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1 **Oral sodium butyrate impacts brain metabolism and hippocampal**
2 **neurogenesis, with limited effects on gut anatomy and function in**
3 **pigs**

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16

17 **Short title:** Effect of oral butyrate on pig brain

18

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21

22 **Authors contribution:** DVL, CHM, JPL, PG, PP and JPS designed the research and secured
23 funding; DVL, SG, IN, JPL and PG performed the animal experiments; DVL, SG and CHM

24 performed brain imaging or provided analytical strategies; DVL, NC, SG and GR performed
25 brain post-mortem analyses; MF, VR, LLN, PG and JPL performed gut measurements, blood
26 and pancreas analyses; DVL, NC, PM, and JPL performed the statistical analyses; DVL wrote
27 the paper; and all co-authors read and revised the paper.

28

29 **Abbreviations**

30 CA-CP: commissura anterior-commissura posterior

31 CGM: cerebral glucose metabolism

32 CTI: computed tomography imaging

33 DCX: doublecortin

34 FDG: fluorodeoxyglucose

35 GCL: granular cell layer

36 PET: positron emission tomography

37 rCGM: regional cerebral glucose metabolism

38 ROI: region of interest

39 SGZ: sub-granular zone

40 SPM: statistical parametric mapping

41 SVC: small volume correction

42

43

44 **Abstract**

45 Butyrate can improve gut functions while HDAC inhibitors might alleviate neurocognitive
46 alterations. Our aim was to assess whether oral butyrate could modulate brain metabolism and
47 plasticity and if this would relate to gut function. Sixteen pigs were subjected to sodium
48 butyrate (SB) supplementation *via* beverage water or water only (C). All pigs had blood
49 sampled after 2 and 3 weeks of treatment, and were subjected to a brain positron emission
50 tomography after 3 weeks. Animals were euthanized after 4 weeks to sample pancreas,
51 intestine, and brain for gut physiology and anatomy measurements, as well as hippocampal
52 histology, Ki67 and DCX immunohistochemistry. SB compared to C treatment triggered
53 basal brain glucose metabolism changes in the nucleus accumbens and hippocampus
54 ($P = 0.003$), increased hippocampal GCL volume ($P = 0.006$) and neurogenesis (Ki67:
55 $P = 0.026$; DCX: $P = 0.029$). After 2 weeks of treatment, plasma levels of glucose, insulin,
56 lactate, GLP-1 and PYY remained unchanged. After 3 weeks, plasma levels of lactate were
57 lower in SB compared to C animals ($P = 0.028$), with no difference for glucose and insulin.
58 Butyrate intake impacted very little gut anatomy and function. These results demonstrate that
59 oral SB impacted brain functions with little effects on the gut.

60

61 **Keywords:** sodium butyrate, pancreas, gut, brain metabolism, cognition, hippocampus

62

63

64 **1. Introduction**

65

66 Brain and behavior alterations are observed in various pathologies including anxiety,
67 depression, Parkinson's or Alzheimer diseases, and are also suspected to operate in diet-
68 dependent metabolic disorders and obesity (1, 2). Histone deacetylase (HDAC) inhibitors
69 such as butyrate have been repeatedly shown to prevent or even treat alterations in brain
70 functions (*e.g.* memory) and behavioral disorders in various rodent models of diseases like
71 stress-induced anxiety (3) or Parkinson (4). Underlying mechanisms most often involve
72 factors entailed in brain neurogenesis and synapse plasticity, including brain- and glial cell
73 line-derived neurotrophic factors and nerve growth factor (3, 4). Interestingly, HDAC have a
74 memory-suppressant effect that can be neutralized by HDAC inhibitors such as butyrate (5-7).
75 In addition, long-term memory processes depend on H3 histone acetylation at the
76 hippocampus level, which is favored by butyrate (5, 7, 8). However, butyrate could also
77 prevent memory impairment and improve cognition through the upregulation of neurotrophic
78 factors (3, 9). Butyrate also functions as a ligand for a subset of G protein-coupled receptors
79 and as an energy metabolite (10).

80 Butyrate is a short-chain fatty acid naturally produced in the gut through bacterial
81 fermentation of dietary fibers that plays an essential role in gastrointestinal tract functions.
82 Butyrate is the primary energy source for colonocytes through mitochondrial β -oxidation and,
83 as already mentioned, modulates gene expression through HDAC inhibition. Dietary butyrate
84 enhanced growth rate and improved feed conversion rate before and during weaning in calves
85 (11, 12) and in piglets (13, 14). Different studies performed in piglets or calves showed that
86 butyrate also delayed gastric emptying (15, 16), increased the density of gastric parietal cells
87 (17) and colonic goblet cells (18), enhanced HSP27 and HSP70 expression in the stomach and

88 colon (12), and increased the activities of many brush-border enzymes in the small intestine
89 (12). Bioactive peptides or hormones secreted into the blood and/or locally produced in the
90 gastrointestinal tract were suggested to be responsible for butyrate effects on gastrointestinal
91 tract development. Altogether, these results are in favor of an effect of butyrate
92 supplementation on the gastrointestinal tract development in mammals especially as butyrate
93 is ingested soon after birth (19).

