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Feeding gilthead sea bream with increasing dietary bacterial single cell protein level: Implication on growth, plasma biochemistry, gut histology, and gut microbiota

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24 **Feeding gilthead sea bream with increasing dietary bacterial single cell protein level:**
25 **implication on growth, plasma biochemistry, gut histology, and gut microbiota**

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36

37 **Abstract**

38 Bacterial single cell protein (SCP) is considered a promising circular protein ingredient
39 for aquafeed, due to the high protein content, and for the possibilities to grow them on
40 different substrates such as organic waste thus leading to low environmental footprint and
41 affordable production costs. Their use as raw material has been assessed in several farmed
42 species, however, research on Mediterranean ones is still scarce. Hence, a study was
43 undertaken to evaluate growth, plasma biochemistry, gut histology and gut microbiota
44 (GM) response of gilthead sea bream (*Sparus aurata*) fed diets with increasing levels of
45 bacterial SCP in comparison to a control without SCP. Three isonitrogenous and
46 isolipidic extruded diets (44% protein; 19% lipid) were formulated with different
47 bacterial SCP (derived from *Corinebacterium glutamicum*) level (10% SCP, SPC10; 15%

48 SCP, SCP15; 20% SCP, SCP20) to replace vegetable protein ingredients (total
49 replacement of soy protein concentrate and partial replacement of corn gluten), while a
50 control diet (CTRL) was formulated without SCP. Fish groups of 45 individuals (initial
51 weight: 75 g) were fed to visual satiation over 108 days. At the end of the trial there were
52 no significant differences on growth, feed intake, feed conversion rate, protein efficiency
53 ratio and protein apparent digestibility. Most of plasma parameters were found to be equal
54 for all treatments, except for those related to nucleic acids molecules degradation such as
55 phosphorus and urea which were higher in SCP10 and SCP20 compared to CTRL,
56 respectively. No morphological alterations were found in the intestines of any fish
57 analysed. Different responses of the overall GM structure in relation to the bacterial SCP
58 inclusion level were detected. Specifically, SCP exerted a positive effect on GM internal
59 diversity which increased at increasing dietary SCP inclusion level. In addition, SCP
60 inclusion lead to increase in the abundance of *Bacillus* spp, taxa which can potentially
61 support nutrition, immune system, and disease resistance. In conclusion, it seems feasible
62 to include up to 20% of SCP dietary level for gilthead sea bream without compromising
63 growth, feed efficiency and health parameters.

64

65 **Keywords**

66 Single cell protein, gilthead sea bream, growth, plasma biochemistry, gut microbiota

67

68 **Introduction**

69 Single cell protein (SCP) are dehydrated cells of unicellular organisms such as fungi,
70 bacteria, yeast and microalgae deriving from fermentation processes of biomass from
71 industry and agriculture (i.e. molasses, whey, starch, alkanes, hydrocarbons, celluloses,
72 ammonia, nitrate, natural gas) (Sharif et al., 2021). SCP are used for animal feed due to
73 many factors: high growth rate of microorganisms, the wide variety of substrates they can
74 metabolize, the low processing costs, and the production of derived nutrients and
75 functional molecules. They are mainly used in human or animal nutrition as protein
76 sources, due to high protein content (60–82% on dry matter), with a suitable amino acid
77 profile. In addition, the content of beneficial lipids, carbohydrates, vitamins and minerals
78 may promote their higher nutritional values in comparison to conventional protein
79 sources (Zepka et al., 2010; Aruna et al., 2017; Sharif et al., 2021). Moreover SCP also
80 requires low water demand compared to the plant sources and are not affected by
81 environmental conditions (Sharif et al., 2021). In aquaculture, SCP has been considered
82 as a protein replacement to standard protein commodities (i.e. fishmeal, FM and soy
83 products) and most of the studies carried out have shown positive impacts on increasing
84 growth (Guo et al., 2019).

85 Yeast SCPs, are a protein source rich in vitamins and micronutrients which can provide
86 several benefits such as enhancing the immune response, reducing stress, and modulating
87 gut microbiota (Rawling et al., 2019; Rimoldi et al., 2020; Ciji and Akhtar, 2021) . These
88 effects were tested on several fish species, such as Atlantic salmon, *Salmo salar* (Hansen
89 et al., 2021), rainbow trout, *Oncorhynchus mykiss*, gilthead seabream *Sparus aurata*, and
90 European seabass *Dicentrarchus labrax* (Agboola et al., 2021).

91 Microalgae SCP are a good crude protein source, (60 %) but is mostly used for
92 production of omega-3 fatty acids (EPA and DHA) and carotenoids (Glencross et al.,
93 2020). Many studies have been conducted on the health, immune response and
94 digestibility of microalgae SCP of several species such as rainbow trout (Zhang et al.,
95 2020), Atlantic salmon (Hart et al., 2021), gilthead sea bream (Carvalho et al., 2020), and
96 European sea bass (Messina et al., 2019).

97 Bacteria SCP are considered to be a promising protein source for aquafeed, due to its
98 ability to alter their composition according to different production setting. They have a
99 high content of raw proteins (80%), high growth rate, and they are able to grow on
100 different substrates such as organic waste and petrochemicals i.e. ethanol, methane,
101 methanol and nitrogen, syngas, CO₂ and H₂ (Delamare-Deboutteville et al., 2019; Jones
102 et al., 2020). Among the other previously mentioned SCP sources, minor attention has
103 been devoted to exploring the use of bacterial SCP as an aquafeed ingredient. Some
104 studies were performed on Atlantic salmon (Aas et al., 2006), rainbow trout (Hardy et al.,
105 2018), Nile tilapia *Oreochromis niloticus* (Smarason et al., 2019), Japanese yellowtail
106 *Seriola quinqueradiata* (Biswas et al., 2020), and African catfish *Clarias gariepinus*
107 (Adeoye et al., 2021). Most of them highlight the possibility to replace 5-20% FM in
108 rainbow trout, 30% FM in catfish, 4-36% FM in Atlantic salmon, and 50% FM in Nile
109 tilapia. To the best of our knowledge, only one study reported the application of dietary
110 bacterial SCP in Mediterranean fish species (Solé-Jiménez et al. 2021). The authors were
111 able to successfully replace 50% of FM using a commercial protein source made of
112 bacterial and processed animal proteins without compromising growth performance of
113 gilthead seabream.

114 In recent years, in the aquaculture sector significant amounts of dietary FM have been
115 successfully replaced with alternative ingredients mainly derived from commodity
116 agricultural crops proteins such as plant-based proteins including various legumes such
117 as soya bean meal. As consequence, the increasing demand for plant proteins for animal
118 feed production, has been also associated to environmental impact concerns over
119 deforestation, land-use displacement and eutrophication (Woodgate et al., 2022).

120 The aim of this study was to explore the efficacy of dietary inclusion level of bacterial
121 SCP to replace plant protein sources. Growth, plasma biochemistry, gut histology, and
122 gut microbiota during the on-growing of gilthead sea bream are investigated.

123

124 **Materials and methods**

125

126 *2.1 Experimental diets*

127

128 Three diets (44% protein; 19% lipid) were formulated to contain increasing level of
129 single cell protein (SCP, Gordini srl, Italy) from *Corinebacterium glutamicum* (10% SCP
130 SPC10, 15% SCP SCP15, 20% SCP SCP20) in order to replace vegetable ingredients
131 (total replacement of soy protein concentrate and partial replacement of corn gluten)
132 while a control diet (CTRL) without SCP and containing the same amount of FM was
133 used (Table 1). Diets were formulated with FM and a mixture of vegetable ingredients
134 currently used for sea bream in aquafeed (Parma et al., 2016). The diets were produced
135 via extrusion (pellet size = 3.0 mm) by SPAROS Lda (Portugal). Proximate composition
136 of the diets and amino acid composition of the SCP are reported in Tables 1-2.

137

138 *2.2 Fish and experimental conditions*

139

140 A growth trial was performed at the Laboratory of Aquaculture, Department of
141 Veterinary Medical Sciences of the University of Bologna (Cesenatico, Italy). Gilthead
142 sea breams were obtained from an Italian hatchery and acclimatized to the facilities for 7
143 days before the beginning of the trial. Forty-five fish per tank were randomly distributed
144 in twelve 500 L tanks. Each diet was administered to triplicate tanks over 108 days. Tanks
145 were provided with natural seawater and connected to a closed recirculating system
146 (overall water volume: 7000 L; Oxygen level $8.0 \pm 1.0 \text{ mg L}^{-1}$; Temperature $23 \pm 1.0 \text{ }^\circ\text{C}$,
147 Salinity 25 g L^{-1} , pH 7.8-8.0) according to Busti et al. (2020a). Fish were hand fed to
148 visual satiation twice a day (8:30, 16:00) for six days a week. Feeding procedures were
149 made to prevent any feed losses, however in cases of uneaten feed, pellets were collected,
150 dried overnight at 105°C , and weighted for overall calculation.

