and
235 09:00am after 12h of fasting) revealed a high inter-individual variability in both
236 concentrations as well as composition. For the sum of all BA, a 12-fold difference across
237 the 72 volunteers was found. Glycine-conjugated BA accounted for around 47% and
238 taurine-conjugated species for around 7% of the total BA pool, while unconjugated species
239 represented 46% of total (Figure 2 and Figure 3G and 3H time point 0). Primary BA were
240 more abundant than the sum of secondary + tertiary and corresponded to approximately
241 60% of the total BA pool.

242 **Post-prandial responses of plasma BA to OGTT and MMTT**

243 The mean total plasma BA concentration increased 3.3-fold in response to the MMTT
244 (Figure 3A) with maximal levels reached after 1 hour and remaining constant for up to 6
245 hours. Even at 8 hours after consumption of MMTT test drink, concentrations were on
246 average 1.7-fold higher than in overnight fasting state. Bile acid concentrations also
247 increased during the OGTT but here peak concentrations were reached after 30 minutes
248 with levels on average 2-fold higher than in fasting state (Figure 3A). After 90 minutes BA
249 concentrations started to decline to reach fasting state levels after 240 minutes.

250 Different classes of BA displayed different kinetic behaviors. Based on mean values of all
251 volunteers, primary BA increased by 3.9-fold during the MMTT and by 2.1-fold during the
252 OGTT. Secondary + tertiary BA showed a lower response with a maximal increase of 2.8-

253 fold during the MMTT and 1.8-fold during the OGTT compared to fasting levels (Figures 3B
254 and 3C). In the OGTT and MMTT, levels of glycine- and taurine-conjugated BA increased
255 around 5-fold in the MMTT and around 3-fold in the OGTT (Figures 3D and 3E). Plasma
256 concentration of unconjugated BA showed only a small increase during MMTT (1.6-fold at
257 240 min) and even a 15% decrease at the end of the OGTT (Figure 3F). A shift in the
258 composition of circulating BA was observed in the transition from fasting to postprandial
259 state. Glycine-conjugated BA – comprising 47% of the total BA pool in fasting state -
260 increased to 68% during the MMTT and the taurine-conjugated species went from 7% to
261 11%, while unconjugated entities declined from 46% to 20% after 2h in the MMTT (Figure
262 3H). A very similar change in composition was observed during the OGTT, although in this
263 case, the increase in glycine-conjugated BA (and concomitant decline in unconjugated
264 species) was less pronounced (Figure 3G).

265 **Gender-specific differences in post-prandial BA responses**

266 During OGTT, plasma BA concentrations increased similarly in men and women, reaching
267 maximum values after 30 and 60 minutes, respectively. Women thereafter sustained
268 maximal concentrations of BA for at least 2 hours, whereas in men concentrations were
269 significantly reduced already at 90 minutes ($P=0.0011$) as shown in Figure 4A. The gender-
270 specific differences in response to the OGTT were particularly evident when inspecting the
271 glycine- and taurine-conjugated BA ($P<0.0001$ and $P=0.0098$ for differences in AUC for
272 taurine- and glycine-conjugated BA, respectively), as demonstrated in Figures 4B and 4C.
273 Although less pronounced than in the OGTT, gender-related differences were also
274 observed in response to the MMTT, but only for taurine-conjugated BA ($P=0.0085$ for
275 difference in AUC) as shown in Figure 4F. Despite these post-prandial gender differences,
276 overnight fasting BA concentrations were similar between men and women.

277 **Clusters of post-prandial BA responses**

278 Not only fasting BA concentrations but also post-prandial changes in plasma BA displayed
279 large inter-individual variability. There was no correlation between the AUC of post-prandial
280 total BA with their fasting concentration (data not shown).

281 As compared to an average 3.3-fold increase in circulating total BA levels following the
282 MMTT, some subjects responded with a more than 10-fold increase, whereas others barely
283 showed any change. A hierarchical clustering approach based on BA concentration profiles
284 in response to the MMTT identified 3 subgroups of volunteers with an almost equal

285 distribution of men and woman in each cluster. These significant differences in plasma BA
286 kinetics are shown in Figures 5A and 5B. Different classes of BA displayed distinct kinetic
287 behaviors, particularly when unconjugated BA were compared to conjugated species. In
288 cluster 1 (n=35), post-prandial BA concentrations increased on average 4.3-fold with a
289 maximum reached within the first hour. Even after 8 hours, in this cluster BA concentrations
290 were 2.3-fold higher than in fasting state. In cluster 3 (n=21), fasting levels of BA were
291 almost twice as high as in other clusters but here subjects displayed a slower increase in
292 plasma profiles with maximal values of around 2.9-fold of fasting state after 4 hours,
293 returning to fasting state concentrations after 8 hours. Subjects in cluster 2 (n=13)
294 displayed a completely different kinetic profile with plasma concentrations increasing to a
295 maximum of 4.6-fold of fasting levels at 6 hours, and even after 8 hours levels were still 3.4-
296 fold higher than in fasting state.

297 **Determinants of BA profiles**

298 To identify some of the determinants underlying the different BA kinetics, host genome and
299 fecal microbiota sequencing approaches were used. Exome sequencing data were
300 analyzed for genetic heterogeneity in target genes identified by knowledge and GO-
301 classification for a subset of 60 genes involved in hepatic BA synthesis, enterohepatic
302 recirculation, and transmembrane transport of BA. A list of these genes screened for
303 variations is provided on Table 1. Subjects were grouped according to the presence of
304 variants known or predicted to impair protein functionality defined as “deleterious” based on
305 SIFT and Polyphen-2 algorithms. Although various gene variants revealed associations with
306 plasma BA profiles (Table 2), those in the *SLCO1A2* gene were observed in a number of
307 volunteers large enough to allow a reliable association with phenotype. *SLCO1A2* encodes
308 the organic anion-transporting polypeptide-1 A2 (OATP1A2), which mediates BA uptake
309 across the apical membrane of enterocytes in upper small intestine but also in distal
310 nephrons, cholangiocytes and at the blood brain barrier(10, 21). In this gene, a previously
311 described functional polymorphism rs11568563 (c.A516C, p.E172D, MAF=0.03) was found
312 in 9 out of 72 subjects and an additional rare variant rs368672331 (c.G727A, p.G243S) was
313 found in another subject. All rs11568563 carriers displayed a delayed post-prandial
314 increase of conjugated BA in plasma compared to the other volunteers not bearing this
315 variant (Figure 6). Variants in the *SLC10A2* gene that encodes the apical sodium-
316 dependent bile acid transporter (ASBT) in ileum (considered the prime site for BA
317 reabsorption) such as rs71640248, rs117447044 and rs56398830 were also found in 4

318 volunteers that as well showed reduced postprandial increases in BA concentrations (data
319 not shown).

320 Multivariate analyses providing marginal R^2 showed that a substantial proportion of the
321 variation in BA concentration profiles could be attributed to the time point of measurement
322 during the MMTT, gender, selected SNPs and microbiota (Table 2). Across all individual
323 parameters, the lowest cumulative values were found for the cholic acid conjugated species
324 TCA and GCA with 37 and 43% of total variation explained, whereas the highest values
325 were calculated for DCA and its glycine and taurine-conjugates with 69, 63 and 64% of
326 variance explained. Up to 50% of total variation could be attributed to the selected SNPs,
327 particularly in case of DCA and its taurine and glycine conjugates. Variants in the gene
328 encoding Epoxide hydrolase 1 (EPHX1) were found in 39 subjects and this gene alone
329 accounted for 14% of variation in DCA concentrations and approximately 30% of variation
330 in concentrations of its conjugates GDCA and TDCA. Additional individual effects
331 (particularly for unconjugated primary BA) were found for the canalicular multispecific
332 organic anion transporter ABCC3, the racemase AMACR that can interconvert BA as well
333 as for the BA receptor TGR5 encoded by GPBAR1 (variants found in 6 subjects) and for a
334 putative phospholipid-transporting ATPase encoded by ATP8B1, which displayed
335 deleterious variants in 26 volunteers. Given the small size of the study population and the
336 small number of carriers of individual variants no other attempts were made to associate
337 phenotype and genotypes.

338 Gender was particularly associated with the concentrations of taurine-conjugated BA. Most
339 of the selected gut bacterial families had only a marginal effect on BA concentrations, with
340 the exception of *Coriobacteriaceae*, *Erysipelotrichaceae*, and most notably
341 *Peptostreptococcaceae*, which accounted for 12% of variation in CA concentrations.
342 Whereas the family *Coriobacteriaceae* is known to include bile acid-converting species(46,
343 62), results pertaining to the *Peptostreptococcaceae* require additional studies.

344 To further assess the possible influence of gut microbiota in a non-supervised manner, *de*
345 *novo* clustering of fecal microbial profiles based on phylogenetic distances revealed the
346 presence of three distinct clusters of individuals (Figure 7). Each of these clusters was
347 characterized by significant differences in alpha-diversity and in the relative abundance of
348 specific taxa (data not shown), but supervised multidimensional analysis of BA kinetics
349 parameters on the basis of these microbiota-derived clusters followed by statistical tests on
350 individual BA parameters did not reveal any significant associations. Likewise, comparison

351 of the fecal microbiota of individuals belonging to the three different BA-specific clusters
352 (Figure 5) did not show significant differences in any of the bacterial taxonomic groups
353 identified (data not shown).

354

355 **Discussion**

356 The high inter-individual variability of plasma BA levels and composition in the fasting and
357 post-prandial states found in our study confirms previous observations(9, 18, 52). This high
358 variability has intrinsic and extrinsic origins. BA synthesis rate, intestinal absorption and
359 hepatic processing, as well as bacterial metabolism are just a few factors that may
360 contribute to the variability. In addition, acute and chronic effects of diet or drug use and
361 also diurnal variation(14, 18) are known to affect the BA pool. Given the fact that dozens of
362 proteins are involved in BA synthesis and handling in the mammalian system, it can be
363 anticipated that there are also numerous underlying genetic factors. Nies et al. attributed
364 the high inter-individual variability specifically to differences in the expression of
365 transporters involved in hepatic BA transport such as OATP1B1, OATP1B3 and
366 OATP2B1(42). Although no systematic study has explored SNPs or haplotypes in relation
367 to post-prandial plasma BA profiles, a recent study identified epistatic effects in primary BA
368 biosynthesis by employing an Empirical Bayesian Lasso approach for analysis of pathway-
369 based GWAS data(25).