94 Whether dietary butyrate can modulate brain functions and metabolic activity in physiological
95 situation is poorly documented. A large body of evidence suggests that butyrate can be
96 absorbed from the gut and entirely metabolized either in the gut mucosa or in the liver, which
97 makes it very difficult to detect butyrate in systemic blood circulation (18). But butyrate is
98 known to cross the blood-brain barrier and to affect epigenetic machineries in the brain (20).
99 Accordingly, monocarboxylate transporters (MCTs) that mediate the transport of
100 monocarboxylates such as lactate, pyruvate, short-chain fatty acids and ketone bodies (21) are
101 present in the blood-brain barrier, glial cells and neurons. Although glucose is the major fuel
102 for normal, activated brain at the adult age (22), it is well established that ketones such as β -
103 hydroxybutyrate and acetoacetate are alternative energy substrates during suckling, starvation
104 or fasting periods and in situations of ketosis induced by high-fat diet (23-25). Several human
105 and animal studies using ^{18}F -fluorodeoxyglucose (FDG) and positron emission tomography
106 (PET) provided evidence that cerebral metabolism of glucose is reduced during ketosis (25-
107 27). However, it is not known whether butyrate, produced in the gastrointestinal tract and
108 present at low levels (μM) in blood, can modulate regional brain glucose metabolism.

109 The aim of the present study was to test the hypothesis that oral sodium butyrate
110 supplementation modulates brain metabolic activity and hippocampal structural plasticity, and
111 that part of this modulation stems in the effects of butyrate on gut biology and barrier

112 function. We tested this hypothesis in growing pigs as we developed brain imaging in this
113 species for investigating diet-induced brain and behavior modulations (2, 28, 29). The pig is
114 now recognized as a valuable alternative model to rodents for investigating human nutrition,
115 gastrointestinal tract development and gut-brain axis interplay (19, 30). Besides, butyrate
116 supplementation was shown to improve development of gastrointestinal tract anatomy and
117 functions in young mammals (13, 31). Exploring different pancreatic and intestinal
118 parameters might contribute to identify potential gut-brain communication pathways for the
119 putative effects of oral butyrate supplementation on brain biology.

120

121 **2. Materials & Methods**

122 The present experiment was conducted in accordance with the current ethical standards
123 of the European Community (Directive 2010/63/EU), agreement No. A35-622 for the
124 experimental facilities, and authorization No. 35-88 to experiment on live animals for the
125 principal investigator of the study. The Regional Ethics Committee in Animal Experiment of
126 Brittany (France) validated the entire procedure described in this paper (R-2012-DVL-01).

127 ***Animals and housing***

128 Four successive batches of four Large White/Landrace × Piétrain 2.5-month-old female
129 growing pigs of 28.9 ± 1.2 kg (mean \pm SEM) body weight (BW) at the beginning of the study
130 were used. The piglets were weaned at 28 days of age and subsequently housed in individual
131 pens (length 80 × width 60 × height 68 cm) equipped with a feeding trough and a drinking
132 nipple. The room temperature was kept at $23.4 \pm 0.1^\circ\text{C}$ with a 13:11-h light-dark cycle.

133 ***Diet and experimental beverages***

134 The pigs were fed a pelleted pre-starter diet from weaning (day 1) to 15 days after
135 weaning and then a starter diet until the end of the study (at approximately 3.5 months after
136 birth). The diets were formulated so that the nutrient composition (other than total sulfur
137 amino acids) met or exceeded recommendations of the NRC (1998) throughout the
138 experimental period (see 32 for the composition of the diets). Animals were divided into
139 two experimental groups. Sodium Butyrate group, SB: 28.7 ± 0.9 kg; Control group, C:
140 29.2 ± 0.9 kg), with two animals per group in each batch ($n = 8$ animals per group in total),
141 and were weighed weekly. During the whole experiment, all the pigs received 40 g of feed per
142 kg BW, which corresponds to the usual allowance in the pig facilities. *Ad libitum* feeding was
143 not used in this study to limit feed intake variability across individuals and to prevent feed
144 refusals. Indeed, the amount of butyrate supplied to the SB pigs was based on the amount of
145 feed ingested. The daily feed allowance was distributed once a day at 0930 h. One hour before
146 meal provision, at 0830 h, the animals received either water alone (control, C group) or a 10-
147 mM sodium butyrate solution (SB group). Access to tap water was permitted only once the
148 amount of water or butyrate solution was completely consumed, and was then presented again
149 in the mid-afternoon until the next day. The present dosage of butyrate (110 mg/kg BW/day)
150 is within the physiological range used in other studies with pigs (13, 31) and rodents (33, 34).
151 Control animals received only tap water.

152 ***Surgical implantation of jugular catheter***

153 Animals were fitted with a jugular catheter under general anaesthesia performed as
154 previously described (35) to achieve 2.2 Minimum Alveolar Concentration (MAC) for
155 isoflurane. Analgesia was achieved with an additional infusion of Fentanyl (30 to 100
156 $\mu\text{g}/\text{kg}/\text{hr}$ IV, Renaudin, Paris, France) during surgery. A silicon catheter was introduced into
157 the external jugular vein and tunnelled subcutaneously to exit in the interscapular space.

158 Animals were allowed to recover for one week after the surgical procedure before any
159 observation or treatment was made.

160 ***Blood sampling and analyses***

161 All the jugular blood samples were collected in tubes (10 ml) containing ethylene
162 diamine tetraacetate (EDTA) (VT-100 STK, 0.1 ml EDTA, 0.47 ml/l: 21 w/v %, CML,
163 Nemours, France). Moreover, a protease inhibitor cocktail (ref P2714-1BTL, SIGMA, L'isle
164 d'Abeau Chenes, Saint-Quentin Fallavier, France) was added to tubes aimed at GLP-1 assays.
165 Two weeks after the beginning of treatments, samples were collected just before (0830 h) the
166 distribution of the experimental treatments (tap water alone for the C group, sodium butyrate
167 solution for the SB group) and 1h after (0930 h), *i.e.* just before feeding (for lactate, insulin,
168 peptide YY – PYY, GLP-1 and osmolality measurements). Three weeks after the beginning
169 of the treatments, other samples were also collected in anaesthetized animals just before brain
170 imaging (for glucose, lactate and insulin measurements). All blood samples were kept in ice
171 before plasma preparation by centrifugation (2300 x g, 10 min, + 4°C). Plasma was recovered,
172 aliquoted and stored frozen at - 40°C until analysis.