151

152 *2.3 Sampling*

153

154 Before each sampling procedures fish were anaesthetized (100 mg L^{-1}) or euthanised
155 (300 mg L^{-1}) by tricaine methanesulfonate MS-222 (Sigma-Aldrich). At the beginning
156 and the end of the trial individual fish weight was measured in each tank. The approximate
157 composition of the carcasses was determined at the beginning of the trial on a pooled
158 sample of 15 fish and on a pooled sample of 5 fish per tank at the end of the trial. Blood
159 from 5 fish per tank ($n=15$ fish per diet treatment) was collected at the end of the trial
160 from the caudal vein. Samples were then centrifuged ($3000 \times g$, 10 min, 4°C) and plasma
161 aliquots were stored at $-80 \text{ }^\circ\text{C}$ until analysis (Bonvini et al., 2018). On the same
162 specimens, three fish (12 fish for each experimental group) were used for histology. For

163 each fish, the gut was gently removed from the celom cavity and the anterior and middle
164 part of the intestine were fixed in buffered formalin and then processed for morphological
165 evaluation. Digesta content (n=5 fish per tank) from distal intestine was also individually
166 sampled at the end of the trial and immediately stored at -80°C for gut microbiota
167 investigation according to Parma et al. (2020).

168

169 *2.4 Digestibility assessment*

170

171 After the end of the trial, the remaining groups of fish were used to determine the
172 apparent digestibility coefficient (ADC) of dry matter and protein, by the indirect method
173 with diets containing yttrium oxide according to Busti et al. (2020b). Fish were fed
174 according to the different diets and after that at 8 h post-prandial fish were euthanised by
175 overdose of anaesthetic and faeces were collected after fish dissection and stripping distal
176 intestine. Faeces were then pooled (N=3) for each tank and kept at -20°C until analysis
177 for yttrium, dry matter and protein. ADC was calculated as follows: $\text{ADC} =$
178 $100 * (1 - (\text{dietary Y2O2 level} / \text{faecal Y2O2 level})) * ((\text{faecal nutrient} / \text{dietary nutrient}))$.

179 All experimental procedures were evaluated and approved by the Ethical-Scientific
180 Committee for Animal Experimentation of the University of Bologna, in accordance with
181 European directive 2010/63/UE on the protection of animals used for scientific purposes.

182

183 *2.5 Gut histology*

184

185 For each sample, 24 intestinal paraffin sections (both anterior and middle part) were
186 obtained. To avoid morphometric evaluations in serial sections and, consequently, to

187 analyse the same characteristics, the first, sixth, eighteenth and twenty-fourth sections
188 were processed and stained with haematoxylin and eosin. Since the mucosa of the anterior
189 intestinal tract showed very complex and branching folds, it was decided to evaluate the
190 absorbent surface using the binarization method. Binarization was done in a blind fashion
191 by 2 expert investigators. The intestinal sections were scanned with the Nikon DS-Qi1Nc
192 digital camera at 10X magnification, using NIS Elements software BR 4.20.01 (Nikon
193 Instruments Europe BV, Amsterdam, Netherlands) with an interactive tool, Scan Large
194 Image, suitable for subsequent image analysis. This tool acquires an image with an area
195 of interest that exceeds the camera's field of view, capturing a large image made up of
196 multiple image frames stitched by an automatic algorithm that cannot be loaded in one
197 piece. Automated Image Binarization was applied to area of each selected intestinal image
198 by means of the software NIS Elements software BR 4.20.01. Image Binarization is a
199 widely used method that allow distinguishing objects of interest from background.
200 Indeed, determines a grey threshold and assigns each pixel of a digital image to one class
201 (image objects) if its grey value is greater than the determined threshold and otherwise to
202 the other class (image background). Specifying correct threshold limits is a crucial
203 procedure of the automated image analysis. The point is to determine which pixels will
204 and which will not be included in the binary layer and thereby distinguish objects to be
205 analyzed from background. By threshold, its possible highlighted the absorbent surface
206 of the intestinal tract. In our case, using binarization we were able to separate the pixels,
207 which represent absorbent surface (brighter pixels) from those, that represent the rest of
208 the layers of the intestinal sections (Fig. 1A and B). The area of measurement can be
209 restricted by a user-defined region of interest ROI (Fig. 1C). ROI is a strong tool used
210 mainly to measure varying image intensity inside the ROIs or number of binary objects

211 inside each ROI, Object Count to restrict binary objects to areas of interest only (Fig. 1C
212 and D). Consequently, this allowed us to quantify the cellular absorbent surface covered
213 by villi. The measured obtained were expressed in square millimetres.

214

215 *2.6 Plasma biochemistry*

216

217 Glucose (GLU), urea, creatine, uric acid, total bilirubin, cholesterol (CHOL),
218 triglycerides (TRIG), high density lipoprotein (HDL), total protein (TP), albumin (ALB),
219 aspartate aminotransferase (AST), alanine transaminase (ALT), alkaline phosphatase
220 (ALP), creatine kinase (CK), lactate dehydrogenase (LDH), lactate (LAC), calcium
221 (Ca²⁺), phosphorus (P), potassium (K⁺) sodium (Na⁺), iron (Fe), chloride (Cl),
222 magnesium (Mg) were determined in the plasma using samples on an automated analyser
223 (AU 480; Olympus/Beckman Coulter, Brea, CA, United State). OSR (Olympus system
224 reagent) method was utilized to evaluate the reported variables according to Pelusio et al.
225 (2021). The ALB/globulin (GLOB), Na/K ratio and Ca x P were calculated.

226

227 *2.7 Calculations*

228

229 The following formulae were used to calculate different performance parameters:
230 specific growth rate (SGR) (% day⁻¹) = 100 * (ln FBW - ln IBW) / days (where FBW and
231 IBW represent the final and the initial body weights, respectively). Feed Intake (FI) (g kg
232 ABW⁻¹ day⁻¹) = ((100 * total ingestion) / (ABW)) / days (where average body weight,
233 ABW = (IBW + FBW) / 2. Feed conversion ratio (FCR) = feed intake / weight gain. Protein
234 efficiency rate (PER) = (FBW - IBW) / protein intake. Gross protein efficiency (GPE)

235 (%) = 100 * [(% final body protein * FBW) - (% initial body protein * IBW)] / total
236 protein intake fish. Economic conversion ratio (ECR) (€/kg fish⁻¹) = FCR * feed cost
237

238 *2.8 Proximate composition analysis*

239

240 Diets and whole body of sampled fish were analysed for an approximate composition
241 as reported in Parma et al. (2020). In brief, the moisture content was obtained by
242 observing the weight loss after drying the samples in a stove at 105 °C until a constant
243 weight was achieved. Crude protein was determined as total nitrogen (N) by using the
244 Kjeldahl method and multiplying N by 6.25. Total lipids were determined according to
245 Bligh and Dyer's (1959) extraction method. Ash content was estimated by incineration to
246 a constant weight in a muffle oven at 450 °C.

247

248 *2.9 Gut Bacterial Community DNA Extraction and Sequencing*

249

250 Total DNA was extracted and analysed from individual distal intestine content
251 obtained from 5 fish per tank (300 mg per fish) at the end of the trial, as previously
252 reported in Pelusio et al. (2021) and from 4 samples of the different diet using the DNeasy
253 PowerSoil Kit (Qiagen, Hilden, Germany). DNA was quantified with NanoDrop ND-
254 1000 (NanoDrop Technologies, Wilmington, DE) and stored at -20 °C until further
255 processing. To target the transient bacterial community, the amplification of the V3-V4
256 hypervariable regions of the 16S rRNA gene was carried out using the 341F and 785R
257 primers carrying Illumina overhang sequencing adapters and 2 × KAPA HiFi HotStart
258 ReadyMix (KAPA Biosystems). The thermal cycle was performed as already described

259 by Pelusio et al. (2021) using 30 amplification cycle. PCR products were purified, and
260 indexed libraries were prepared following Illumina protocol “16S Metagenomic
261 Sequencing Library Preparation”. Libraries were normalized to 4 nM and pooled. Pooled
262 libraries were denatured with 0.2 N NaOH and diluted to 6 pM with 20% PhiX control.
263 Sequencing was performed on Illumina MiSeq platform using 2 x 250 bp paired-end
264 protocol according to the manufacturer’s instructions (Illumina, San Diego, CA). At the
265 end of the sequencing process, raw sequences were processed combining PANDAseq and
266 QIIME2 pipelines (Bolyen et al., 2019; <https://qiime2.org>). High-quality reads, obtained
267 after a filtering step for length (minimum/maximum = 350/550 bp) and quality with
268 default parameters, were cleaned using DADA2 (Callahan et al., 2016) and clustered into
269 amplicon sequence variants (ASVs) using VSEARCH algorithm (Rognes et al., 2016).
270 Taxonomy was assigned using RDP classifier against SILVA database (Quast et al.,
271 2013). The 4 feed samples were discarded for subsequent analysis due to the high number
272 of “unassigned taxa”. Three different metrics were used to evaluate internal ecosystem
273 diversity (alpha-diversity) – Faith’s Phylogenetic Diversity (PD_whole_tree), Chao1
274 index for microbial richness, and number of observed ASVs. UniFrac distances were
275 computed to estimate inter-sample ecosystem diversity (beta-diversity) and used as input
276 for Principal Coordinates Analysis (PCoA).