370 Based on blood sampling up to 4 (OGTT) or even 8 hours postprandially in the case of
371 MMTT, distinct kinetic profiles of individual BA species were observed. Maximal increases
372 in plasma levels exceeding fasting concentrations up to 4.6-fold for taurine- as well as
373 glycine-conjugated BA were found at 30 min in the OGTT with sustained levels for over 2
374 hours. These changes match well to those reported previously by others(49)(65). In a study
375 with 73 young volunteers (26 ± 4 years-old), Matysik et al. demonstrated a rise in glycine-
376 and taurine-conjugated BA with a peak at 60 minutes after glucose ingestion, although
377 there was no blood sampling at earlier time points(40). Interestingly, Zhao et. al. reported a
378 biphasic increase in plasma BA, with the first peak at 30 minutes and a second peak at 120
379 minutes(65). Although the mean plasma levels in our study did not reveal a biphasic
380 behavior, some volunteers clearly presented 2 distinct peaks in BA profiles. The fast
381 appearance of the BA in blood during the OGTT - which in essence can be superimposed
382 onto the rise in blood glucose levels (Figure 6C and 6F) - suggests a very rapid uptake of
383 BA in the upper small intestine. Absorption in the duodenum and jejunum is generally not
384 considered as relevant and is postulated to occur by passive diffusion of protonated BA
385 species(13, 54). The main site of BA reuptake and delivery into portal circulation is thought
386 to be the ileum with apical influx via the Na⁺-dependent *SLC10A2* (ASBST) transporter. Our
387 results however strongly argue for absorption of a considerable quantity of BA already in

388 duodenum and jejunum. This applies also for the conjugated BA, since maximal plasma
389 concentrations were observed here at 30 minutes in response to during the OGTT and at
390 60 minutes in response to the MMTT, when the majority of ingested nutrients not even have
391 reached the ileum (Figure 3).

392 The presence of glucose in the intestinal lumen is known to elicit a cholecystokinin (CCK)
393 secretion(37) followed by gallbladder contraction. A rise in plasma BA levels in response to
394 the OGTT is thus per se not surprising. Since BA are not required for glucose absorption,
395 BA secretion may therefore be taken as an archaic evolutionary response with glucose
396 sensing as a surrogate for ingested food. CCK secretion is far stronger in response to a
397 mixed meal and was shown to increase proportional to the lipid content of the meal(37, 50).
398 Increasing CCK-output translates into differences in gallbladder emptying and
399 corresponding plasma BA changes(50, 51). Although we did not determine CCK-levels and
400 gallbladder ejection fraction, it is very reasonable to assume the responses to the OGTT
401 and MMTT in our volunteers was very similar to those described in other studies⁴⁴.
402 Conjugated BA are the most abundant in bile(3, 47) and consequently the post-prandial
403 plasma changes of these BA as compared to the other species were much more
404 pronounced. Concentrations of unconjugated BA only mildly increased in a small subset of
405 subjects after 6 hours in response to the MMTT.

406 It was previously reported that the increase in plasma levels of conjugated BA in response
407 to OGTT and oral lipid tolerance tests are higher in women than in men(23, 48), while no
408 differences are reported in most studies for the fasting state(9, 19, 63). Also in mice, gender
409 effects in the BA pool were observed and those were related to different expression levels
410 of BA biosynthetic enzymes(17, 55). Although all women enrolled in our study were post-
411 menopausal, significant differences in post-prandial levels of conjugated BA were still
412 detectable between men and women, particularly for taurine-conjugated BA during the
413 OGTT, but also during the MMTT.

414 Bile acids quantified in systemic circulation are basically the spillover of a non-complete
415 hepatic extraction of BA in first pass(6, 24). Their concentration in peripheral blood is thus
416 the result of a series of connected processes with secretion from the gallbladder into
417 intestine, transit and absorption across apical and basolateral membranes of enterocytes
418 into portal blood followed by hepatic uptake and biliary secretion. Variability in individual
419 components within this sequence of events will eventually define the shape of the temporal
420 plasma BA profile following a meal. We could group our volunteers based on the post-

421 prandial kinetic profiles into 3 main clusters with different velocities in plasma appearance
422 and absolute concentrations of BA.

423 It can be expected that these differences in plasma BA profiles in response to dietary
424 challenges lead to alterations in physiological processes sensitive to BA. It is known that
425 GLP-1(22, 27, 56) and insulin(1) secretion as well as substrate oxidation (demonstrated in
426 rodents) are subject of regulation by BA. And, indeed, we observed an association of post-
427 prandial total plasma BA and GLP-1 concentrations during the OGTT (Figure 8). When
428 volunteers were clustered according to the increase in plasma levels of glycine-conjugated
429 BA during the first hour of the OGTT, significant differences between the high and low
430 responders were observed for circulating GLP-1 levels. This suggests that the rise in GLP-1
431 initiated by glucose administration may be modulated by BA released into the intestine of by
432 circulating BA reaching enteroendocrine cells (Figure 8C). The effect was most prominent
433 for glycine-conjugated BA as the responsive BA category which also has the highest
434 circulating concentrations. This relationship between BA secretion and corresponding GLP-
435 1 levels was however not observed in the MMTT (Figure 8 D). The MMTT drink also
436 contained lipids and proteins that can also promote GLP-1 secretion, resulting in a 2-fold
437 higher plasma GLP-1 and BA levels as compared to the OGTT. Lipids and proteins seem to
438 promote a stronger GLP-1 output that may overrule the BA modulatory effects on GLP-1
439 secretion during the MMTT.

440 One of the most important outcomes of the present study is that genetic variants in the BA
441 transporter OATP1A2 associate with differences in postprandial BA kinetics but not the
442 overnight fasting plasma BA levels. Carriers of the nonsynonymous SNP (c.A516C,
443 p.E172D) in the *SLCO1A2* gene displayed significantly lower plasma BA concentrations in
444 response to the OGTT and during the first hours of the MMTT, indicating reduced and
445 delayed absorption of BA from intestine. Since OATP1A2 is expressed in the duodenum(4,
446 20) it is the candidate transporter that could mediate the early phase of BA absorption
447 leading to rapid changes in plasma BA levels as observed in response to the OGTT in
448 which peak BA levels can be superimposed on glucose peaks. The A516C variant of
449 *SLCO1A2* was demonstrated to markedly reduce the transport capacity for different
450 substrates when expressed heterologously(8, 34) although this has not been shown for BA
451 as substrates. Post-prandial concentrations of unconjugated BA, which are thought to be
452 taken up into enterocytes via diffusion (not depending on transporters) were not affected in
453 volunteers with the A516C variant, corroborating the importance of this transporter for

454 duodenal BA transport. Although BA concentrations could not be measured in portal blood
455 in the present study, BA in systemic circulation are considered as a surrogate of
456 enterohepatic circulation (5), corroborating our hypothesis of a slower intestinal BA uptake
457 in carriers of the deleterious variation in *SLCO1A2*.

458 Taken together, our study describes different BA kinetics following an OGTT and a MMTT
459 and identified key determinants underlying the large inter-individual variability in
460 postprandial BA profiles. The different patterns of post-prandial BA responses, associated
461 with fasting concentrations of BA allowed the classification of the 72 subjects into the 3
462 major clusters or “metabotypes”. The finding that heterogeneity in 60 preselected genes of
463 BA synthesis and transport explained most of the BA variance argues for these
464 metabotypes as mainly genetically determined and not too much dependent on gut
465 microbiota structure as measured by amplicon sequencing in spite of the known
466 involvement of specific bacterial species in bile acid metabolism in the intestine. Finally, the
467 present study is the first to demonstrate an association between a common genetic variant
468 in the OATP1A2 transporter and post-prandial BA kinetics, despite the absence of any
469 effects in the fasting state. Understanding the metabolism of BA in the post-prandial state is
470 essential to better understand their roles in human physiology.

471

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481

482 **References**

- 483 1. **Adrian TE, Gariballa S, Parekh KA, Thomas SA, Saadi H, Kaabi Al J, Nagelkerke**
484 **N, Gedulin B, Young AA.** Rectal taurocholate increases L cell and insulin secretion,
485 and decreases blood glucose and food intake in obese type 2 diabetic volunteers.
486 *Diabetologia* 55: 2343–2347, 2012.
- 487 2. **Adzhubei I, Jordan DM, Sunyaev SR.** Predicting functional effect of human
488 missense mutations using PolyPhen-2. *Curr Protoc Hum Genet* Chapter 7: Unit7.20,
489 2013.
- 490 3. **Alnouti Y, Csanaky IL, Klaassen CD.** Quantitative-profiling of bile acids and their
491 conjugates in mouse liver, bile, plasma, and urine using LC-MS/MS. *J Chromatogr B*
492 *Analyt Technol Biomed Life Sci* 873: 209–217, 2008.
- 493 4. **Amelsberg A, Jochims C, Richter CP, Nitsche R, Fölsch UR.** Evidence for an
494 anion exchange mechanism for uptake of conjugated bile acid from the rat jejunum.
495 *Am J Physiol* 276: G737–42, 1999.
- 496 5. **Angelin B, Björkhem I, Einarsson K, Ewerth S.** Hepatic uptake of bile acids in
497 man. Fasting and postprandial concentrations of individual bile acids in portal venous
498 and systemic blood serum. *J Clin Invest* 70: 724–731, 1982.
- 499 6. **Angelin B, Einarsson K, HELLSTROM K.** Evidence for the absorption of bile acids
500 in the proximal small intestine of normo- and hyperlipidaemic subjects. *Gut* 17: 420–
501 425, 1976.
- 502 7. **Arumugam M, Raes J, Pelletier E, Le Paslier D, Yamada T, Mende DR,**
503 **Fernandes GR, Tap J, Bruls T, Batto J-M, Bertalan M, Borrueal N, Casellas F,**
504 **Fernandez L, Gautier L, Hansen T, Hattori M, Hayashi T, Kleerebezem M,**
505 **Kurokawa K, Leclerc M, Levenez F, Manichanh C, Nielsen HB, Nielsen T, Pons**
506 **N, Poulain J, Qin J, Sicheritz-Ponten T, Tims S, Torrents D, Ugarte E, Zoetendal**
507 **EG, Wang J, Guarner F, Pedersen O, de Vos WM, Brunak S, Doré J, MetaHIT**
508 **Consortium, Antolín M, Artiguenave F, Blottiere HM, Almeida M, Brechot C,**
509 **Cara C, Chervaux C, Cultrone A, Delorme C, Denariáz G, Dervyn R, Foerstner**
510 **KU, Friss C, van de Guchte M, Guedon E, Haimet F, Huber W, van Hylckama-**
511 **Vlieg J, Jamet A, Juste C, Kaci G, Knol J, Lakhdari O, Layec S, Le Roux K,**
512 **Maguin E, Mérieux A, Melo Minardi R, M'rini C, Muller J, Oozeer R, Parkhill J,**
513 **Renault P, Rescigno M, Sanchez N, Sunagawa S, Torrejon A, Turner K,**
514 **Vandemeulebrouck G, Varela E, Winogradsky Y, Zeller G, Weissenbach J,**
515 **Ehrlich SD, Bork P.** Enterotypes of the human gut microbiome. *Nature* 473: 174–
516 180, 2011.
- 517 8. **Badagnani I, Castro RA, Taylor TR, Brett CM, Huang CC, Stryke D, Kawamoto**
518 **M, Johns SJ, Ferrin TE, Carlson EJ, Burchard EG, Giacomini KM.** Interaction of
519 methotrexate with organic-anion transporting polypeptide 1A2 and its genetic
520 variants. *J Pharmacol Exp Ther* 318: 521–529, 2006.
- 521 9. **Bathena SPR, Thakare R, Gautam N, Mukherjee S, Olivera M, Meza J, Alnouti Y.**
522 **Urinary bile acids as biomarkers for liver diseases I. Stability of the baseline profile in**
523 **healthy subjects.** *Toxicol Sci* 143: 296–307, 2015.