173 Plasma concentrations of GLP-1 (pM) were measured by means of a commercial
174 Glucagon Like Peptide-1 (Active) Elisa Immuno Assay Kit (ref. EGLP-35 K, Merck KGaA,
175 Darmstadt, Germany). Peptide YY plasma concentrations (ng/ml) were determined using a
176 commercial PYY Elisa Immuno Assay Kit (ref. EK-059-04, Phœnix Pharmaceuticals, INC., ,
177 Burlingame, CA 0, USA). Insulin concentration was measured by RIA (Insulin-CT;
178 CisbioBioassays, Codolet, France). Plasma levels of glucose and lactate were measured in
179 duplicate by automated spectrophotometry (Konelab 20i; Thermo Fisher Scientific, Waltham,
180 MA, USA). Plasma osmolality was measured on a micro-osmometer (Hermann Roebling
181 12/12 DR, Berlin, Germany).

182 ***Brain imaging procedure***

183 After three weeks of treatment, the animals underwent a brain-imaging session to
184 investigate basal brain metabolism. The brain imaging modality used to investigate the
185 regional cerebral glucose metabolism (rCGM) was the positron emission tomography (PET)
186 of ^{18}F -fluorodeoxyglucose (^{18}F -FDG, CIS bio international, France). The procedure for
187 animal preparation and anesthesia was similar to that described in our previous publications
188 (36, 37). The animals were placed in a head first supine position on the bed of a whole body,
189 high-resolution PET. A venous catheter was then inserted into their left ear in order to inject
190 the ^{18}F -fluoro-deoxyglucose radiolabel (^{18}F -FDG, 200 MBq in 20 ml), at least 30 min after the
191 beginning of anesthesia. Brain images were acquired on a computed tomography imaging
192 machine in 3D mode (CTI/Siemens ECAT, 962, HR+) and processed as described in Clouard
193 et al. (36).

194 ***Euthanasia, organ and tissue collection***

195 After four weeks of treatment, the pigs were euthanized with intravenous administration
196 of T61® following a pre-anesthesia with ketamine (5 mg/kg intramuscularly, Rhône Mérieux,
197 Lyon, France), and then exsanguinated.

198 *Brain collection, fixation and conservation.* After euthanasia, the pigs were decapitated
199 and the head was perfused *via* the carotids with 2 L of 1% sodium nitrite and 5 L of buffered
200 4% formalin at 4°C. The brains were kept in buffered 4% formalin during 24h and then
201 transferred in 30% sucrose with 0.1% sodium azide at 4°C until being sliced with a
202 cryomicrotome. Brain slices were deposited on glass slides and stored at -20°C.

203 *Gut tissue collection.* Two pigs per afternoon (one C and one SB) were randomly
204 slaughtered at 30 min intervals, 2.5 h after the last treatment (tap water *vs.* sodium butyrate
205 solution, but no meal). After laparotomy, the entire gastrointestinal tract was carefully
206 removed and the small intestine was isolated by ligation before removal. The pancreas was

207 weighed and three pieces (about one g per sample) were gently collected, frozen in liquid
208 nitrogen and then stored at -20°C until enzyme activity analysis. The mesentery was removed
209 all along the small intestine that was then measured. Twenty five-centimeter segments of
210 duodenum (20 cm distal to Treitz ligament), mid-jejunum and distal ileum (ending 10 cm
211 proximal to the ileo-cecal valvula) were cut, emptied of digesta contents and gently rinsed
212 with cold saline. For each collected segment, cross-sectional tissue samples were made for the
213 following preparations or analysis: 15 cm for immediate use in Ussing chambers (not
214 duodenum) and quickly transported to the laboratory in cold Ringer bicarbonate solution (see
215 below); 5 cm for fixation (in buffered formalin 10%), paraffin embedding and histology; and
216 1 cm of whole tissue (cut in small pieces) for HSP analysis was snap-frozen in liquid nitrogen
217 and then stored at -80°C. Part of the remaining collected tissues was scraped using a glass
218 slide (small intestine, 4-5 cm) or cut into small pieces (colon, 1 cm) for enzyme
219 determination, snap-frozen and then stored at -20°C.

220 ***Brain immunohistochemical analysis***

221 Brain immunohisto-fluorescence was performed on 30- μ m brain sections of the left
222 hippocampus. One section every 25 sections was selected for antigen Ki67 and doublecortin
223 (DCX) immunohistochemistry. Only the first 10 selected sections were gathered and used for
224 immunohistochemistry, which covers approximately half of the hippocampus, *i.e.* 7.5 mm
225 over around 15 mm (see 38 for comparison with whole hippocampus sampling). Ki67 and
226 DCX staining was performed on adjacent sections for each hippocampal progression site.
227 Ki67 protein is expressed in proliferating cells while DCX is a microtubule-associated protein
228 expressed by neuronal precursor cells and immature neurons (39).