277

278 *2.10 Statistical analysis*

279

280 All data are presented as mean \pm standard deviation (SD). A tank was used as the
281 experimental unit for analysing growth performance and a pool of five fish were
282 considered the experimental unit for analysing carcass composition. Individual fish were

283 used for analysing plasma biochemistry. Data on growth, nutritional indices, apparent
284 digestibility, plasma biochemistry were analysed by a one-way analysis of variance
285 (ANOVA) with Tukey's post hoc test. The normality and/or homogeneity of variance
286 assumptions were validated for all data preceding ANOVA. The gut morphometric
287 evaluation was expressed as mean \pm SD. The data obtained was analysed by *t*-test. The
288 differences among treatments were considered significant at $p \leq 0.05$. All microbiota
289 analysis and respective plots were produced using R software ([https://www.r-](https://www.r-project.org/)
290 [project.org/](https://www.r-project.org/)) with “vegan” (<http://www.cran.r-project.org/package=vegan/>), “Made4”
291 (Culhane et al., 2005) and “stats” packages ([https://stat.ethz.ch/R-manual/R-](https://stat.ethz.ch/R-manual/R-devel/library/stats/html/00Index.html)
292 [devel/library/stats/html/00Index.html](https://stat.ethz.ch/R-manual/R-devel/library/stats/html/00Index.html)). Data separation was tested by a permutation test
293 with pseudo-F ratios (function “Adonis” in “vegan” package). When required, Wilcoxon
294 and Kruskal–Wallis test were used to assess significant differences in alpha diversity and
295 taxon relative abundance between groups. *P*-values were adjusted for multiple
296 comparisons using the false discovery rate (FDR) (function *p.adjust* in the “stats”
297 package), and a *p*-value ≤ 0.05 was considered statistically significant, while a *p*-value
298 between 0.05 and 0.1 was considered as a trend.

299

300 **3. Results**

301

302 *3.1 Growth, nutritional indices, and protein digestibility*

303

304 Growth performance parameters are reported in Table 3. No significant differences on
305 FBW, weight gain and SGR were detected between dietary treatments. Similarly, no
306 significant differences on FI, FCR, ECR and survival were also observed (Table 3).

307 Data on body composition, nutritional indices and apparent digestibility are shown in
308 Table 4. Protein body content was lower in the SCP15 diet compared to the CTRL diet
309 while no differences on moisture, lipid and ash content were detected between treatments.
310 GPE was lower in SCP15 compared to the CTRL diet while no significant differences on
311 PER and apparent digestibility of dry matter and protein was detected. Concerning
312 somatic indices, CF was lower in the SCP20 diet compared to SCP10, while no significant
313 differences between treatments were detected for HSI and VSI.

314

315 3.2 *Gut histology*

316

317 Anterior and middle intestine were lined by a tunica mucosa constituted by epithelium
318 and lamina propria forming folds/villi along all tracts. The intestinal mucosa of the
319 anterior intestine tract was organized in mucosal folds formed by tall primary everting
320 constituted by the mucosa and submucosa: from their main axis, secondary everting of
321 epithelium and *lamina propria* formed other folds/villi-like. Along the entire length of
322 the middle intestine, the complex folds were rare/absent than to anterior tract: most of
323 these folds resemble normal villi. No inflammation features were observed, such as villi
324 shortening and nuclear positioning disparity. Regarding morphometric analysis no
325 significant differences in both anterior and middle intestinal tracts were observed (Table
326 5).

327

328 3.3 *Plasma biochemistry*

329

330 Plasma parameters are shown in Table 6. Urea was higher in SCP20 compared to the
331 other groups. TRIG were higher in SCP15 compared to CTRL. HDL was higher in SCP20
332 compared to CTRL. Ca^{+2} and Cur Ca^{+2} were significantly higher in the SCP20 diet than
333 in the CTRL diet, while P and CaxP were higher in SCP10 compared to CTRL. No
334 significant differences among treatments were detected for GLU, CREA, Uric Ac, Tot
335 Bil, CHOL, TP, ALB, AST, ALT, ALP, CK, LDH, LACT, K^+ , Na^+ , Fe, Cl, Mg,
336 ALB/GLO, Na/K.

337

338 *3.4 Faecal bacterial community profiles related to dietary groups*

339

340 The 16S rRNA gene sequencing was performed on a total of 60 distal intestine content
341 samples, yielding 401'841 high-quality reads (mean \pm SD, 6'182 \pm 2'574) and clustered
342 into a total of 6'252 ASVs. To assess whether the treatments with increasing SCP could
343 exert an effect on the gut bacteria community during the growth process of gilthead
344 seabream, the gut microbiota (GM) was analysed for each dietary group at the end of the
345 trial. The GM variations between samples (beta-diversity) were assessed by the Principal
346 Coordinates Analysis (PCoA) based on Unweighted UniFrac distances, with the taxa
347 most explaining sample segregation being superimposed on the bidimensional space. In
348 addition, the gut microbial community internal diversity was represented with three
349 different metrics for each dietary group: PD_whole_tree, Chao1 index, and
350 observed_ASVs. According to the findings (Figure 2A), all SCP groups showed a
351 significant variation compared to the control group, in terms of overall GM composition
352 ("pairwise Adonis permutation test", $p < 0.01$). Focusing on dietary group, a significant
353 overall GM composition variation between all dietary groups was observed (SCP10 vs

354 SCP15, SCP10 vs SCP20, SCP 15 vs SCP20; “pairwise Adonis permutation test”, $p <$
355 0.01). As for internal ecosystem diversity, SCP10 dietary group showed a significant
356 reduction of the alpha-diversity compared to control group in all the 3 metrics considered
357 (SCP10 vs CTRL; Wilcoxon rank-sum test, $p = 0.005$) (Figure 2B). The SCP15 group
358 showed a significant reduction of the microbial internal ecosystem diversity compared to
359 control group only when considering PD_whole_tree metric, (SCP15 vs CTRL;
360 Wilcoxon rank-sum test, $p = 0.007$) (Figure 2B). On the contrary, SCP20 group showed
361 a significant increase of alpha-diversity, as for chao1 index and observed_ASVs metrics,
362 when compared to the control group (SCP20 vs CTRL; Wilcoxon rank-sum test, $p < 0.05$)
363 (Figure 2B). When considering only the different dietary groups, a significant increase of
364 microbial internal ecosystem diversity was observed associated with the increase of SCP
365 concentration, in chao1 index and observed_ASVs metrics (Figure 2B) (SCP10 vs
366 SCP15, SCP10 vs SCP20, SCP 15 vs SCP20; Wilcoxon rank-sum test, $p = 0.005$, $p =$
367 0.0002 , $p = 0.02$, respectively). To further assess the GM composition of gilthead sea
368 bream fed with different SCP concentrations, the overall composition at different
369 phylogenetic levels was investigated, as reported in Figure 3: at phylum (Figure 3A) and
370 family level (Figure 3B). More specifically, the most abundant phylum and the most
371 represented families in the gilthead sea bream GM of the 4 experimental groups were
372 showed in the Table 7. Moving to a lower taxonomic level, some compositional
373 differences were observed at genus level among dietary groups (Wilcoxon rank-sum test
374 $p < 0.05$) (Figure 4). According to the data, the relative abundance of *Bacillus*,
375 *Escherichia-shigella* and *Oceanobacillus* genera was significantly lower in CTRL group
376 compared to fish fed with different SCP concentrations (Wilcoxon $p < 0.05$). On the other
377 hand, the relative abundance of [*Eubacterium*] *coprostanoligenes* group, *Weissella*,

378 *Ruminococcaceae* UCG-10, *Ruminococcaceae* UCG-013, *Ruminococcaceae* UCG-014,
379 *Ruminococcaceae* UCG-005, *Christensenellaceae* R-7 group genera was generally
380 significant lower in fish fed with different SCP concentrations compared to control group
381 (Wilcoxon $p < 0.05$). With a focus on *Staphylococcus* genus, a direct effect of the SCP
382 concentration on its relative abundance was observed. More specifically, a significantly
383 higher abundance of *Staphylococcus* genus in fish fed with SCP10 diet compared to
384 control group (Wilcoxon, $p = 0.029$) was reported. Higher SCP concentrations were
385 instead associated with a significant reduction of the relative abundance of this genus
386 compared to SCP10 group, thus showing a significant relative abundance reduction in
387 fish fed with SCP15 diet compared to SCP10 diet (Wilcoxon, $p = 0.04$). SCP20 group
388 showed a significant reduction of the relative abundance of *Staphylococcus* genus
389 compared to both SCP15 and SCP10 groups (Wilcoxon, $p = 0.0004$, $p = 0.0005$,
390 respectively) and a tendency in abundance reduction compared to CTRL group
391 (Wilcoxon, $p = 0.09$) (Figure 4)