- 524 10. **Bronger H, König J, Kopplow K, Steiner H-H, Ahmadi R, Herold-Mende C,**
525 **Keppler D, Nies AT.** ABCC drug efflux pumps and organic anion uptake transporters
526 in human gliomas and the blood-tumor barrier. *Cancer Res* 65: 11419–11428, 2005.
- 527 11. **Caporaso JG, Lauber CL, Walters WA, Berg-Lyons D, Huntley J, Fierer N,**
528 **Owens SM, Betley J, Fraser L, Bauer M, Gormley N, Gilbert JA, Smith G, Knight**
529 **R.** Ultra-high-throughput microbial community analysis on the Illumina HiSeq and
530 MiSeq platforms. *ISME J* 6: 1621–1624, 2012.
- 531 12. **Chen J, Bittinger K, Charlson ES, Hoffmann C, Lewis J, Wu GD, Collman RG,**
532 **Bushman FD, Li H.** Associating microbiome composition with environmental
533 covariates using generalized UniFrac distances. *Bioinformatics* 28: 2106–2113, 2012.
- 534 13. **Dawson PA, Lan T, Rao A.** Bile acid transporters. *J Lipid Res* 50: 2340–2357, 2009.
- 535 14. **Duane WC, Levitt DG, Mueller SM, Behrens JC.** Regulation of bile acid synthesis in
536 man. Presence of a diurnal rhythm. *J Clin Invest* 72: 1930–1936, 1983.
- 537 15. **Edgar RC, Haas BJ, Clemente JC, Quince C, Knight R.** UCHIME improves
538 sensitivity and speed of chimera detection. *Bioinformatics* 27: 2194–2200, 2011.
- 539 16. **Edgar RC.** UPARSE: highly accurate OTU sequences from microbial amplicon reads.
540 *Nat Methods* 10: 996–998, 2013.
- 541 17. **Fu ZD, Csanaky IL, Klaassen CD.** Gender-divergent profile of bile acid homeostasis
542 during aging of mice. *PLoS ONE* 7: e32551, 2012.
- 543 18. **Gälman C, Angelin B, Rudling M.** Bile acid synthesis in humans has a rapid diurnal
544 variation that is asynchronous with cholesterol synthesis. *Gastroenterology* 129:
545 1445–1453, 2005.
- 546 19. **Gälman C, Angelin B, Rudling M.** Pronounced variation in bile acid synthesis in
547 humans is related to gender, hypertriglyceridaemia and circulating levels of fibroblast
548 growth factor 19. *J Intern Med* 270: 580–588, 2011.
- 549 20. **Glaeser H, Bailey DG, Dresser GK, Gregor JC, Schwarz UI, McGrath JS,**
550 **Jolicoeur E, Lee W, Leake BF, Tirona RG, Kim RB.** Intestinal drug transporter
551 expression and the impact of grapefruit juice in humans. *Clin Pharmacol Ther* 81:
552 362–370, 2007.
- 553 21. **Hagenbuch B, Stieger B.** The SLCO (former SLC21) superfamily of transporters.
554 *Molecular Aspects of Medicine* 34: 396–412, 2013.
- 555 22. **Hansen M, Scheltema MJ, Sonne DP, Hansen JS, Sperling M, Rehfeld JF, Holst**
556 **JJ, Vilsbøll T, Knop FK.** Effect of chenodeoxycholic acid and the bile acid
557 sequestrant colesevelam on glucagon-like peptide-1 secretion. *Diabetes Obes Metab*
558 18: 571–580, 2016.
- 559 23. **Ho JE, Larson MG, Vasan RS, Ghorbani A, Cheng S, Rhee EP, Florez JC, Clish**
560 **CB, Gerszten RE, Wang TJ.** Metabolite profiles during oral glucose challenge.
561 *Diabetes* 62: 2689–2698, 2013.

- 562 24. **Houten SM, Watanabe M, Auwerx J.** Endocrine functions of bile acids. *EMBO J* 25:
563 1419–1425, 2006.
- 564 25. **Huang A, Martin ER, Vance JM, Cai X.** Detecting genetic interactions in pathway-
565 based genome-wide association studies. *Genet Epidemiol* 38: 300–309, 2014.
- 566 26. **Jost L.** Partitioning diversity into independent alpha and beta components. *Ecology*
567 88: 2427–2439, 2007.
- 568 27. **Katsuma S, Hirasawa A, Tsujimoto G.** Bile acids promote glucagon-like peptide-1
569 secretion through TGR5 in a murine enteroendocrine cell line STC-1. *Biochem*
570 *Biophys Res Commun* 329: 386–390, 2005.
- 571 28. **Kawamata Y, Fujii R, Hosoya M, Harada M, Yoshida H, Miwa M, Fukusumi S,**
572 **Habata Y, Itoh T, Shintani Y, Hinuma S, Fujisawa Y, Fujino M.** A G protein-
573 coupled receptor responsive to bile acids. *J Biol Chem* 278: 9435–9440, 2003.
- 574 29. **Kelder T, Stroeve JHM, Bijlsma S, Radonjic M, Roeselers G.** Correlation network
575 analysis reveals relationships between diet-induced changes in human gut microbiota
576 and metabolic health. *Nutr Diabetes* 4: e122, 2014.
- 577 30. **Kreymann B, Williams G, Ghatei MA, Bloom SR.** Glucagon-like peptide-1 7-36: a
578 physiological incretin in man. *Lancet* 2: 1300–1304, 1987.
- 579 31. **Kumar P, Henikoff S, Ng PC.** Predicting the effects of coding non-synonymous
580 variants on protein function using the SIFT algorithm. *Nat Protoc* 4: 1073–1081, 2009.
- 581 32. **Lagkourdos I, Fischer S, Kumar N, Clavel T.** Rhea: a transparent and modular R
582 pipeline for microbial profiling based on 16S rRNA gene amplicons. *PeerJ* 5: e2836,
583 2017.
- 584 33. **Lagkourdos I, Kläring K, Heinzmann SS, Platz S, Scholz B, Engel K-H,**
585 **Schmitt-Kopplin P, Haller D, Rohn S, Skurk T, Clavel T.** Gut metabolites and
586 bacterial community networks during a pilot intervention study with flaxseeds in
587 healthy adult men. *Mol Nutr Food Res* 59: 1614–1628, 2015.
- 588 34. **Lee W, Glaeser H, Smith LH, Roberts RL, Moeckel GW, Gervasini G, Leake BF,**
589 **Kim RB.** Polymorphisms in human organic anion-transporting polypeptide 1A2
590 (OATP1A2): implications for altered drug disposition and central nervous system drug
591 entry. *J Biol Chem* 280: 9610–9617, 2005.
- 592 35. **Lefebvre P, Cariou B, Lien F, Kuipers F, Staels B.** Role of bile acids and bile acid
593 receptors in metabolic regulation. *Physiol Rev* 89: 147–191, 2009.
- 594 36. **Li H, Durbin R.** Fast and accurate long-read alignment with Burrows-Wheeler
595 transform. *Bioinformatics* 26: 589–595, 2010.
- 596 37. **Liddle RA, Goldfine ID, Rosen MS, Taplitz RA, Williams JA.** Cholecystokinin
597 bioactivity in human plasma. Molecular forms, responses to feeding, and relationship
598 to gallbladder contraction. *J Clin Invest* 75: 1144–1152, 1985.
- 599 38. **Ma K, Saha PK, Chan L, Moore DD.** Farnesoid X receptor is essential for normal

- 600 glucose homeostasis. *J Clin Invest* 116: 1102–1109, 2006.
- 601 39. **Maruyama T, Miyamoto Y, Nakamura T, Tamai Y, Okada H, Sugiyama E,**
602 **Nakamura T, Itadani H, Tanaka K.** Identification of membrane-type receptor for bile
603 acids (M-BAR). *Biochem Biophys Res Commun* 298: 714–719, 2002.
- 604 40. **Matysik S, Martin J, Bala M, Scherer M, Schäffler A, Schmitz G.** Bile acid
605 signaling after an oral glucose tolerance test. *Chem Phys Lipids* 164: 525–529, 2011.
- 606 41. **McKenna A, Hanna M, Banks E, Sivachenko A, Cibulskis K, Kernytsky A,**
607 **Garimella K, Altshuler D, Gabriel S, Daly M, DePristo MA.** The Genome Analysis
608 Toolkit: a MapReduce framework for analyzing next-generation DNA sequencing
609 data. *Genome Res* 20: 1297–1303, 2010.
- 610 42. **Nies AT, Niemi M, Burk O, Winter S, Zanger UM, Stieger B, Schwab M,**
611 **Schaeffeler E.** Genetics is a major determinant of expression of the human hepatic
612 uptake transporter OATP1B1, but not of OATP1B3 and OATP2B1. *Genome Med* 5:
613 1, 2013.
- 614 43. **Parks DJ, Blanchard SG, Bledsoe RK, Chandra G, Consler TG, Kliewer SA,**
615 **Stimmel JB, Willson TM, Zavacki AM, Moore DD, Lehmann JM.** Bile acids: natural
616 ligands for an orphan nuclear receptor. *Science* 284: 1365–1368, 1999.
- 617 44. **Popescu IR, Helleboid-Chapman A, Lucas A, Vandewalle B, Dumont J,**
618 **Bouchaert E, Derudas B, Kerr-Conte J, Caron S, Pattou F, Staels B.** The nuclear
619 receptor FXR is expressed in pancreatic beta-cells and protects human islets from
620 lipotoxicity. *FEBS Lett* 584: 2845–2851, 2010.
- 621 45. **Renga B, Mencarelli A, Vavassori P, Brancaleone V, Fiorucci S.** The bile acid
622 sensor FXR regulates insulin transcription and secretion. *Biochim Biophys Acta* 1802:
623 363–372, 2010.
- 624 46. **Ridlon JM, Kang D-J, Hylemon PB.** Bile salt biotransformations by human intestinal
625 bacteria. *J Lipid Res* 47: 241–259, 2006.
- 626 47. **Rossi SS, Converse JL, Hofmann AF.** High pressure liquid chromatographic
627 analysis of conjugated bile acids in human bile: simultaneous resolution of sulfated
628 and unsulfated lithocholyl amidates and the common conjugated bile acids. *J Lipid*
629 *Res*.
- 630 48. **Schmid A, Neumann H, Karrasch T, Liebisch G, Schäffler A.** Bile Acid
631 Metabolome after an Oral Lipid Tolerance Test by Liquid Chromatography-Tandem
632 Mass Spectrometry (LC-MS/MS). *PLoS ONE* 11: e0148869, 2016.
- 633 49. **Shaham O, Wei R, Wang TJ, Ricciardi C, Lewis GD, Vasan RS, Carr SA,**
634 **Thadhani R, Gerszten RE, Mootha VK.** Metabolic profiling of the human response
635 to a glucose challenge reveals distinct axes of insulin sensitivity. *Mol Syst Biol* 4: 214,
636 2008.
- 637 50. **Sonne DP, Rehfeld JF, Holst JJ, Vilsbøll T, Knop FK.** Postprandial gallbladder
638 emptying in patients with type 2 diabetes: potential implications for bile-induced
639 secretion of glucagon-like peptide 1. *Eur J Endocrinol* 171: 407–419, 2014.