229 Sections were thawed during 10 min at room temperature and washed three times for
230 10 min in phosphate buffered saline (PBS). After the last wash, 150 μ L of blocking buffer

231 (PBS solution, Gibco by Life Technologies; 10% horse serum, Sigma; 0.3% Triton, Sigma)
232 was then dropped on the region of interest (ROI) and incubated during 1 h at room
233 temperature in a humidity chamber. Blocking buffer excess was carefully removed with
234 absorbent tissue and 150 μ L of primary antibody (Ki67: Anti-Ki67 antibody ab15580,
235 dilution 1:200, Abcam; DCX: Anti-doublecortin (C-18) antibody sc8066 1:200, Santa Cruz)
236 was added on the ROI. Parafilm was carefully laid on tissue slice and slides were stored at
237 4°C for at least 16 h until the next morning. Sections were then washed three times with
238 blocking buffer for 10 min and 150 μ L of secondary antibody (Ki67: Alexa Fluor 488
239 Conjugate 1/200, Cell Signaling Technology; DCX: CyTM3-conjugated AffiniPure Donkey
240 Anti-Goat IgG (H+L) 1:500, Jackson ImmunoResearch Laboratories, West Grove, PA, USA)
241 was added for a 2-h incubation at room temperature. Slides were finally mounted with
242 Fluoroshield Mounting Medium with 4',6-diamidino-2-phenylindole (2 drops per slide, ref
243 17513, Abcam, Cambridge, UK). Sections were examined under a fluorescent microscope
244 (Eclipse 80i, Nikon), digitized and large-field mosaics were performed with micro-manager
245 (ImageJ plugin) in order to get the whole hippocampal section on one image.

246 Determination of hippocampus and granular cell layer (GCL) volumes, number of
247 Ki67-positive nuclei in the sub-granular zone (SGZ), number of DCX-positive cells in the
248 GCL were performed for each selected hippocampal section and reported as an inferred value
249 per site considering that each site is a block of 25 sections.

250 ***Gut physiology measurements ex vivo (Ussing chambers)***

251 *Electrophysiology ex vivo.* Six Ussing chambers (2 per gut site; P2250 model;
252 Physiological Instruments, San Diego, CA, USA) were used for investigating gut
253 electrophysiology (and permeability, see below) in each pig. Jejunal, ileal and colonic
254 mucosae were stripped off the muscular layers, cut into two adjacent pieces and immediately

255 mounted in chambers (1 cm² mucosa in contact with 2.5 ml Ringer bicarbonate buffers). Gut
256 electrophysiology (and permeability) parameters in basal conditions were measured in the two
257 chambers and values averaged per tissue. Then, one chamber was kept in basal condition
258 while the other was used for measurements under oxidative stress using monochloramine
259 (final concentration 1 mM, both sides of the tissue) (40). The Ringer-bicarbonate buffer
260 contained 16 mmol/L of glucose and 16 mmol/L of mannitol on the serosal and mucosal
261 sides, respectively. Chambers were kept at 39°C and buffers oxygenated all the time
262 automatically. Tissue viability was checked every 30 min by recording tissue electrical
263 potential difference automatically. Gut mucosae were left equilibrating for 20 min before
264 short-circuit current (I_{sc}) and the trans-epithelial electrical resistance (TEER) were
265 determined under clamp condition (3 mV for 300 ms every 30 s) using Ohm's law (41).
266 Colonic mucosa sodium-dependent glucose absorption capacity and chloride secretion
267 capacity were determined after addition of D-glucose (16 mM, mucosal side) and the
268 cholinergic agonist carbachol (1 mM, serosal side), respectively (41).

269 *Trans- and para-cellular permeability ex vivo.* Gut mucosa permeability to FITC-
270 dextran 4000 (FD4, MW 4 kDa; para-cellular permeability marker) and horseradish
271 peroxidase (HRP type II, MW 40 kDa; trans-cellular permeability probe) was assessed
272 exactly as previously described for the small intestine and the colon (42, 43). When tissue
273 electrophysiology was stabilized (see above), FD4 and HRP were added to the mucosal side
274 of tissues and serosal fluid (200 µL, replaced by 200 µL Ringer-glucose) was collected
275 kinetically (every 30 min for 120 min). Fluid FD4 and HRP concentrations were determined
276 by fluorimetry and spectrophotometry, respectively and marker transmucosal flows were
277 calculated as detailed previously (44).

278 ***Gut anatomy, enzyme activities and heat shock proteins (HSP)***

279 *Intestinal length and villous-crypt architecture.* Small intestinal length was measured,
280 pieces of tissues were collected at various sites along the gut (see above) and were dehydrated
281 and then embedded in paraffin as reported previously (42, 43). Five-micrometer tissue
282 sections were prepared, dewaxed, rehydrated and stained with eosin and haematoxylin, and
283 villi and crypts (10–15 per section, well oriented, full size) were measured for their length,
284 width and surface area (42, 43). Intestinal villus height-to-crypt depth (VH:CD) ratio, and ‘M’
285 ratio for estimating three-dimension mucosal surface area (45) were also calculated. Villous
286 and crypt morphology parameters were averaged per site per pig before statistical analysis.

287 *Pancreatic and intestinal enzyme activities.* After thawing, the samples coming from
288 pancreas tissue and intestinal mucosa, protein content was measured and three pancreatic
289 enzymes (trypsin, E.C. 2.3.21.4; lipase, E.C. 3.1.1.3; amylase, E.C. 3.2.1.1.) were analyzed as
290 reported before (46). The activities of five enzymes – two peptidases (amino-peptidase N,
291 E.C. 3.4.11.2; dipeptidyl-peptidase IV, DPP4, E.C. 3.4.14.5), two disaccharidases (sucrase,
292 E.C. 3.2.1.48; maltase, E.C. 3.2.1.20) and one phosphatase (intestinal alkaline phosphatase,
293 E.C. 3.1.3.1) – were determined in small intestinal mucosa homogenates as previously
294 reported (42, 43). Alkaline phosphatase activity was also determined in a piece of tissue from
295 proximal colon. Specific (/g protein) and total (/kg of BW for pancreas, and /g fresh mucosa
296 for intestine) enzymes activities (or activity concentrations for intestinal alkaline phosphatase)
297 were calculated as previously reported.