392

393 **Discussion**

394

395 The growth parameters (FBW, WG, SGR, FCR and FI) observed during the trial,
396 showed similar results regardless the level of SCP. Focusing on the feed intake, the
397 absence of significant differences has a positive implication considering its correlation
398 with appetite and palatability of feed. Most of the substances that increase the
399 attractiveness and palatability of feed in fish are characterized by low molecular weight
400 including nitrogenous and amphoteric components, amino acids, betaines, and
401 nucleotides. High nucleotide content derived from bacterial SCP could enhance

402 palatability and feed intake. It is recognized that nucleotides may act as a taste enhancer
403 specifically due to inosine and inosine monophosphate which were identified as a feeding
404 stimulant (Gamboa-Delgado et al., 2018; Hossain et al., 2020). This could be particularly
405 relevant when high level of plant protein sources is used to replace FM since they contain
406 significantly less nucleotides and the present of plant anti-nutritional factors may interfere
407 with palatability. Our results are in contrast with previous studies conducted on rainbow
408 trout (Hardy et al., 2018) and black sea bass (Chen et al., 2020), proving that the
409 replacement of soy protein with bacterial SCP could reduce FI due to low palatability.
410 The authors postulated that the source of bacterial protein meal and the process by which
411 it was dried or the presence of flavour compounds could have been responsible for low
412 palatability. The absence of statistical difference on final body weight, weight gain and
413 SGR, suggests that the inclusion of bacterial SCP to up to 20% could replace soy protein
414 concentrate and corn gluten without affecting growth performance. In Atlantic salmon
415 and rainbow trout, up to 30% replacement of soy protein with SCP did not affect animal
416 growth (Romarheim et al., 2011; Hardy et al., 2018). This is in accordance with a study
417 conducted on Nile Tilapia by Maulu et al. (2021), providing that dietary increasing level
418 of *Clostridium autoethanogenum* protein could replace up to 20% of soybean meal
419 improving growth performance. However, lower body protein content was observed in
420 SCP15. Lower protein content in SCP15 was also reflected in the lower values achieved
421 for protein efficiency (GPE) in the same treatment. However, differences in protein
422 efficiency were minimal and were not reflected in the FCR obtained. Overall, the absence
423 of growth differences, FCR, protein digestibility, and PER, suggest an optimal
424 digestibility, nutrients absorption, and utilization of bacterial SCP in gilthead sea bream
425 up to 20% of dietary inclusion level. Concerning somatic indices, no differences were

426 evaluated in HSI and VSI, assuming no effect on assimilation and distribution of nutrients
427 within the animal body's tissues.

428 Most of the plasma biochemistry results, such as TP, ALB, Glu, TC, and ALT, did not
429 show any statistical differences, demonstrating that SCP could guarantee optimal
430 nutrition and general health status (including liver health status as indicated by AST and
431 ALT) in agreement with previous plasma values on this species and with the results
432 obtained in Nile Tilapia fed bacterial SCP (Parma et al., 2020; Maulu et al., 2021).
433 However, Ca^{+2} and $CurCa^{+2}$ values were higher in SCP20 and, CaxP was higher in SCP10
434 compared to control diet. Differences when compared to the standard calcium values
435 could represent a stress indicator, causing imbalance plasma ion level and increasing
436 plasma osmolality (Mancera et al., 2002; Mateus et al., 2017). Despite this, calcium
437 values shown in this work, are in line with previous works of the same species and within
438 values of healthy fish (Peres et al., 2013; Pelusio et al., 2021). It is also worth highlighting
439 the higher value of plasmatic urea in SCP20. In agreement with Oliva-Teles et al. (2006)
440 high levels of SCP can lead to high levels of non-protein nitrogen content, such as nucleic
441 acid, mostly represented by RNA. Ammonia is the final product of pyrimidines
442 catabolism, which are one of the main components of RNA molecules. Even the higher
443 values of inorganic phosphorus detecting in SCP10 could be related to the degradation of
444 RNA molecules, even if this trend was not observed at the higher inclusion level (SCP15,
445 SCP20) tested. Plasma triglyceride and HDL were higher in the fish fed SCP15 and
446 SCP20 compared with the control diet, respectively. According to Maulu et al. (2021) the
447 inclusion of 200g kg of *Clostridium autoethanogenum* bacterial SCP increased plasma
448 triglycerides and cholesterol in Nile Tilapia. Although it is not clear how SCPs could
449 affect this process, the authors suggested an improvement in lipid metabolism and a role

450 of bacterial SCP in glucolipid metabolism to maintain whole-body energy homeostasis
451 through the adenosine monophosphate-activated protein kinase (AMPK) signalling
452 pathway. Interestingly, dietary nucleotide inclusion also increased blood triglycerides in
453 red sea bream *Pagrus major*, but further research is needed to illustrate lipid transport in
454 fish administered with nucleotides (Hossain et al., 2016a; Hossain et al., 2020).

455 Histology is considered a valid method for evaluating aquafeed ingredients since
456 several raw materials (mainly of vegetable origin) are known to induce morphological
457 changes thus altering the processes of nutrient digestion, absorption and pathogen
458 resistance (Rey et al., 2020). Gut histology revealed no histopathological changes of SCP
459 in the intestines of the animals examined indicating that SCP could replace soy derived
460 proteins, without altering the anatomic structure of the intestine. The literature is lacking
461 regarding the effects of bacterial SCP-enriched diets on gut histology in fish species.
462 However, dietary nucleic acid supplementation has a positive influence on intestinal
463 morphology such as increased enterocyte height and compensatory of intestinal
464 morphology damage. This is due to high inclusion of alternative vegetal protein that were
465 reported in several fish species (Hossain et al., 2016b; Hossain et al., 2020). It is worth to
466 mention that the level of soybean meal used in the present study in all the treatments was
467 within standard practical levels, which are known to not induce intestinal inflammatory
468 process in this species, as reported by Bonaldo et al. (2008) and Parma et al. (2016).

469 A growing number of researchers have addressed the study of gut microbiota in fish
470 species of commercial interest, since it is recognized as a powerful tool for assessing
471 digestive condition and gut health. In gilthead sea bream the inclusion of different protein
472 ingredients such as soy, insect, yeast and eggs peptide, have recently shown potential for
473 a GM reconfiguration (Parma et al., 2016; Antonopoulou et al., 2019; Rimoldi et al., 2020;

474 Naya-Català et al., 2021a, 2021b). According to our findings, GM was dominated by
475 Firmicutes, at phylum level while *Staphylococcaceae*, *Bacillaceae*, *Lactobacillaceae*,
476 *Leuconostocaceae* were the most represented taxa at family level. These data are in
477 agreement with previous findings on faecal GM of gilthead sea bream fed practical
478 aquafeed ingredients. Firmicutes and lactic acid bacteria (LAB) have been associated to
479 vegetal ingredients and considered a beneficial taxa able to promote nutrient digestion
480 and counteract pathogen invasion (Parma et al., 2016; Parma et al., 2020; Panteli et al.,
481 2021). Different responses of the overall GM structure in relation to the bacterial SCP
482 inclusion level were detected as evidenced by a significant separation in the PCOA
483 analyses of all the SCP inclusion level compared to the control diet. In addition, dietary
484 inclusion level exerted an effect on GM internal diversity which increased along with
485 higher SCP levels. In agreement with our findings, in gilthead sea bream the replacement
486 of 50 and 100% of FM with a mix of processed animal proteins (PAPs) and bacterial SCP,
487 leads to increased in alpha diversity indexes (Solé-Jiménez et al., 2021). Increase in GM
488 diversity may have positive implications for gut health due to increased competition
489 against opportunistic pathogens (Parma et al., 2020; Apper et al., 2016). In addition, a
490 wider range of bacteria supported by a higher diversity may promote a more diverse
491 number of host functions (Solé-Jiménez et al., 2021). However, low level of SPC (SCP10)
492 lead to a lower alpha diversity indexes compared to the control diet which further supports
493 the SCP dose effects on microbial diversity. In agreement with our study, a low dietary
494 inclusion level (equal to 5%) of autolysed yeast tended to reduce gut microbial alpha
495 diversity compared to a vegetable-based control diet in gilthead sea bream (Rimoldi et
496 al., 2020). In addition, the inclusion of brewer's yeast hydrolysate at 0.1 and 0.2% reduced
497 microbial diversity in largemouth bass (*Micropterus salmoides*) (Zhou et al., 2018). The