- 640 51. **Sonne DP, Samuel van Nierop F, Kulik W, Soeters MR, Vilsbøll T, Knop FK.**
641 Postprandial Plasma Concentrations of Individual Bile Acids and FGF-19 in Patients
642 with Type 2 Diabetes. *J Clin Endocrinol Metab* (June 7, 2016). doi: 10.1210/jc.2016-
643 1607.
- 644 52. **Steiner C, Othman A, Saely CH, Rein P, Drexel H, Eckardstein von A, Rentsch**
645 **KM.** Bile acid metabolites in serum: intraindividual variation and associations with
646 coronary heart disease, metabolic syndrome and diabetes mellitus. *PLoS ONE* 6:
647 e25006, 2011.
- 648 53. **Tagliacozzi D, Mozzi AF, Casetta B, Bertucci P, Bernardini S, Di Ilio C, Urbani A,**
649 **Federici G.** Quantitative analysis of bile acids in human plasma by liquid
650 chromatography-electrospray tandem mass spectrometry: a simple and rapid one-
651 step method. *Clin Chem Lab Med* 41: 1633–1641, 2003.
- 652 54. **Trauner M, Boyer JL.** Bile Salt Transporters: Molecular Characterization, Function,
653 and Regulation. *Physiol Rev* 83: 633–671, 2003.
- 654 55. **Turley SD, Schwarz M, Spady DK, Dietschy JM.** Gender-related differences in bile
655 acid and sterol metabolism in outbred CD-1 mice fed low- and high-cholesterol diets -
656 Turley - 2003 - Hepatology - Wiley Online Library. *Hepatology*.
- 657 56. **Vanitha Bala SRDPKADNSMAJSJRGKSM.** Release of GLP-1 and PYY in response
658 to the activation of G protein-coupled bile acid receptor TGR5 is mediated by
659 Epac/PLC- ϵ pathway and modulated by endogenous H₂S. *Frontiers in Physiology* 5,
660 2014.
- 661 57. **Wang H, Chen J, Hollister K, Sowers LC, Forman BM.** Endogenous bile acids are
662 ligands for the nuclear receptor FXR/BAR. *Mol Cell* 3: 543–553, 1999.
- 663 58. **Wang K, Li M, Hakonarson H.** ANNOVAR: functional annotation of genetic variants
664 from high-throughput sequencing data. *Nucleic Acids Res* 38: e164, 2010.
- 665 59. **Wang Q, Garrity GM, Tiedje JM, Cole JR.** Naive Bayesian classifier for rapid
666 assignment of rRNA sequences into the new bacterial taxonomy. *Appl Environ*
667 *Microbiol* 73: 5261–5267, 2007.
- 668 60. **Watanabe M, Houten SM, Matakai C, Christoffolete MA, Kim BW, Sato H,**
669 **Messaddeq N, Harney JW, Ezaki O, Kodama T, Schoonjans K, Bianco AC,**
670 **Auwerx J.** Bile acids induce energy expenditure by promoting intracellular thyroid
671 hormone activation. *Nature* 439: 484–489, 2006.
- 672 61. **Watanabe M, Houten SM, Wang L, Moschetta A, Mangelsdorf DJ, Heyman RA,**
673 **Moore DD, Auwerx J.** Bile acids lower triglyceride levels via a pathway involving
674 FXR, SHP, and SREBP-1c. *J Clin Invest* 113: 1408–1418, 2004.
- 675 62. **Wegner K, Just S, Gau L, Mueller H, Gérard P, Lepage P, Clavel T, Rohn S.**
676 Rapid analysis of bile acids in different biological matrices using LC-ESI-MS/MS for
677 the investigation of bile acid transformation by mammalian gut bacteria. *Anal Bioanal*
678 *Chem* 409: 1231–1245, 2017.
- 679 63. **Xiang X, Backman JT, Neuvonen PJ, Niemi M.** Gender, but not CYP7A1 or

680 SLCO1B1 polymorphism, affects the fasting plasma concentrations of bile acids in
681 human beings. *Basic Clin Pharmacol Toxicol* 110: 245–252, 2012.

682 64. **Yamagata K, Daitoku H, Shimamoto Y, Matsuzaki H, Hirota K, Ishida J,**
683 **Fukamizu A.** Bile acids regulate gluconeogenic gene expression via small
684 heterodimer partner-mediated repression of hepatocyte nuclear factor 4 and Foxo1. *J*
685 *Biol Chem* 279: 23158–23165, 2004.

686 65. **Zhao X, Peter A, Fritsche J, Elcnerova M, Fritsche A, Häring H-U, Schleicher ED,**
687 **Xu G, Lehmann R.** Changes of the plasma metabolome during an oral glucose
688 tolerance test: is there more than glucose to look at? *Am J Physiol Endocrinol Metab*
689 296: E384–93, 2009.

690 66. **Zietak M, Kozak LP.** Bile acids induce uncoupling protein 1-dependent
691 thermogenesis and stimulate energy expenditure at thermoneutrality in mice. *Am J*
692 *Physiol Endocrinol Metab* (December 29, 2015). doi: 10.1152/ajpendo.00485.2015.

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705 **Figure Legends**

706

707 **Table 1** – List of selected genes known as involved in BA synthesis and transport that were
708 screened for deleterious variations. Genes selected represent hormones, membrane
709 receptors, transcription factors and enzymes involved in BA synthesis as well as
710 transporters expressed in hepatocytes, bile canaliculi and intestine.

711

712 **Table 2** – Effect of different parameters identified by multivariate analyses as responsible
713 for the observed variation in the most abundant bile acids in plasma in postprandial state.
714 The numbers in each cell indicate the proportion of variation (in percentage) in each BA
715 explained by the corresponding variable. The numbers in brackets after gene names
716 indicate the number of subjects carrying variations in the respective gene. Highlighted cells
717 indicate significant associations ($P < 0.05$).

718

719 **Figure 1 – The energy restriction did not affect BA profiles during OGTT and MMTT**
720 **or after a 12-hour fasting.** A – Total BA in plasma during the OGTT. B – Total BA in
721 plasma during the MMTT. Continuous lines represent BA profile before and dotted lines, the
722 BA profile after a 12 weeks period of intervention with 20% dietary energy restriction. Only
723 the volunteers that abide to the energy restriction are represented (in control group no
724 changes due to the energy restriction were observed as well). C – Total BA and BA classes
725 in plasma after a 12-hour fasting. N = 40.

726

727 **Figure 2 – Fasting plasma bile acids.** Plasma concentrations of the most abundant BA in
728 healthy subjects in the overnight fasting state (average of sampling performed at $t=0$ of
729 OGTT and MMTT before and after the weight loss intervention). Each differently colored
730 segment in bars represents the concentration of an individual BA.

731

732 **Figure 3 – Plasma BA concentration profiles in response to the OGTT and MMTT.** A to
733 F - Plasma concentrations of different BA classes during the OGTT (dotted line) and MMTT
734 (full line). G and H – Changes in the composition of the plasma BA pool according to their
735 conjugation type during the OGTT (G) and MMTT (H). Results are shown as the mean \pm
736 SEM of all volunteers averaging the results of the two challenges performed before and
737 after 12 weeks ($n=72$).

738

739 **Figure 4 – Gender effects on plasma BA concentration profiles during the OGTT and**
740 **MMTT.**

741 **A to C** – Plasma concentrations of total, glycine- and taurine-conjugated BA in men (dotted
742 lines, n=35) and women (full lines, n=37) during the OGTT. **D to F** – Plasma concentrations
743 of total, glycine- and taurine-conjugated BA in men (dotted lines, n=35) and women (full
744 lines, n=37) during the MMTT. Results are shown as the mean \pm SEM, averaging the
745 results of challenges performed before and after intervention. Significant differences in the
746 AUC after t- test are indicated in the graphs.

747

748 **Figure 5 – Clustering of volunteers according to postprandial plasma BA kinetics.**

749 **A** – Clustering of volunteers based on plasma concentration profiles of bile acids during the
750 MMTT. **B** – Changes in total and different classes of bile acids during the MMTT in
751 participants of the 3 clusters. Results are expressed as mean \pm SEM. n=35 in cluster 1,
752 n=13 in cluster 2 and n=21 in cluster 3. **C** – Summary of differences found among the 3
753 clusters for the different bile acid classes as represented in pane B ($P \leq 0.05$).

754

755 **Figure 6 – Effects of the functional polymorphism A516C in *SLCO1A2* (*OATP1A2*) on**
756 **plasma BA kinetics. A to C** – Plasma concentrations of bile acids in A516C carriers
757 (dotted line, n=9) and non-carriers (full line, n=62) during the MMTT. **D to F** – Plasma
758 concentrations of bile acids in A516C carriers and non-carriers during the OGTT. Results
759 are the mean \pm SEM of BA levels averaging the results of challenges performed before and
760 after the weight loss intervention. Statistical differences (multiple T-tests) are indicated in
761 each graph. The grey area in panes C and F represents the profile of glucose during the
762 MMTT and OGTT, respectively.

763

764 **Figure 7 – *De novo* clustering analysis of fecal microbiota profiles as obtained by**
765 **high-throughput sequencing of 16S rRNA gene amplicons.** Partitioning Around
766 Medoids was performed as described previously. The clustering with best support was
767 visualized with an NMDS plot (**A**). Associated changes in alpha-diversity (**B**) and differential
768 abundances of taxonomic groups among the clusters (**C**) were tested statistically in the R
769 programming environment using Rhea.

770

771 **Figure 8 – Relationship between plasma bile acids and GLP-1 concentrations in the**
772 **post-prandial state. A and B** – Plasma concentrations of glycine-conjugated BA (black

773 lines) and GLP-1 (grey lines) in response to the OGTT (A) and MMTT (B) (N=72). **C and D**
774 – Plasma concentration of glycine-conjugated BA (black lines) and GLP-1 (grey lines).
775 Subjects were ranked according to post-prandial increase of glycine-conjugated BA
776 concentrations during the first hour of the OGTT (C) and MMTT (D) (N=36 in each group).
777 Results are expressed as mean \pm SEM of BA fold-change from t=0.