298 *Gut heat shock proteins.* Inducible heat shock proteins HSP27 and HSP70 were
299 determined in gut tissues as reported before (42, 43).

300 ***Statistical analysis***

301 *Statistical analysis of body weight, pancreatic enzymes and blood plasma data.* Data
302 were tested for normal distribution and homogeneity of variances. Pancreatic enzymes and

303 blood plasma data were analyzed with a simple ANOVA or an ANOVA for repeated
304 measurements over time to compare data obtained before and after water *vs.* butyrate intake.
305 Interactions between treatment and time were also investigated (SPSS Statistics software
306 version 20.0; IBM, Somers, NY, USA). All data are reported as means \pm SEMs. The level of
307 significance for all analyses was set as $P < 0.05$ and trends were considered at $0.05 \leq P \leq$
308 0.10.

309 *Brain immunodetection data.* For the immunohistochemistry data, animal numbers are
310 N=7 for Control group (C) and N=6 animals for Butyrate group (SB). Two animals (1 C and 1
311 SB) were excluded because of an incorrect slicing axis or unsuccessful
312 immunohistochemistry, respectively. Hippocampal site-based statistical analyses were
313 performed with an ANOVA (SPSS Statistics software version 20.0; IBM, Somers, NY, USA),
314 with hippocampal site progression and animal as covariates. Data are presented as mean
315 values for the entire block of 25 slices for each hippocampal site.

316 *Statistical brain image analysis.* The regional ^{18}F -FDG uptake was standardized to the
317 brain global uptake using proportional scaling in order to suppress individual differences in
318 global cerebral glucose metabolism (CGM). Brain images were analyzed using independent *t*
319 tests to compare the Butyrate (SB) and Control (C) groups with the two following contrasts:
320 SB > C and SB < C. After a whole-brain first-level analysis without *a priori*, we performed
321 small volume correction (SVC) analyses on ROIs, upon *a priori* hypotheses, *i.e.* in areas
322 associated with learning and memory, attentional processes, reward and cognitive control
323 (anterior prefrontal cortex, dorsolateral prefrontal cortex, orbitofrontal cortex, cingulate
324 cortex, hippocampus, parahippocampal cortex, amygdala, caudate, putamen, nucleus
325 accumbens, globus pallidus). With this analysis, allowing for voxel-to-voxel comparisons
326 within restricted ROIs, we managed to identify the voxels for which the activity was
327 statistically different between groups (treatments) in the ROIs. For the SVC analyses, the

328 value of $P = 0.05$ (uncorrected for multiple comparisons) was set as the significance threshold
329 with a cluster size $k \geq 25$ voxels. The statistical analysis with Statistical Parametric Mapping 8
330 (SPM8) produced a listing of voxels for which the activation differed between treatments.
331 Each voxel was associated with a set of coordinates (x y z) corresponding to its spatial
332 location in the *commissura anterior-commissura posterior* (CA-CP) plane with CP set as the
333 origin. The ROIs chosen for the SVC analysis were anatomically identified on the basis of a
334 3D digitized pig brain atlas developed in our laboratory (47).

335 *Statistical analysis of intestinal parameters.* Intestinal parameter data were analyzed
336 using the Statistical Analyzing System (version 8.1, 2000; SAS Institute Inc., Cary, NC) using
337 models that included the treatment (C vs. SB) the site and the treatment by site interaction.
338 MIXED procedures with REPEATED statement for gut sites were used. Results are presented
339 as least-square means and pooled SEM. Least-square means comparisons for each
340 combination of treatment and site were made only when a tendency ($P \leq 0.10$) for a treatment
341 by site interaction was observed. In these cases, means were separated using Bonferroni post-
342 hoc comparison test. Permeability data displaying non-Gaussian distribution were log-
343 transformed before statistical analysis. Effects were considered significant at $P < 0.05$ and as
344 trends at $0.05 \leq P \leq 0.10$.

345

346 **3. Results**

347 *Body weight and blood analyses*

348 During the whole experiment, there was no significant difference in BW between
349 groups (from 29.2 ± 0.9 to 42.0 ± 1.0 kg in C pigs, from 28.7 ± 0.9 to 41.9 ± 1.5 kg in SB

350 animals, from $P = 0.494$ to $P = 0.949$). All the pigs consumed their daily ration and there was
351 no difference in feed consumption between groups.

352 *Blood analyses before and after drinking (after two weeks of treatment).* Lactate (Fig
353 1A), PYY (Fig 1B) and GLP-1 (Fig 1C) jugular plasma levels significantly decreased after
354 drinking, independently from the treatment (lactate, from 922 ± 93 to 695 ± 48 $\mu\text{mol/l}$,
355 $P = 0.040$; PYY, from 2.9 ± 0.2 to 2.7 ± 0.2 ng/ml , $P = 0.0140$; GLP-1, from 14.4 ± 1.8 to
356 13.4 ± 1.7 pM , $P = 0.023$). There was an interaction between treatment and time ($P = 0.025$)
357 for osmolality that increased in SB pigs and decreased in C pigs after beverage drinking
358 (Fig 1D). There were no other effects for these four parameters. No effect of the treatment or
359 sampling time nor interaction was observed for plasma glucose and insulin (data not shown).

360 *Blood analyses before brain imaging (after three weeks of treatment).* SB pigs had
361 lower lactate jugular plasma levels than C pigs (782 ± 52 vs. 1087 ± 114 $\mu\text{mol/l}$; $P = 0.028$)
362 (Fig 1E). There was no difference between groups for the glucose and insulin jugular plasma
363 levels (data not shown).