498 authors postulated a direct effect of specific molecules in the yeast such as nucleotides,
499 mannan oligo saccharide and β -glucan which could inhibit or promote specific bacterial
500 taxa (Zhou et al., 2018). In line with this hypothesis more recently Song et al. (2022)
501 observed in *Litopeneus vannamei* an unexpected decrease of gut microbiota diversity
502 when the guanosine 5'-monophosphate nucleotide was supplemented at 0.1% in diets
503 containing fermented soy in comparison to a non-supplemented diet. Concerning the
504 specific gut microbiota compositional changes, SCP inclusion led to increased *Bacillus*,
505 *Escherichia-shigella* and *Oceanobacillus*. In addition, taxa belonging to *Clostridiaceae* 1
506 family were also responsible for the separation between SCP treatments and control diet.
507 Interestingly, sea bream fed 5% dietary inclusion level of autolysed dried yeast in low
508 FM diet showed an enrichment in *Bacillales* and *Clostridiales* as compared to a control
509 vegetable-based diet (Rimoldi et al., 2020). *Bacillus* is one of the most important
510 beneficial taxa in fish species, which can make a positive contribution to nutrition, to the
511 immune system, and to disease resistance (Busti et al., 2020b; Soltani et al., 2019). In
512 particular *Bacillus spp.* showed growth and feed digestibility improvement mediated by
513 the production of exogenous enzymes (protease, lipase, phytase, chitinase), by the
514 degradation of plant-derived anti-nutritional factors, and by increasing nutrient levels
515 through microbial synthesis of essential bio-molecules (i.e. amino acids, fatty acids, and
516 vitamins). Against pathogens, bacteriocins from *Bacillus spp.* possess a broader spectrum
517 of inhibition that may include Gram-negative and Gram-positive bacteria of genera of
518 *Aeromonas*, *Edwardsiella*, *Streptococcus*, *Pseudomonas*, and *Vibrio*. In addition, oral
519 administration of *Bacillus* strains as probiotics has increased immune parameters (Ringo
520 et al., 2020). This includes lysozyme, phagocytosis, nitric oxide, bactericidal activity,
521 immune genes expression, humoral skin mucus parameters (Soltani et al., 2019) or may

522 affect the immune system by decreasing inflammation via the up-regulated secretion of
523 anti-inflammatory cytokines (Busti et al., 2020b). Overall these effects of SCP on GM
524 could partially explain the maintenance of the optimal productive results achieved under
525 all SCP dietary inclusion level.

526

527 **Conclusion**

528

529 In conclusion, the results of feed intake, growth, feed utilization and gut histology
530 indicate that bacterial SCP from *Corinebacterium glutamicum* can be successfully
531 incorporated up to 20% in practical aquafeed diets to reduce vegetable protein ingredients
532 (total replacement of soy protein concentrate and partial replacement of corn gluten)
533 without any negative effects on growth, protein utilization and gut health during the on-
534 growing phase of gilthead sea bream. The inclusion of SCP at each level tested,
535 determined a shift in the gut microbiota structure promoting taxa such as *Bacillus spp*,
536 which is considered one of the most important beneficial taxa in fish species.

537

538 **Author's contributions**

539 Conceptualization A.B., L.P., S.F., P.P.G.; Methodology A.M., A.B., L.P., S.F., D.S.,
540 M.C., M.M., G.L., P.C., A.D.M., F.D.; Investigation A.M., A.B., L.P., S.F., D.S., M.C.,
541 M.M., G.L., P.C., F.D., A.D.M.; Writing-original draft preparation A.M., D.S., L.P.;
542 Writing-review and Editing A.M., A.B., L.P., S.F., D.S., M.C., M.M., G.L., P.C., F.D.
543 All authors have read and agreed to the published version of the manuscript.

544

545 **Data availability**

546 All data are available in this manuscript

547

548 **Declaration of Competing Interest**

549 The authors claim that there is no conflict of interest

550

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556

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Table 1. Ingredients and proximate composition of the experimental diets

	CTRL	SCP 10	SCP 15	SCP 20
Ingredients, % of the diet				
Fishmeal Super Prime (Peruvian)	18.0	18.0	18.0	18.0
Soy protein concentrate	17.0	8.00	4.00	0.00
Corn gluten	16.8	11.0	8.00	5.50
Soybean meal 48	14.2	12.8	12.4	12.4
Wheat meal	7.60	8.10	8.40	9.00
Fish oil	4.00	4.00	4.00	4.00
Salmon oil	12.0	12.0	12.0	12.0
*Vit & Min Premix INVIVO 1%	1.00	1.00	1.00	1.00
MCP	0.60	0.40	0.30	0.30
L-Lysine	0.10	0.27	0.40	0.50
DL-Methionine	0.04	0.09	0.11	0.11
L-Taurine	0.05	0.06	0.06	0.06
Sunflower meal concentrate	8.61	14.28	16.33	17.13
#NT 70 Gordini	-	10.0	15.0	20.0
Yttrium oxide	0.01	0.01	0.01	0.01
Proximate composition, % on a wet weight basis				
Moisture	6.55	6.78	6.85	6.91
Protein	44.41	44.13	44.44	44.59
Lipid	19.01	18.83	19.04	18.73
Ash	7.12	7.12	7.20	7.33
Gross energy MJ kg ⁻¹	22.41	22.42	22.42	22.42

*Vitamins and mineral premix (IU or mg kg⁻¹ diet; Invivo NSA, Portugal); DL-alpha tocopherol acetate, 200 mg; sodium menadione bisulphate, 10 mg; retinyl acetate, 16,650 IU; DL-cholecalciferol, 2000 IU; thiamine, 25 mg; riboflavin, 25 mg; pyridoxine, 25 mg; cyanocobalamin, 0.1 mg; niacin, 150 mg; folic acid, 15 mg; L-ascorbic acid monophosphate, 750 mg; inositol, 500 mg; biotin, 0.75 mg; calcium panthotenate, 100 mg; choline chloride, 1000 mg; betaine, 500 mg; copper sulphate heptahydrate, 25 mg; ferric sulphate monohydrate, 100 mg; potassium iodide, 2 mg; manganese sulphate monohydrate, 100 mg; sodium selenite, 0.05 mg; zinc sulphate monohydrate, 200 mg.

Single cell protein, SCP from *Corinebacterium glutamicum*

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Table 2. Amino acid composition, g/100g of the bacterial single cell protein (NT 70 Gordini)

Aspartic acid (including Asparagina)	5.48
Glutamic acid (including Glutamine)	7.50
Hydroxyproline	<0.1
Serine	2.26
Glycine	4.15
Histidine	1.18
Arginine	3.36
Threonine	12.6
Alanine	4.16
Proline	2.11
Tyrosine	2.20
Valine	3.26
Methionine	1.36
Isoleucine	2.85
Leucine	4.75
Phenylalanine	2.54
Lysine	3.05
Tryptophane	0.915
Cysteine	0.526

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Table 3. Growth performance measured in gilthead seabream fed increasing dietary single cell protein levels.

	CTRL	SCP10	SCP15	SCP20	<i>P</i> - value
IBW	75.1 ± 2.67	75.5 ± 0.95	75.0 ± 0.56	75.3 ± 2.02	0.98
FBW	216.3 ± 11.41	216.6 ± 5.95	212.2 ± 2.48	214.6 ± 3.69	0.831
WG	141.2 ± 8.74	141.2 ± 5.95	136.9 ± 2.69	139.3 ± 4.46	0.776
SGR	0.98 ± 0.02	0.98 ± 0.01	0.96 ± 0.01	0.97 ± 0.03	0.653
FCR	1.29 ± 0.04	1.28 ± 0.05	1.35 ± 0.01	1.35 ± 0.05	0.122
FI	1.16±0.03	1.16±0.04	1.20±0.02	1.22±0.03	0.124
Survival	97.80 ± 2.2	96.30 ± 1.3	98.5 ± 2.6	97.80 ± 0.00	0.531
ECR	1.63 ± 0.06	1.53 ± 0.06	1.58 ± 0.02	1.54 ± 0.05	0.136

Data are given as the tanks mean (n=3) ± SD. In each line, different superscript letters indicate significant differences among treatments ($P \leq 0.05$).

IBW= Initial body weight (g).

FBW = Final body weight (g).

WG = Weight gain (g).

SGR = Specific growth rate (% day⁻¹) = 100 * (ln FBW- ln IBW) / days.

FCR = Feed conversion rate = feed intake / weight gain.

FI = Feed intake (g kg ABW⁻¹ day⁻¹) = ((100*total ingestion)/(ABW))/days).

ABW = average body weight = (IBW + FBW)/2.

Survival = Survival (%).

Economic conversion ratio (€/kg fish⁻¹) = FCR * feed cost

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Table 4. Body composition, nutritional indices, somatic indices and apparent digestibility measured in gilthead seabream fed increasing dietary single cell protein levels.