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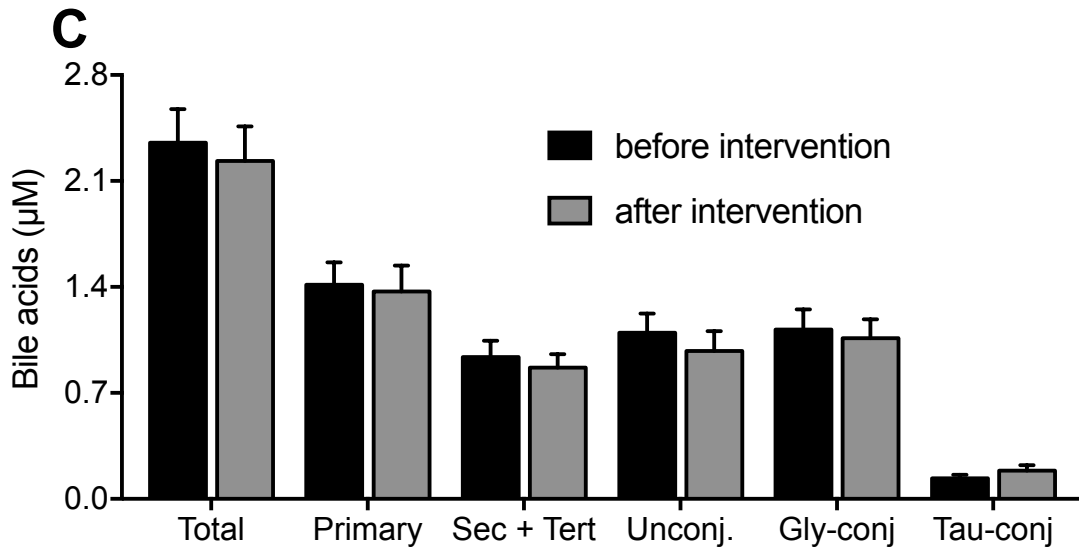
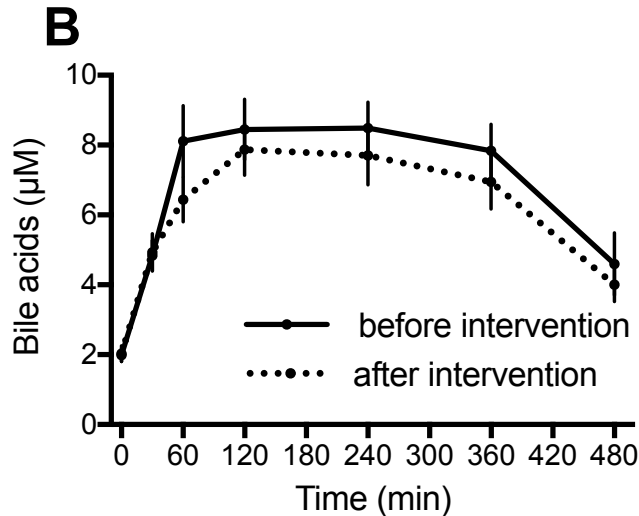
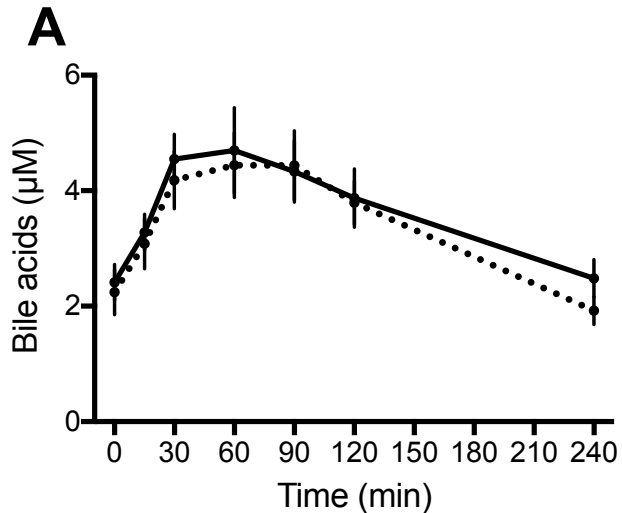
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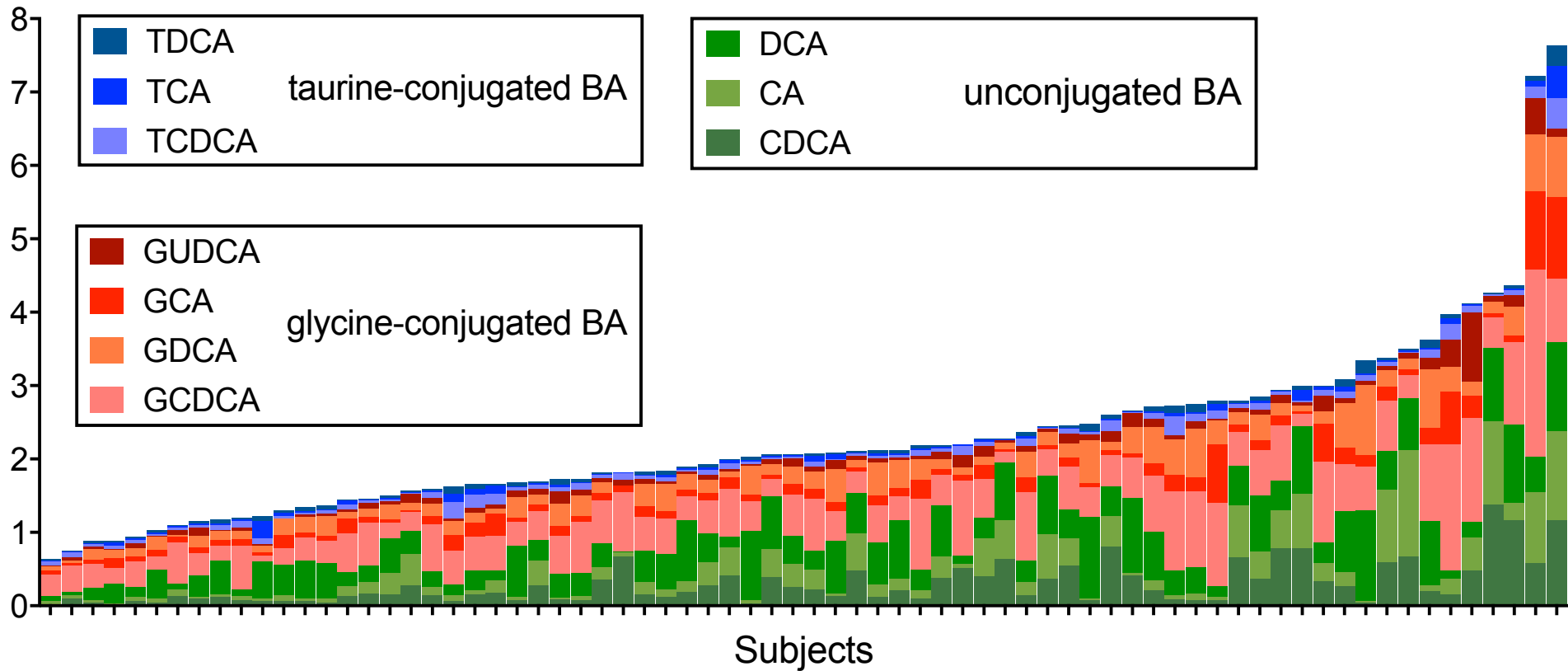
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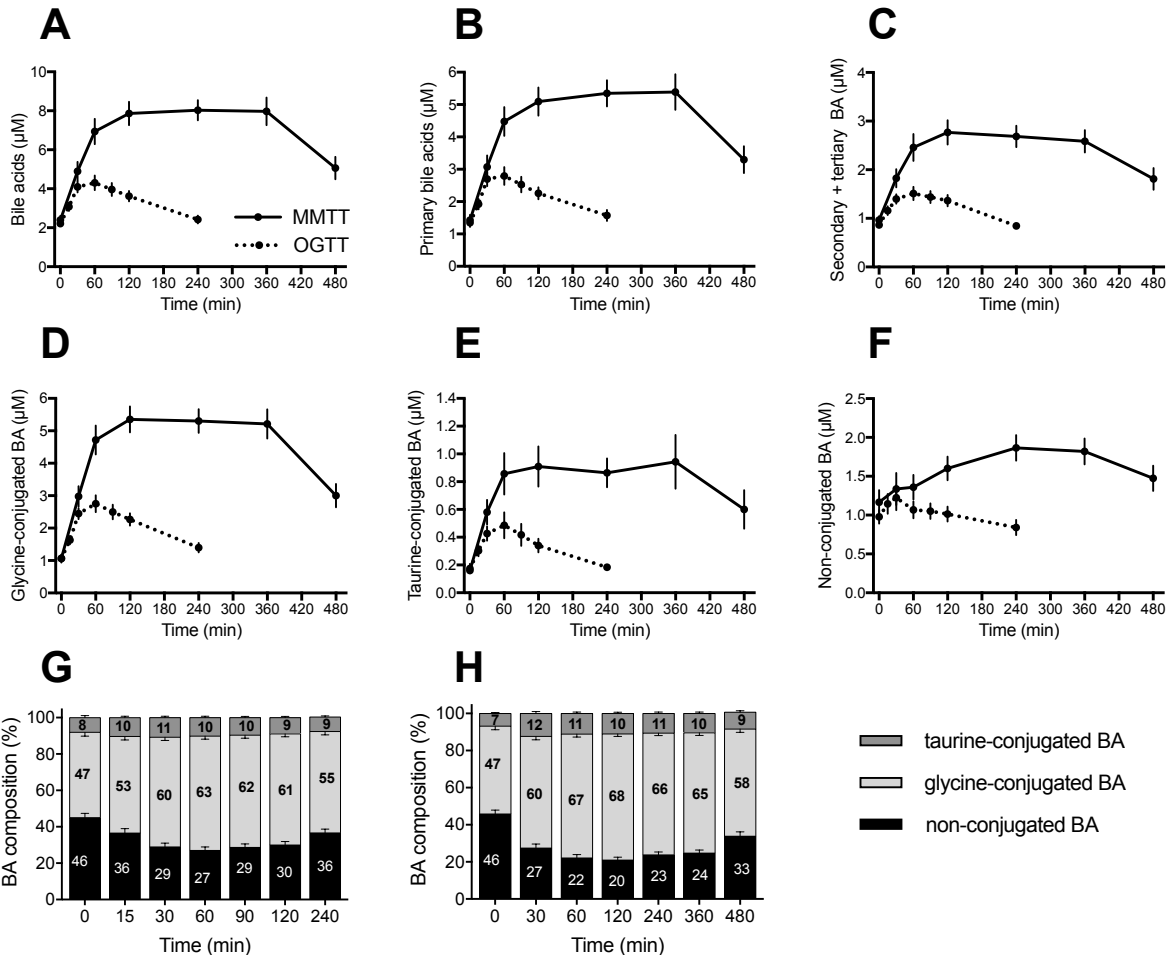
Protein Name	Symbol	Gene symbol	Function
Cholesterol 7 α -hydroxylase	CYP7A1	CYP7A1	Enzyme
Sterol 27-hydroxylase	CYP27A1	CYP27A1	Enzyme
25-hydroxycholesterol 7 α -hydroxylase	CYP7B1	CYP7B1	Enzyme
24-hydroxycholesterol 7 α -hydroxylase	CYP39A1	CYP39A1	Enzyme
Microsomal 3 β -hydroxy- Δ 5-C27-steroid oxidoreductase	HSD3B7	HSD3B7	Enzyme
Microsomal sterol 12 α -hydroxylase	CYP8B1	CYP8B1	Enzyme
Δ 4-3-oxosteroid 5 β -reductase	AKR1D1	AKR1D1	Enzyme
Acyl-CoA Oxidase 1	ACOX1	ACOX1	Enzyme
Acyl-CoA Oxidase 2	ACOX2	ACOX2	Enzyme
Bile Acid-CoA:Amino Acid N-Acyltransferase	BAAT	BAAT	Enzyme
Bile Acid-CoA Ligase	BAL	SLC27A5	Enzyme
α -Methylacyl-CoA Racemase	AMACR	AMACR	Enzyme
D-bifunctional enzyme	DBP	DBP	Enzyme
Peroxisomal Thiolase 2	SCP2	SCP2	Enzyme
Aldo-Keto Reductase Family 1 Member C4	AKR1C4	AKR1C4	Enzyme
Hydroxysteroid Sulfotransferase	HST	SULT2A1	Enzyme
UDP glucuronosyltransferase Family 2 Member B4	UGT2B4	UGT2B4	Enzyme
UDP glucuronosyltransferase Family 2 Member B7	UGT2B7	UGT2B7	Enzyme
UDP glucuronosyltransferase Family 1 Member A3	UGT1A3	UGT1A3	Enzyme
Solute Carrier Family 4 Member 2	SLC4A2	SLC4A2	Enzyme
Microsomal Epoxide Hydrolase	mEH	EPHX1	Enzyme
Cholecystokinin	CCK	CCK	Hormone
Fibroblast Growth Factor 19	FGF19	FGF19	Hormone
Farnesoid X Receptor	FXR	NR1H4	Nuclear receptor
Small Heterodimer Partner	SHP	NR0B2	Nuclear receptor
Liver X Receptor	LXR	NR1H3	Nuclear receptor
Forkhead Box A2	HNF-3 β	FOXA2	Nuclear receptor
Hepatocyte Nuclear Factor-4 α	HNF-4 α	HNF4A	Nuclear receptor
G Protein-Coupled Bile Acid Receptor 1	GPBAR1	GPBAR1	Receptor
Bile Salt Export Pump	BSEP	ABCB11	Transporter
Apical Sodium-dependent BA Transporter	ASBT	SLC10A2	Transporter
Organic Solute Transporter- α subunit	OST α	SLC51A	Transporter
Organic Solute Transporter- β subunit	OST β	SLC51B	Transporter
Sodium/Taurocholate Cotransporting Polypeptide	NTCP	SLC10A1	Transporter
Solute Carrier Organic Anion transporter Family Member 1A2	OATP1A2	SLCO1A2	Transporter
Solute Carrier Organic Anion transporter Family Member 1B1	OATP1B1	SLCO1B1	Transporter
Solute Carrier Organic Anion transporter Family Member 1B3	OATP1B3	SLCO1B3	Transporter
Solute Carrier Organic Anion transporter Family Member 2B1	OATP2B1	SLCO2B1	Transporter
ATP Binding Cassette Subfamily C Member 3	ABCC3	ABCC3	Transporter
ATP Binding Cassette Subfamily A Member 1	ABCA1	ABCA1	Transporter
ATP Binding Cassette Subfamily A Member 3	ABCA3	ABCA3	Transporter
ATP Binding Cassette Subfamily G Member 2	ABCG2	ABCG2	Transporter
ATP Binding Cassette Subfamily G Member 5	ABCG5	ABCG5	Transporter
ATP Binding Cassette Subfamily G Member 8	ABCG8	ABCG8	Transporter
ATP Binding Cassette Subfamily C Member 2	ABCC2	ABCC2	Transporter
ATP Binding Cassette Subfamily B Member 1	ABCB1	ABCB1	Transporter
ATP Binding Cassette Subfamily B Member 4	ABCB4	ABCB4	Transporter
ATPase Phospholipid Transporting 8B1	ATP8B1	ATP8B1	Transporter
Cytosolic Intestinal BA-binding Protein	IBABP	FABP6	Transporter