364 ***Regional brain glucose metabolism***

365 The without *a priori* analysis revealed very few structures with basal activity differing
366 between groups (Table 1). The right hippocampus ($P = 0.003$), the right primary
367 somatosensory cortex ($P < 0.001$) and left inferior colliculus ($P = 0.002$) were more activated
368 in the SB group than in the C group. The left primary somatosensory cortex ($P = 0.002$) and
369 the left inferior temporal gyrus ($P = 0.004$) were less activated in the SB group than in the C
370 group.

371 The SVC analysis identified only two brain structures over 14 (bilaterally) selected for
372 which the regional glucose metabolism was different between groups (Table 1). The right

373 nucleus accumbens was less activated in the SB group than in the C group ($P = 0.010$), while
374 both activated ($P = 0.003$) and deactivated ($P = 0.010$) clusters were found in the right
375 hippocampus in the SB group compared to C group.

376 Fig 2 indicates the activated and deactivated clusters for the image contrast analyses
377 comparing the SB group to C group.

378 ***Hippocampal structural plasticity***

379 Hippocampal and GCL volumes increased along the hippocampus site progression (Fig
380 3). Along hippocampal progression, similar hippocampal volume was measured, whereas the
381 GCL volume was significantly higher in the SB group (Fig 3B, $P = 0.006$).

382 Ki67-positive nuclei and DCX-positive cells were detected in order to assess the
383 structural plasticity of the hippocampus. Both Ki67-positive nuclei density and DCX-positive
384 cells density appeared higher in the SB group compared to C group (Fig 4A). Along
385 hippocampal site progression, significantly higher numbers of Ki67-positive nuclei and DCX-
386 positive cells were noticed in the SB group as compared to C group (Fig 4B; Ki67: $P = 0.026$;
387 DCX: $P = 0.029$).

388 ***Gut physiology ex vivo (Ussing chambers)***

389 *Electrophysiology ex vivo.* Mucosal electrophysiology parameters were determined in
390 both basal and oxidant stress conditions (Table 2). Basal Isc did not display any significant
391 treatment by site interaction, or treatment or site effect. Basal glucose absorption capacity
392 (dIsc-glucose) displayed a significant treatment by site interaction ($P = 0.04$), but no
393 treatment or site effect. The interaction and site effects were not significant for chloride
394 secretion capacity (dIsc-carbachol) but treatment effect in basal condition tended to be
395 significant (SB < C, $P = 0.09$) for this variable.

396 *Trans- and para-cellular permeability ex vivo.* There was no significant treatment by
397 site interaction for both permeability markers and measurement conditions (Table 3).
398 However, HRP (trans-cellular) permeability in basal condition tended to be globally lower in
399 SB compared to C treatment ($P = 0.09$). Treatment differences for FD4 (para-cellular
400 permeability) in basal or stress condition did not reach significance (Table 3).

401 ***Gut anatomy, enzyme activities and heat shock proteins (HSP)***

402 *Intestinal length and villous-crypt architecture.* Small intestinal length did not differ
403 between C and SB groups (respectively 0.34 and 0.35 m/kg BW, SEM = 0.042; $P=0.620$).
404 Similarly, there was no significant treatment by site interaction or significant difference
405 between treatments for all parameters of villous-crypt (small intestine), crypt (colon)
406 architecture, villous height-to-crypt depth ratio or M factor (Table S1, Supplemental
407 Material). By contrast, and as expected, site effects were (highly) significant for all the
408 studied variables, except villous perimeter and crypt depth (Table S1).

409 *Pancreatic and intestinal enzyme activities.* No effect of butyrate ingestion was shown
410 concerning pancreas and intestine protein contents as well as pancreatic enzyme activity
411 levels (Table 4). There was a tendency ($P = 0.063$) for a treatment by site interaction for
412 amino-peptidase N, showing that this enzyme total activity was lower in SB than C group in
413 the jejunum ($P < 0.05$), with no differences in the duodenum or ileum. There was a tendency
414 ($P = 0.07$) for a treatment by site interaction for DPP4 specific activity, though without
415 significant differences at any site. There was no significant treatment by site interaction or
416 treatment differences for disaccharidases and intestinal alkaline phosphatase activities (Table
417 5). Site effect was highly significant ($P < 0.0001$) for all the analyzed intestinal enzymes.

418 *Gut Heat shock proteins.* Inducible HSP27 and HSP70 were assayed using Western
419 blotting in gut tissues. There was no significant treatment by site interaction or treatment

420 differences for these HSPs (Table S2, Supplemental Material). Site effect was (HSP27,
421 $P < 0.0001$) or tended to be (HSP70, $P = 0.07$) significant.

422

423 **4. Discussion**

424

425 The main result of this study is that butyrate supplementation *via* beverage water during 3-4
426 weeks in growing pigs fed a standard diet can trigger regional brain glucose metabolism
427 changes in several brain structures, including the hippocampus. An increased glucose
428 metabolism relative to overall brain metabolism was detected in the right hippocampus with
429 the first-level analysis and then confirmed with the SVC analysis. This second-level analysis
430 performed on specific regions of interest also detected deactivated clusters in the right
431 hippocampus and nucleus accumbens. Increased hippocampal plasticity was also detected in
432 post-mortem histochemical analyses, with more neurogenesis and an increased GCL volume.
433 Blood analyses revealed lower plasma levels of lactate in the SB group compared to the C
434 group at the moment of brain imaging, but no difference in terms of glucose or insulin levels.
435 This result demonstrates that the differences in basal brain activity observed between groups
436 were not attributable to modifications in plasma glycaemia at the time of brain imaging.
437 Incidentally, there was no short-term effect of butyrate intake on lactate, glucose, PYY, GLP-
438 1 and insulin plasma levels either, indicating that the butyrate effect on regional brain
439 metabolism, perceivable at a distance from butyrate ingestion, was not related to systemic
440 glucose control or gut hormones variations. The absence of treatment effect for GLP-1 and
441 PYY plasma levels was quite surprising because butyrate is known to stimulate the release of
442 GLP-1 from intestinal L-cells (48) and increase fasting and postprandial plasma PYY
443 concentrations when infused in the colon (49). GLP-1 can improve learning and memory *via*

444 an action at the hippocampal level (50), but this mechanism was probably not at the origin of
445 the effects observed in our study. It is also possible that butyrate never reached L-cells and
446 was absorbed at the stomach level, which would prevent any significant increase of plasma
447 GLP-1. In this case, the gut-brain communication pathway involved needs further
448 investigation.