	CTRL	SCP10	SCP15	SCP20	<i>P</i> -value
<i>Whole body composition, %</i>					
Moisture	62.0±0.87	61.5±1.03	62.1±0.88	62.2±0.51	0.719
Protein	17.8±0.08 ^b	17.3±0.08 ^{ab}	17.2±0.03 ^a	17.5±0.06 ^{ab}	0.025
Lipid	16.9±1.29	17.0±1.17	16.9±0.80	16.7±0.46	0.990
Ash	3.4±0.16	3.4±0.22	3.5±0.19	3.1±0.19	0.257
<i>Nutritional indices, %</i>					
PER	1.52 ± 0.01	1.54 ± 0.02	1.52 ± 0.01	1.51 ± 0.01	0.178
GPE	28.0 ± 1.30 ^b	26.7 ± 0.22 ^{ab}	25.7 ± 0.72 ^a	26.3 ± 0.45 ^{ab}	0.015
<i>Apparent digestibility, %</i>					
Dry matter digestibility	92.9 ± 3.1	94.1 ± 0.9	93.4 ± 1.4	94.4 ± 1.1	0.761
Protein	95.5 ± 1.57	96.0 ± 0.16	95.5 ± 1.42	96.5 ± 0.90	0.307
<i>Somatic indices</i>					
CF	1.84 ± 0.17 ^{ab}	1.85 ± 0.15 ^b	1.78 ± 0.16 ^{ab}	1.73 ± 0.18 ^a	0.018
VSI	6.17 ± 1.02	5.65 ± 1.32	5.98 ± 0.77	5.93 ± 1.30	0.628
HSI	1.22 ± 0.22	1.35 ± 0.24	1.36 ± 0.25	1.31 ± 0.26	0.403

Data are given as the mean (n=3 ± SD). In each line, different superscript letters indicate significant differences among treatments ($P \leq 0.05$).

PER = Protein efficiency ratio = ((FBW-IBW)/protein intake).

GPE = Gross protein efficiency = $100 * [(\% \text{final body protein} * \text{FBW}) - (\% \text{initial body protein} * \text{IBW})] / \text{total protein intake fish}$.

HSI = Hepatosomatic index (%) = $100 * (\text{liver weight} / \text{FBW})$.

VSI = Viscerosomatic index (%) = $100 * (\text{viscera weight} / \text{FBW})$.

SD = Standard deviation.

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813 **Table 5.** Absorbed surface of the anterior and middle intestine evaluated in gilthead
 814 seabream by means of binarization method

	CTRL	SCP10	SCP15	SCP20	<i>P-value</i>
Anterior absorbent surface	3.68 ± 2.0	4.15 ± 1.5	3.76 ± 1.0	3.71 ± 1.2	> 0.05
Middle absorbent surface	1.38 ± 0.8	1.22 ± 0.4	1.33 ± 0.6	1.60 ± 0.8	> 0.05

815 Data are given as the tanks mean (n=12 ± SD)

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Table 6. Plasma biochemistry values measured in gilthead seabream fed increasing dietary single cell protein levels.

	CTRL	SCP10	SCP15	SCP20	<i>P</i> - value
GLU	160.4 ± 20.03	160.0 ± 16.12	147.3 ± 11.91	147.6 ± 14.41	0.5669
Urea	12.4 ± 0.99 ^a	13.3 ± 1.81 ^a	13.7 ± 0.73 ^a	16.5 ± 1.41 ^b	0.0006
CREA	0.25 ± 0.02	0.24 ± 0.02	0.25 ± 0.02	0.27 ± 0.04	0.3445
Uric Ac	0.21 ± 0.11	0.15 ± 0.03	0.18 ± 0.13	0.24 ± 0.12	0.5495
Tot Bil	0.04 ± 0.00	0.04 ± 0.00	0.04 ± 0.01	0.06 ± 0.01	0.0625
CHOL	264.0 ± 11.8	318.1 ± 17.6	301.8 ± 21.0	321.0 ± 33.4	0.0524
TRIG	555.4 ± 58.1 ^a	756.7 ± 30.6 ^{ab}	775.6 ± 198.0 ^b	754.5 ± 88.2 ^{ab}	0.0172
HDL	92.5 ± 4.14 ^a	109.8 ± 5.23 ^{ab}	106.8 ± 10.6 ^{ab}	114.6 ± 17.9 ^b	0.0337
TP	3.45 ± 0.21	3.60 ± 0.38	3.61 ± 0.28	3.69 ± 0.35	0.2625
ALB	0.87 ± 0.09	0.87 ± 0.13	0.87 ± 0.09	0.89 ± 0.09	0.9538
AST	59.6 ± 8.49	58.8 ± 15.3	66.7 ± 46.3	62.0 ± 12.8	0.9757
ALT	14.6 ± 0.97	16.3 ± 4.11	31.3 ± 36.36	15.7 ± 2.80	0.4859
ALP	230.1 ± 71.3	253.7 ± 40.5	234.3 ± 28.3	273.6 ± 53.0	0.7484
CK	3156 ± 3991	1119 ± 451	1313 ± 1107	2065 ± 1227	0.4770
LDH	504.0 ± 248.6	289.9 ± 188.5	301.3 ± 236.5	458.2 ± 289.1	0.5189
LACT	31.6 ± 5.38	26.2 ± 3.96	30.8 ± 9.52	23.2 ± 6.28	0.1364
Ca ⁺²	12.1 ± 0.50 ^a	12.3 ± 0.76 ^{ab}	12.6 ± 0.30 ^{ab}	12.8 ± 0.63 ^b	0.0492
P	10.9 ± 0.83 ^a	12.6 ± 1.14 ^b	11.9 ± 0.40 ^{ab}	11.9 ± 0.85 ^{ab}	0.0025
K ⁺	5.90 ± 1.18	6.31 ± 1.17	6.34 ± 0.22	6.16 ± 1.37	0.5291
Na ⁺	181.3 ± 3.58	176.8 ± 2.27	178.9 ± 5.51	179.7 ± 0.98	0.1605
Fe	80.0 ± 8.41	75.9 ± 13.1	77.3 ± 8.88	90.5 ± 17.0	0.2830
Cl	153.1 ± 1.83	148.7 ± 1.04	150.5 ± 4.37	151.3 ± 0.42	0.0637
Mg	2.94 ± 0.25	3.01 ± 0.11	3.08 ± 0.12	3.02 ± 0.09	0.8412
ALB/GLO	0.34 ± 0.02	0.32 ± 0.02	0.32 ± 0.01	0.32 ± 0.01	0.1310
CaxP	132.2 ± 14.11 ^a	155.8 ± 23.60 ^b	149.6 ± 8.27 ^{ab}	154.0 ± 18.1 ^{ab}	0.0113
Na/K	31.7 ± 5.90	29.0 ± 5.47	28.7 ± 0.23	30.4 ± 6.95	0.3155
Cur Ca ⁺²	14.8 ± 0.42 ^a	15.0 ± 0.66 ^{ab}	15.2 ± 0.26 ^{ab}	15.4 ± 0.55 ^b	0.0281

Data are given as the mean (n=15 diet⁻¹) ± SD. Different letters indicate significant difference (One-way ANOVA $P \leq 0.05$) between treatments.

GLU, glucose, (mg dL⁻¹); Urea, (mg dL⁻¹); CREA, creatinine, (mg dL⁻¹); Uric Ac, uric acid, (mg dL⁻¹); Tot Bil, total bilirubin, (mg dL⁻¹); CHOL, cholesterol, (mg dL⁻¹); TRIG, triglycerides, (mg dL⁻¹); HDL, high density lipoprotein; TP, total protein, (mg dL⁻¹); Alb, albumin, (g dL⁻¹); Ast, aspartate aminotransferase, (U L⁻¹); Alt, alanine aminotransferase (U L⁻¹); Alp, alkaline phosphatase, (U L⁻¹); CK, creatine kinase, (U L⁻¹); LDH, lactate dehydrogenase, (U L⁻¹); LAC, lactate (mmol L⁻¹); Ca⁺², calcium, (mg dL⁻¹); P, inorganic phosphorus, (mg dL⁻¹); K⁺, potassium, (mEq L⁻¹); Na⁺, sodium, (mEq L⁻¹); Fe, iron, (µg dL⁻¹); Cl, chloride, (mEq L⁻¹); Mg, magnesium, (mg dL⁻¹); ALB/GLO, albumin/globulin; CaxP, calcium*phosphorus; Na/K, sodium/potassium; Cur Ca²⁺, current calcium (mg dL⁻¹); SD, standard deviation.

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Table 7. Relative abundance (%) \pm SD of the most abundant phyla and families (all belonging to Firmicutes phylum) in the 4 experimental diet.

	CTRL	SCP10	SCP15	SCP20
Phylum				
Firmicutes	90.3 \pm 6.8	90.0 \pm 4.9	71.7 \pm 13.1	60.4 \pm 17.3
Family				
<i>Staphylococcaceae</i>	32.1 \pm 13.5	46.4 \pm 9.8	37.4 \pm 8.5	19.6 \pm 7.7
<i>Bacillaceae</i>	1.5 \pm 1.2	20.0 \pm 7.4	14.1 \pm 5.1	8.3 \pm 3.6
<i>Lactobacillaceae</i>	8.2 \pm 8.3	10.2 \pm 9.3	7.2 \pm 6.1	7.3 \pm 5.5
<i>Leuconostocaceae</i>	32.4 \pm 18.1	1.3 \pm 1.6	1.4 \pm 1.3	1.1 \pm 1.3

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826 **Figure 1**

827 A-D Process of image binarization. A) Original acquired image with 10x. B) Specific threshold to highlight
 828 the absorbent surface of the intestinal tract. C) Region of interest (ROI) used to measure number of binary
 829 objects inside each ROI, is indicated by arrow. D) Binary area, sum of areas of all binary objects. It is
 830 shown in square pixels and it's indicated in green. In these anterior and middle gut samples it corresponds
 831 to the absorbent surface of the intestinal tract.