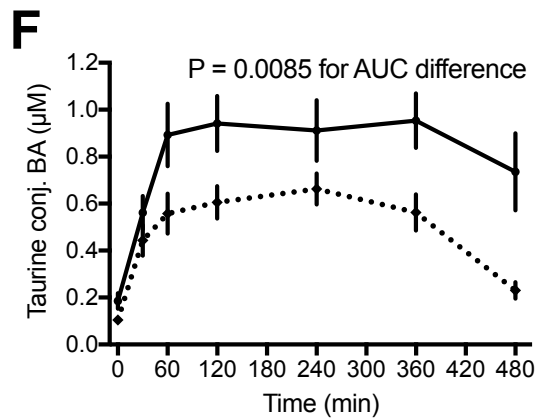
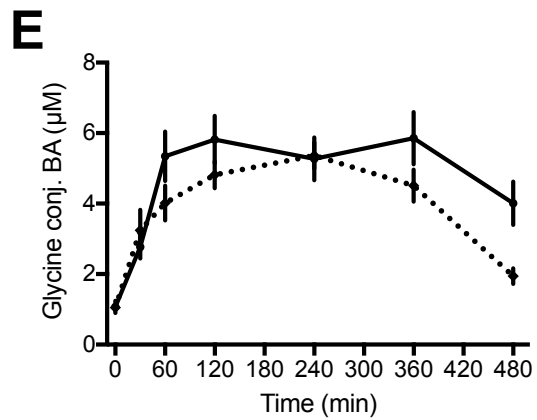
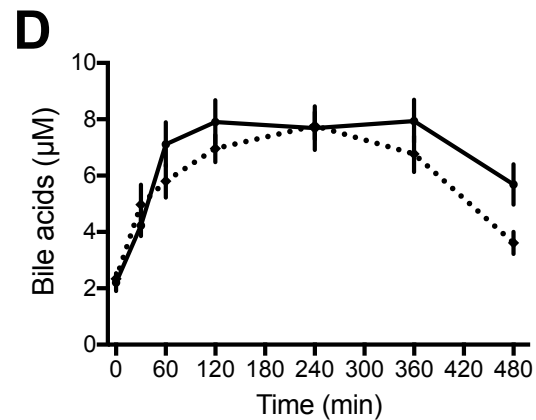
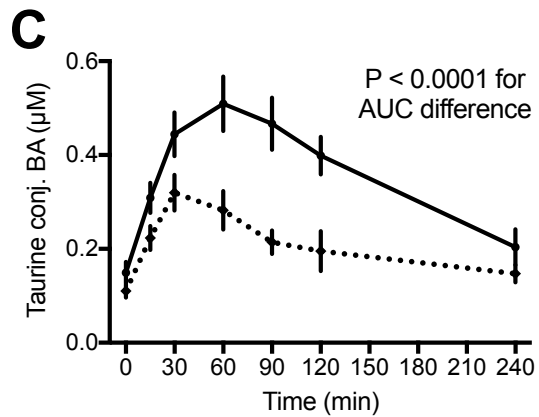
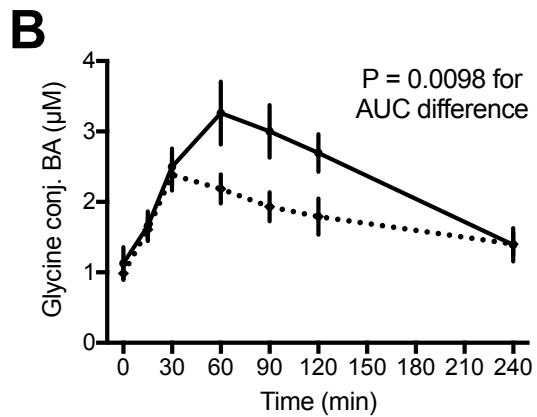
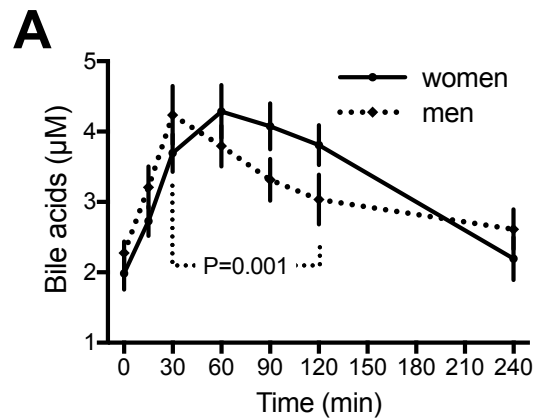
	<i>TIME</i>	<i>GENDER</i>	<i>genetic explained variation</i>	<i>microbiome explained variation</i>	<i>CYP27A1 (4)</i>	<i>CYP39A1 (24)</i>	<i>HSD3B7 (4)</i>	<i>SLC10A2 (4)</i>	<i>SLCO1A2 (9)</i>	<i>SLCO1B1 (30)</i>	<i>SLCO1B3 (24)</i>	<i>ABCC3 (63)</i>	<i>ABCA3 (4)</i>	<i>ABCG5 (10)</i>	<i>ABCG8 (6)</i>	<i>ABCC2 (8)</i>	<i>AMACR (63)</i>	<i>GPBAR1 (6)</i>	<i>EPHX1 (39)</i>	<i>UGT2B4 (49)</i>	<i>ATP8B1 (26)</i>	<i>NR0B2 (10)</i>	<i>Acidaminococcaceae</i>	<i>Alcaligenaceae</i>	<i>Bacteroidaceae</i>	<i>Bifidobacteriaceae</i>	<i>Christensenellaceae</i>	<i>Coriobacteriaceae</i>	<i>Enyspeliobacteriaceae</i>	<i>Lachnospiraceae</i>	<i>Peptostreptococcaceae</i>	<i>Porphyromonadaceae</i>	<i>Prevotellaceae</i>	<i>Rikenellaceae</i>	<i>Ruminococcaceae</i>	<i>Veillonellaceae</i>	TOTAL VARIATION EXPLAINED		
CA	2	1	39	21	0	3	1	5	1	1	3	9	0	0	2	0	2	6	2	1	3	0	0	0	0	0	0	0	0	5	1	12	2	0	0	0	0	1	63
CDCA	6	1	38	10	0	2	2	3	0	0	2	4	0	0	0	0	4	6	3	3	8	1	2	0	1	0	0	1	2	0	3	0	0	0	0	1	0	55	
DCA	4	0	54	11	0	0	0	3	0	3	1	6	2	2	2	4	8	2	14	1	1	5	2	1	2	1	0	2	0	1	0	0	1	0	0	1	69		
GCA	14	0	23	6	0	1	0	0	3	0	2	1	1	0	1	1	3	1	2	3	4	0	1	0	1	1	0	0	0	0	1	0	0	1	1	0	43		
GCDCA	21	1	18	8	0	1	0	0	0	0	2	0	2	0	0	0	3	2	3	0	3	2	1	0	2	1	0	4	0	0	0	0	0	0	0	0	0	48	
GDCA	10	1	50	2	0	0	1	0	1	3	2	2	3	0	3	1	2	0	29	1	1	1	0	0	0	0	0	0	0	0	0	0	0	0	0	1	1	63	
TCA	5	3	25	4	0	0	3	1	2	1	3	1	0	0	0	1	6	1	0	2	4	0	0	0	0	1	1	0	0	0	0	0	0	0	2	0	0	37	
TCDCA	13	5	24	6	0	0	2	3	1	0	2	2	0	1	0	0	4	2	2	1	2	2	1	0	0	0	1	4	0	0	0	0	0	0	0	0	0	48	
TDCA	7	4	50	3	0	0	0	2	0	1	3	2	2	1	2	1	3	1	31	1	0	0	0	0	1	0	0	0	0	0	0	0	0	0	1	1	64		