449 Our results showed that chronic butyrate supplementation led to a decreased regional
450 glucose metabolism in the dorsal hippocampus and nucleus accumbens, as well as an
451 increased regional glucose metabolism in the ventral hippocampus, together with an increased
452 GCL volume and hippocampal neurogenesis. These data suggest that butyrate, which is
453 largely metabolized by the gut and the liver, and thus found at very low concentrations (μM)
454 (18), may influence brain glucose metabolism. It was previously shown in a rat model of
455 spreading depression, that ^{14}C -labelled butyrate, injected intravenously, was readily taken up
456 and metabolized by adult brain, probably by astrocytes (20). Moreover, this study showed that
457 enhanced cortical butyrate uptake parallels local increase in ^{14}C -deoxyglucose utilization (20).
458 However, the increased glucose metabolism found in some brain regions of the SB groups is
459 in contrast with its reduction generally observed under ketogenic diets (26, 27, 51). The
460 reduction of brain glucose (oxidative) metabolism during ketosis correlates with the increase
461 in plasma ketones (acetoacetate and β -hydroxybutyrate) concentration (mM), and is
462 associated with the increase of cerebral metabolic rate of ketones (26, 27, 51). It has been
463 proposed that the tricarboxylic acid cycle (TCA) is fed by acetyl-CoA derived from ketones
464 thereby decreasing glucose oxidation downstream pyruvate dehydrogenase (PDH) (51).
465 However, concerning butyrate, there is no evidence for such a metabolic behaviour in brain
466 cells. One explanation for the discrepancy in our observation of a stimulation of glucose
467 metabolism could be that, in comparison to the high concentration of ketones supplying the
468 brain, butyrate brain levels may be too low to compete with glucose oxidation. Alternatively,

469 butyrate-stimulated brain glucose metabolism could implicate the ability of butyrate to act as
470 an HDAC inhibitor, thereby upregulating the transcription of genes involved in glucose
471 metabolic pathway. Indeed, a transcriptomic study of the human colonic mucosa reported that
472 butyrate stimulated the transcription of PDH, citrate synthase (TCA enzyme) and succinate
473 deshydrogenase (respiratory chain) (52). Furthermore, using germ-free mice, Donohoe et al.
474 (53) suggested that butyrate stimulates mitochondrial oxidation of glucose in colonic
475 epithelial cells. Interestingly, Moretti et al. (9) showed that sodium butyrate treatment in a rat
476 model of mania modified mitochondrial function in prefrontal cortex, hippocampus,
477 amygdala and striatum. Therefore, our data could suggest that increasing colonic production
478 of butyrate, for example with fibre-enriched diets, could influence brain glucose metabolism.

479 The dorsal and ventral parts of the hippocampus are two functionally different
480 structures that are connected to different neuronal networks (54). The dorsal hippocampus
481 performs primarily cognitive functions whereas the ventral hippocampus relates to stress,
482 emotions and affect. For example in rats, lesions in the dorsal hippocampus caused spatial
483 memory impairment in the radial arm maze, contrary to lesions in the ventral hippocampus
484 (55), and stress-related genes were found expressed in the ventral hippocampus of prenatally
485 stressed rats (56). Interestingly, ventral hippocampal afferents to the nucleus accumbens were
486 also found regulating susceptibility to depression (57). If the changes of regional metabolism
487 in the nucleus accumbens and hippocampus further to butyrate supplementation were
488 confirmed in a pig model of anxiety/depression, early malnutrition, or neurocognitive
489 alterations, then behavioral tests might reveal whether these brain modulations are
490 accompanied by changes in cognitive abilities and susceptibility to stress or anxiety.
491 Interestingly, we previously demonstrated that perinatal administration of butyrate to dams
492 improved performance in the spatial memory test on young rats that visited more frequently
493 the novel arm during a Y-maze test (58). Gieling et al. (59) reviewed studies in pig cognition

494 and identified several promising types of tasks such as versatile spatial free-choice type tasks
495 allowing for simultaneous measurement of several behavioural domains. In a recent study, we
496 showed that piglets with unbalanced perinatal nutrition had smaller hippocampal granular cell
497 layer and decreased neurogenesis, as well as different cognitive responses compared to
498 control animals during a spatial memorization and learning task (38). These pig models might
499 be valuable to assess whether butyrate supplementation could reverse the deleterious
500 neurocognitive and emotional effects of early malnutrition and/or mental pathologies.