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833 **Figure 2**

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835 Beta diversity and alpha diversity of gut microbiota of gilthead seabream fed with increasing dietary single
 836 cell protein levels (SCP). (A) Principal Coordinates Analysis (PCoA) based on unweighted UniFrac
 837 distances between gut microbiota structure of animals fed with CTRL diet, SCP10 diet, SCP15 diet and
 838 SCP20 diet. Samples are significantly separated (permutation test with pseudo-F ratios Adonis; $p = 0.001$).
 839 Black arrows are obtained by fitting the family relative abundance values for each sample within the
 840 ordination space (function envfit of the “vegan” R package, with a p -value < 0.001). (B) Boxplots show
 841 alpha diversity values, measured by Faith's Phylogenetic Diversity (PD_{whole tree}), Chao1 index, and
 842 amplicon sequence variants (observed_{ASVs}). All metrics showed a significant variation (Kruskal–Wallis
 843 test $p < 0.01$) of alpha diversity among dietary groups. More specifically, SCP10 and SCP15 groups showed
 844 a significant reduction of the internal ecosystem diversity compared to CTRL group, while the SCP20 group
 845 showed a significant opposite variation compared to control group (Wilcoxon rank-sum test, $p < 0.05$).

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847 **Figure 3**

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849 Microbiota composition (%) of distal gut content of gilthead sea bream fed with increasing dietary single
 850 cell protein levels (SCP). Bar plot summarizing the microbiota composition at phylum (A) and family level
 851 (B) of fish intestinal content. Only phyla with a relative abundance $\geq 1.0\%$ in at least 2 samples, and families
 852 with relative abundance $\geq 1.0\%$ in at least 2 samples are represented.

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854 **Figure 4**

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856 Taxonomic composition (%) of bacterial communities of distal gut content of gilthead sea bream fed with
 857 increasing dietary single cell protein levels (SCP). Distributions of relative abundance of genera that
 858 showed a significant variation between groups fed with different diets (Wilcoxon rank-sum test, *** $p \leq$
 859 0.001 ; ** $p \leq 0.01$; * $p \leq 0.05$). Only genera with a mean relative abundance $\geq 1.0\%$ in at least 2 samples
 860 were represented. The central box of each dataset represents the distance between the 25th and the 75th
 861 percentiles. The median between them is marked with a black line.

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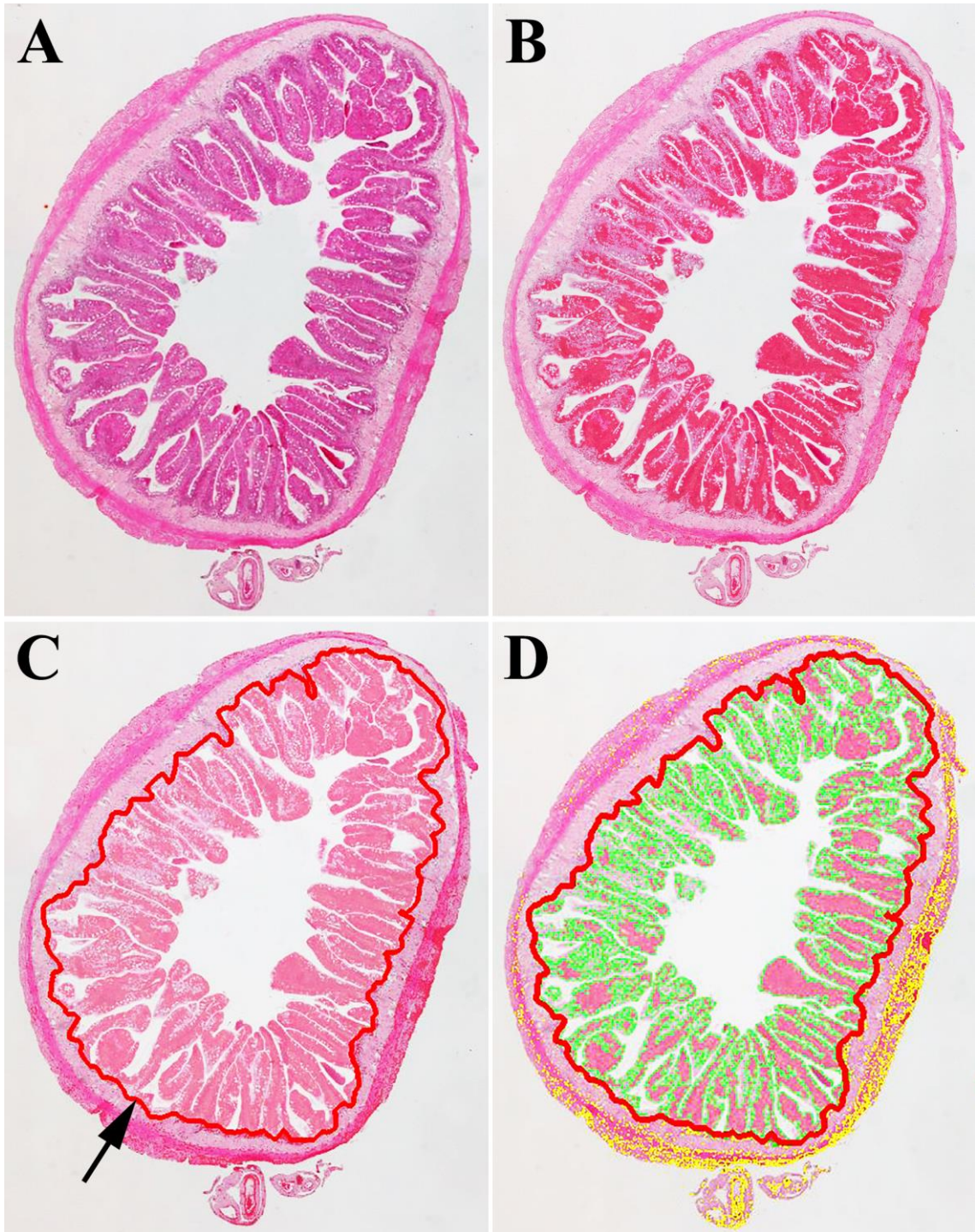
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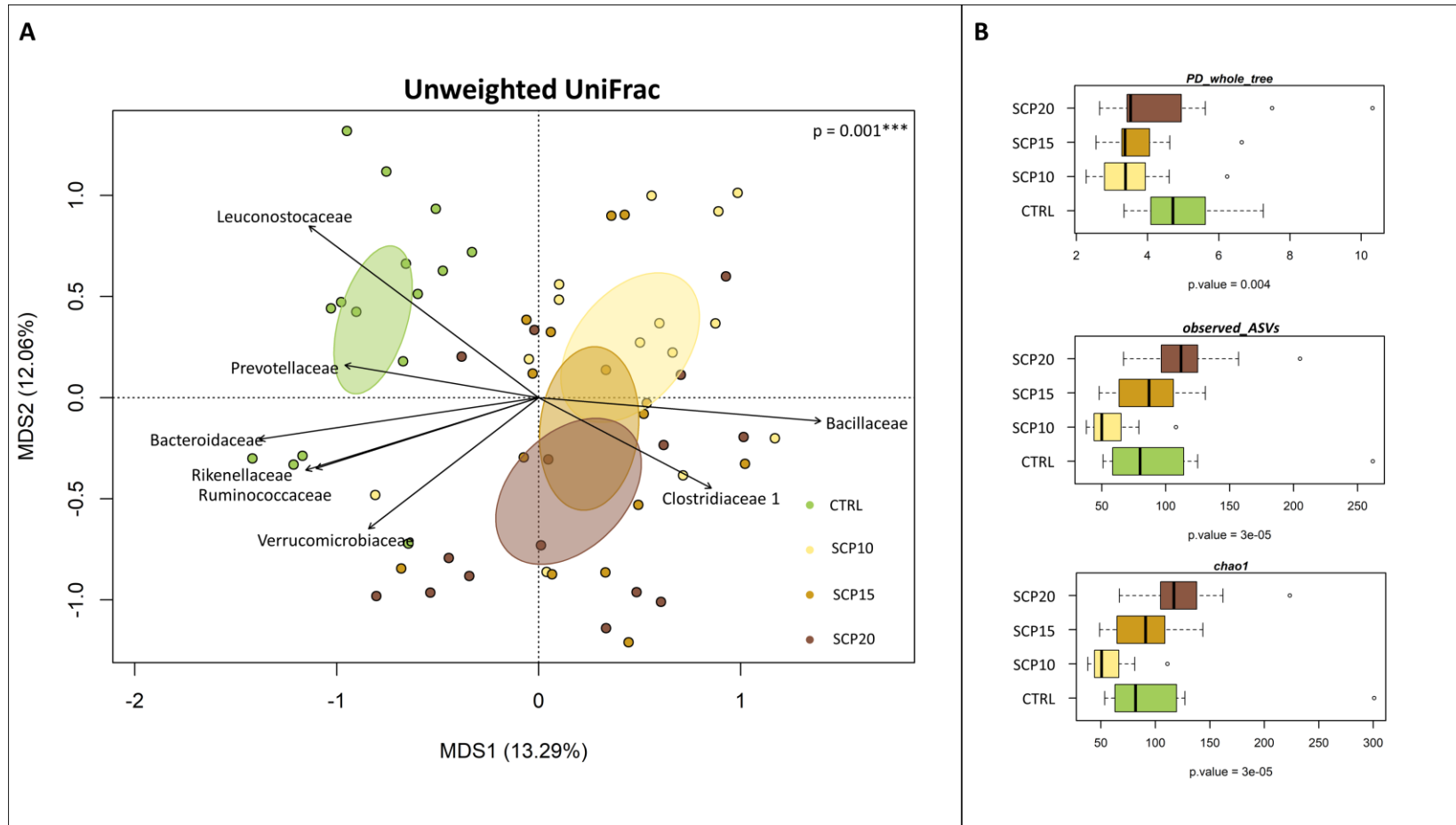
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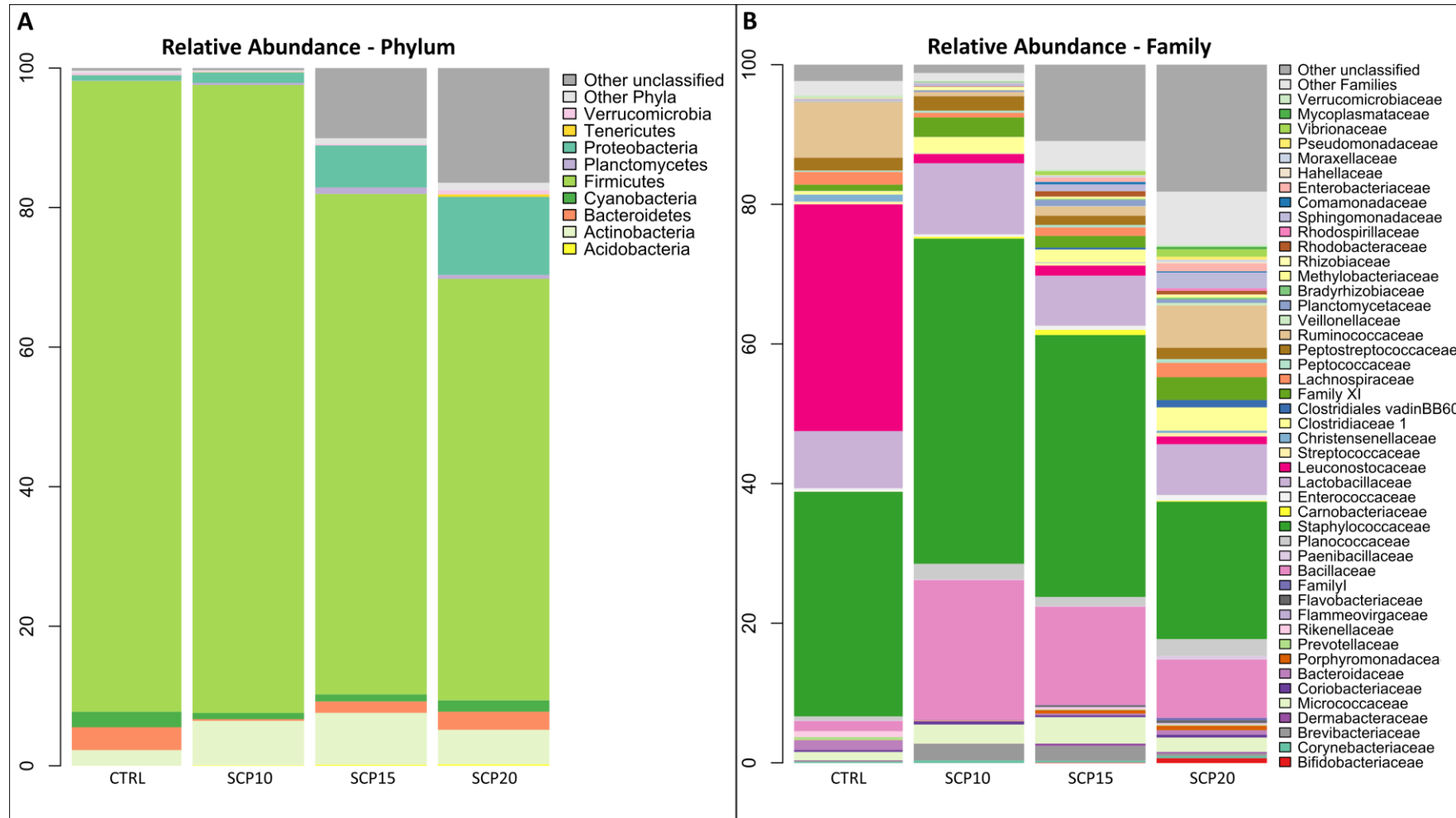
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Figure 2