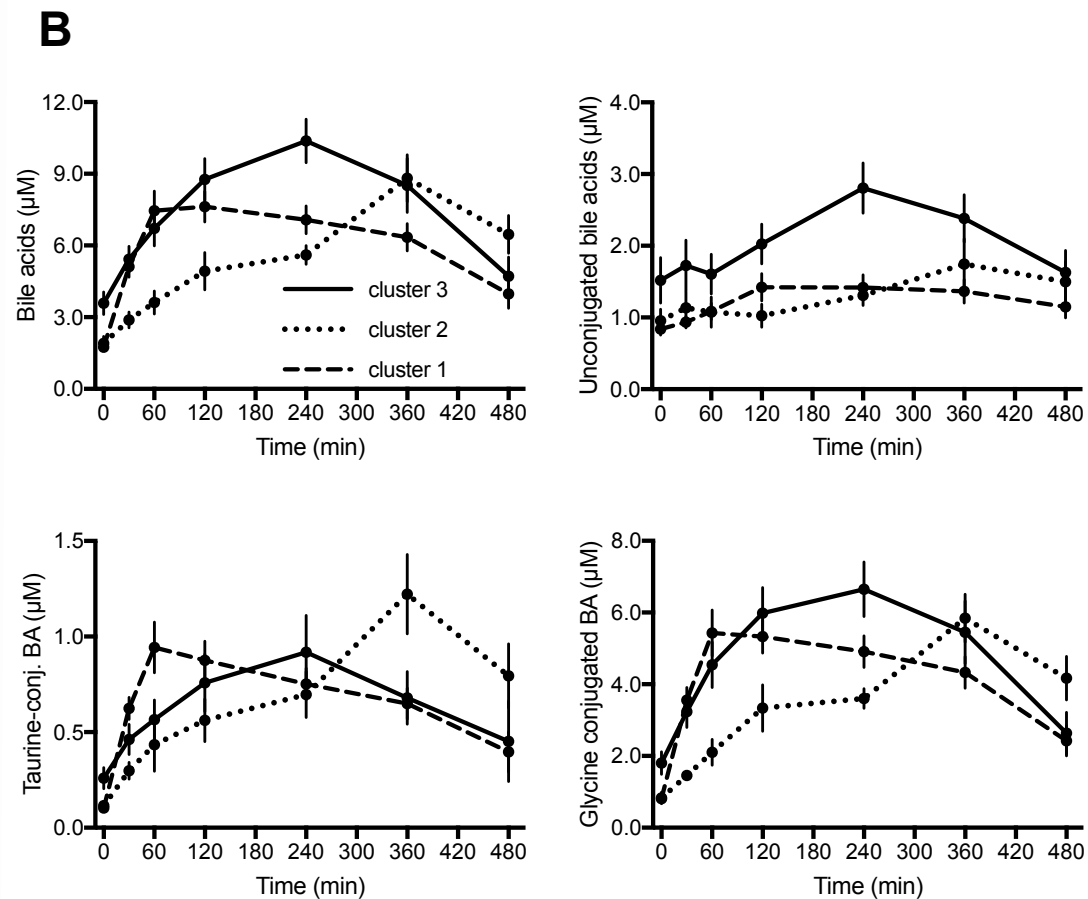
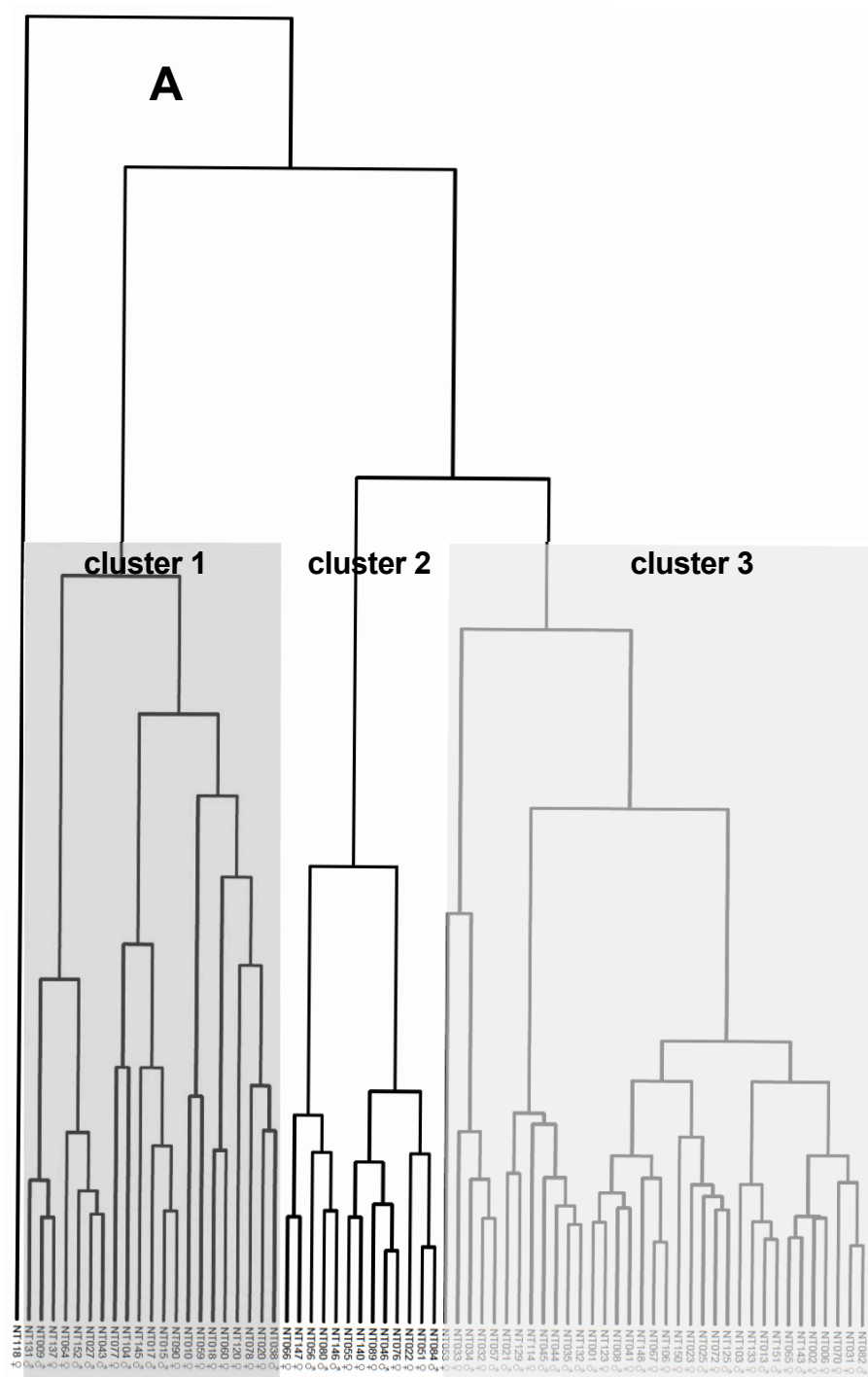


Bile acids (μM)



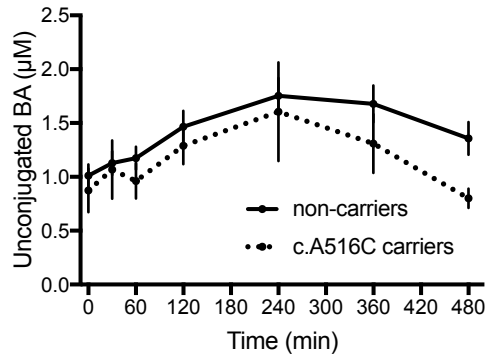
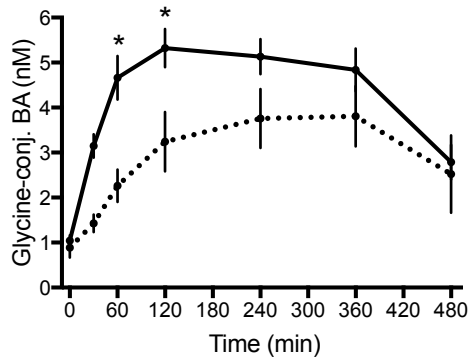
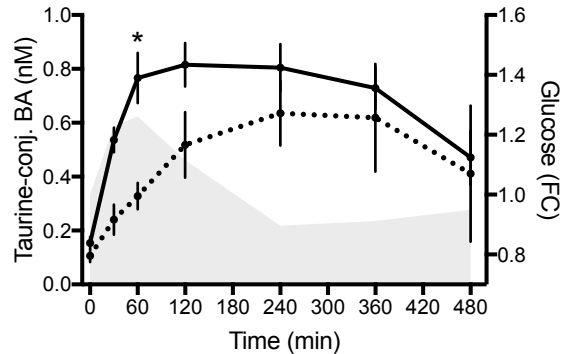
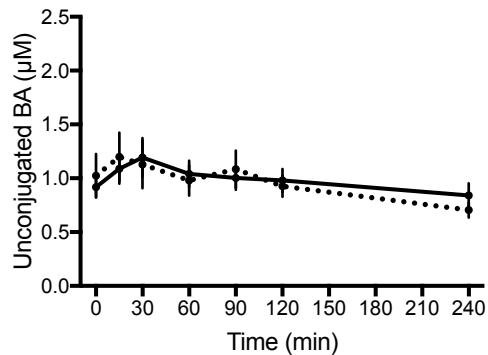
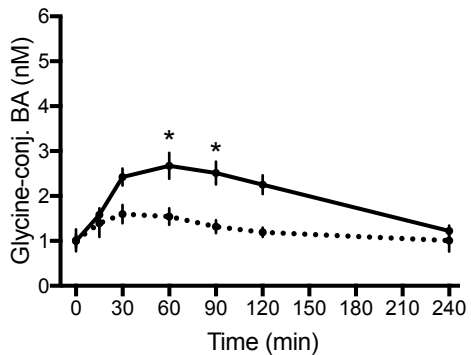
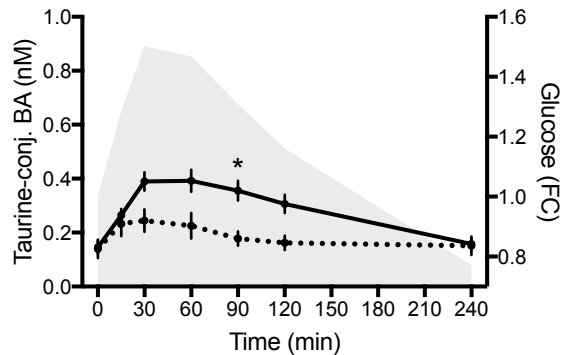