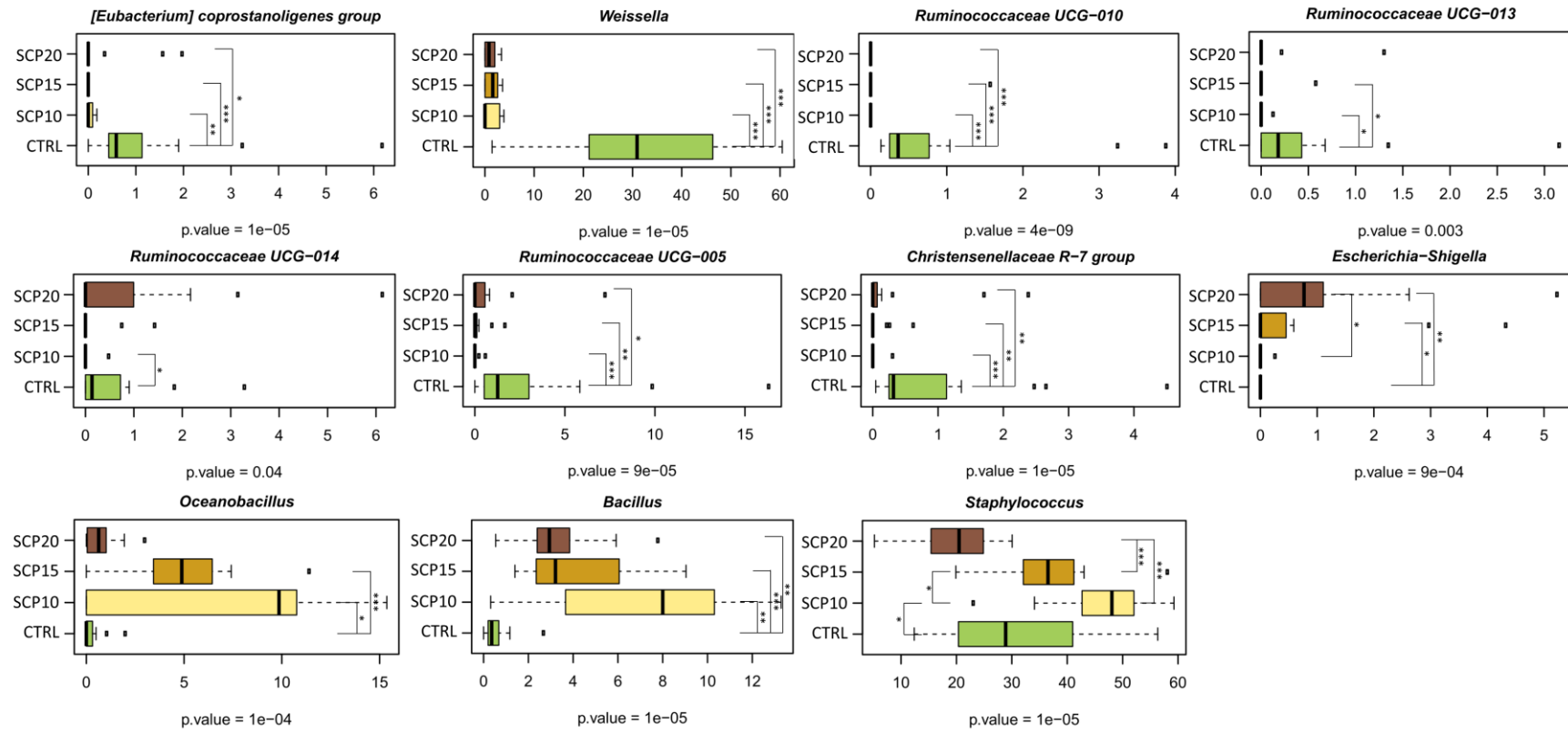
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