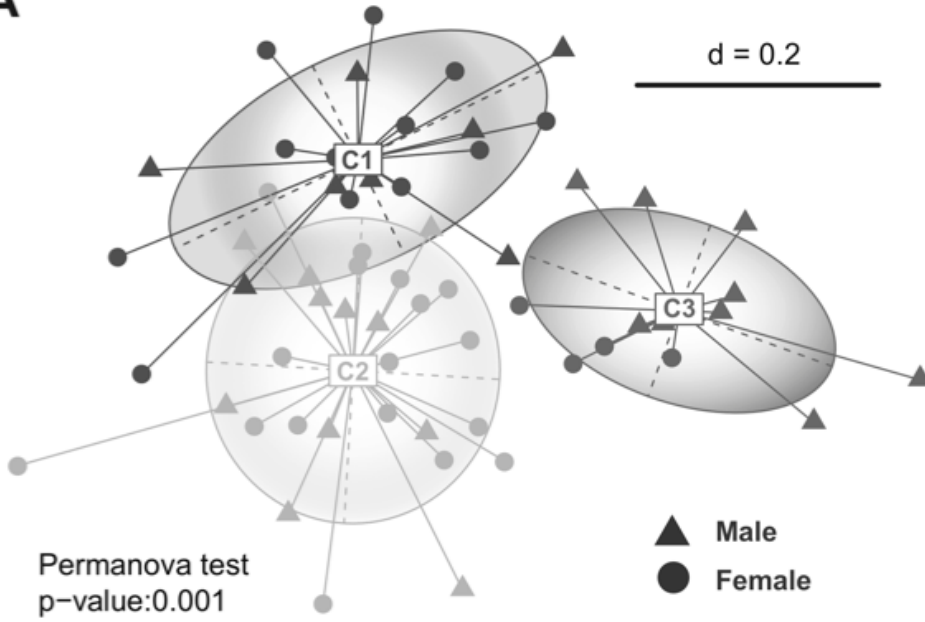
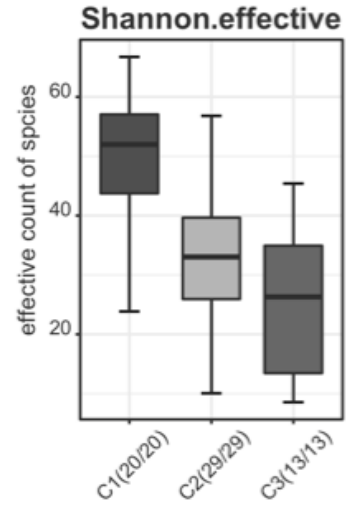




C

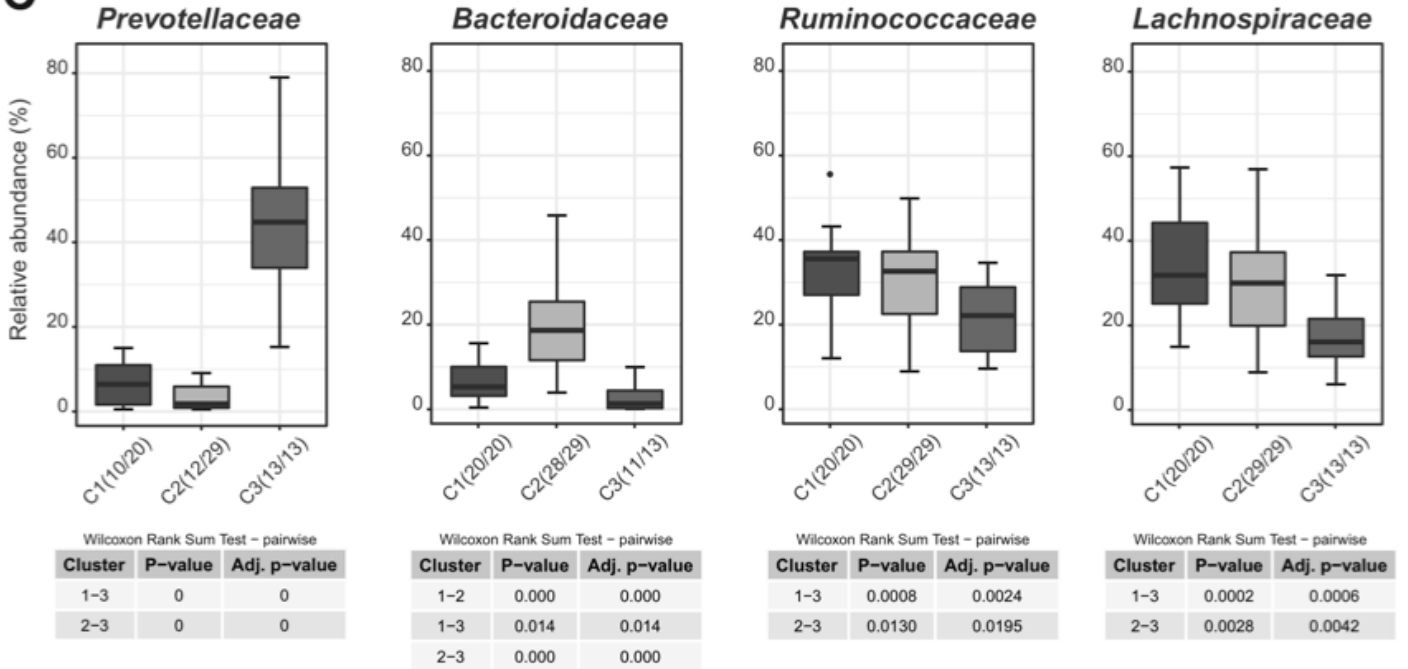
	total	unconj.	glycine-conj.	taurine-conj.
0	3≠1	3≠1	-	-
30	3≠2, 2≠1	3≠1	3≠2, 2≠1	-
60	3≠2, 2≠1	-	3≠2, 2≠1	3≠1, 2≠1
120	3≠2, 2≠1	3≠2, 3≠1	3≠2, 2≠1	-
240	3≠2, 3≠1	3≠2, 3≠1	3≠2, 3≠1	-
360	3≠1, 2≠1	3≠1	-	3≠2, 2≠1
480	2≠1	-	2≠1	2≠1

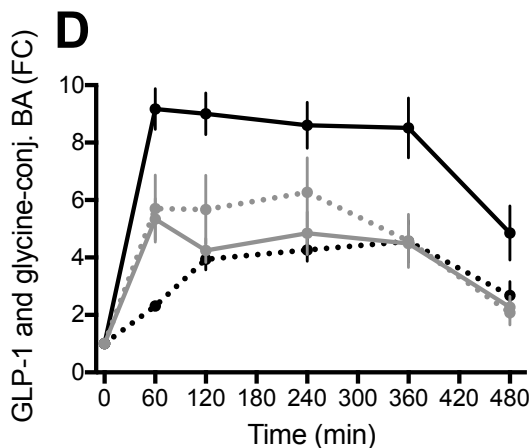
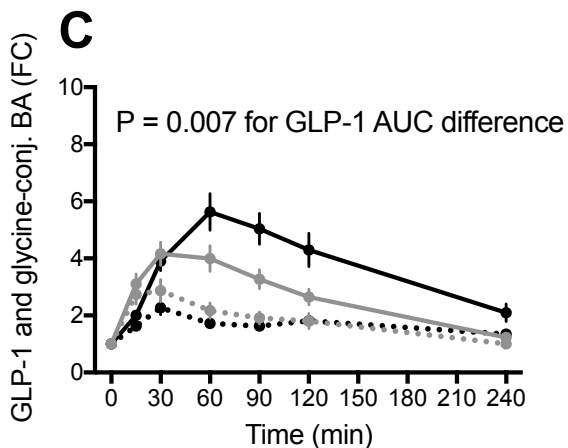
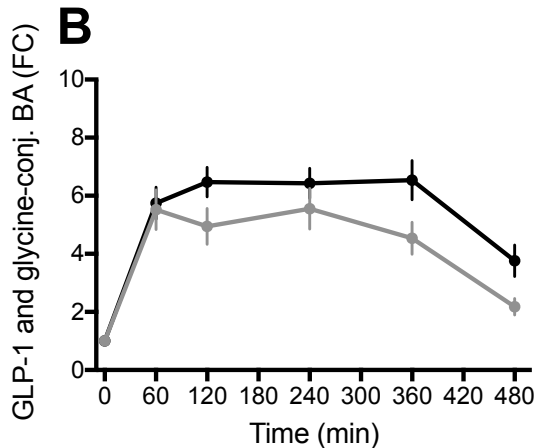
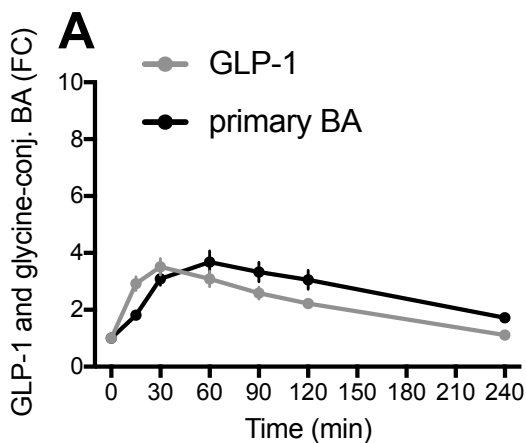
A**B****C****D****E****F**

A**B**

Wilcoxon Rank Sum Test - pairwise

Cluster	P-value	Adj. p-value
1-2	1e-04	2e-04
1-3	1e-04	2e-04

C



- GLP-1 in subjects with **low** post-prandial BA increase
- GLP-1 in subjects with **high** post-prandial BA increase
- BA in subjects with **low** post-prandial BA increase
- BA in subjects with **high** post-prandial BA increase