501 The growing pigs used in our study were approximately 2 month-old at the beginning of
502 the experiment. It is therefore likely that their brain functions and metabolism were still
503 subject to plasticity mechanisms, and that their brain did not completely switch from
504 monocarboxylates consumption during the early postnatal period to glucose consumption
505 later on. Chronic ingestion of sodium butyrate might modify the circulating levels of
506 monocarboxylates, including butyrate and lactate. This could explain the decreased plasma
507 lactate levels in SB pigs just before brain imaging, and the fact that plasma lactate levels were
508 lower in SB than C pigs. In the hypothesis of an increased passage of butyrate and perhaps
509 lactate to the brain, the question of their interaction or competition with glucose can be raised.
510 The radioligand used in our study to investigate regional brain metabolism was a glucose
511 analogue (^{18}F -FDG) that is uptaken by brain cells, including glial cells and neurons. This
512 exogenous glucose competes with endogenous glucose. Aside from the conventional view of
513 brain energy metabolism stating that glucose is consumed preferentially in neurons (22), an
514 alternative model suggests that glucose preferentially taken by astrocytes, is degraded to
515 lactate and then exported into neurons to be oxidized (60). Facing such a complex situation,
516 the only conclusion we can draw up upon our own results is that chronic consumption of
517 sodium butyrate during three weeks was sufficient to decrease basal plasma lactate levels and
518 modify the regional glucose metabolism in the hippocampus and nucleus accumbens.

519 Our data indicate that a 4-week period of sodium butyrate intake impacted very little gut
520 anatomy and function. Indeed, this treatment only tended to reduce gut secretory capacity and
521 trans-cellular permeability in basal condition and lowered amino-peptidase N total activity in
522 the jejunum. By contrast, neither villus-crypt architecture nor para-cellular permeability and
523 inducible HSPs were affected. This is in contrast with our previous investigations in piglets
524 and calves (12, 13), possibly because in this experiment, the pigs were older and heavier than
525 previously. Based on the gut variables measured in the present experiment, this organ appears
526 to contribute very little to the effects of butyrate observed at the brain level, but further
527 explorations are needed to see whether butyrate might modulate other gastrointestinal tract
528 parameters than those investigated in this study, and consequently send information to the
529 brain to modulate its metabolism and plasticity. For example, peripheral and central immunity
530 might be explored, as well as inflammation markers at both levels.

531 In conclusion, this study demonstrated that oral sodium butyrate influences regional
532 brain metabolism and plasticity, and that the gut, systemic glucose control, or gut hormones
533 variations appear to contribute very little to these effects. Further studies are needed to
534 determine whether the butyrate effects on brain metabolism and hippocampal plasticity can
535 improve neurocognition and mental welfare, especially when brain and behavioural disorders
536 are observed. Different pig models are already available to investigate this question in the
537 context of early malnutrition or diet-induced conditions. This would pave the way to a new
538 treatment against nutritional- or stress-related psychological disorders.

539

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545

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752
753

523 **Table 1.** Brain structures presenting a higher (SB > C) or lower (SB < C) glucose metabolism (expressed in Standard Uptake Value) in butyrate-
524 supplemented piglets (SB group) compared to control piglets (C group) using a “without *a priori*” first-level global analysis and a “with *a priori*”
525 second-level small volume correction (SVC) analysis. The SVC analysis included the following regions of interest (ROI): anterior prefrontal
526 cortex, dorsolateral prefrontal cortex, orbitofrontal cortex, cingulate cortex, hippocampus, parahippocampal cortex, amygdala, caudate, putamen,
527 nucleus accumbens, globus pallidus. The peak *t*-value, uncorrected *P*-value and coordinates in the CA-CP (*commissura anterior-commissura*
528 *posterior*) reference plane are indicated for each significant cluster comprising at least 25 voxels, for the left (L) and/or right (R) hemispheres.

Brain structures	Side	SB > C			SB < C		
		Peak t	Peak <i>P</i> (unc)	x,y,z	Peak t	Peak <i>P</i> (unc)	x,y,z
First-level global analysis							
Primary somatosensory cortex	L				3.48	0.002	-14 19 18
	R	4.39	<0.001	8 15 25			
Inferior temporal gyrus	L				3.13	0.004	-26 5 6
Inferior colliculus	L	3.38	0.002	-8 -8 -2			
Hippocampus	R	3.35	0.003	12 3 -3			
Second-level SVC analysis							
Hippocampus	R	3.32	0.003	12 3 -4	2.67	0.010	10 -2 9
Nucleus accumbens	R				2.66	0.010	2 19 -4

529

530 **Table 2.** Gut electrophysiology parameters determined *ex vivo* in Ussing chambers (n=6-8 per treatment).

	<i>Site</i> Jejunum		Ileum		Colon		SEM	Statistics (P=) ¹			
	<i>Treatments</i> ² C	SB	C	SB	C	SB		Tr.	Site	Tr.*Site	
Basal condition ³											
Isc ($\mu\text{A}/\text{cm}^2$)	47	70	37	25	62	57	32	0.96	0.13	0.38	
TEER ($\Omega \times \text{cm}^2$)	50	49	57	51	46	28	10	0.25	0.44	0.79	
dIsc-glucose ($\mu\text{A}/\text{cm}^2$)	8	11	17	9	6	3	8	0.78	0.66	0.41	
dIsc-carbachol ($\mu\text{A}/\text{cm}^2$)	21	14	32	5	51	20	12	0.09	0.27	0.21	
Under oxidant stress (monochloramine, MC) ²											
dIsc-MC ($\mu\text{A}/\text{cm}^2$)	5	-55	6	5	-44	-115	36	0.29	0.19	0.63	
dIsc-glucose ($\mu\text{A}/\text{cm}^2$)	4.7	1.0	4.8	1.2	1.8	2.6	2.5	0.49	0.68	0.04	
dIsc-carbachol ($\mu\text{A}/\text{cm}^2$)	21	11	18	12	32	5	8	0.18	0.96	0.64	

531 ¹ Tr.: Treatment; Tr.*Site: treatment by site interaction.

532 ² Treatments: C: control; SB: sodium butyrate.

533 ³ Isc: short-circuit current; TEER: trans-epithelial electrical resistance; dIsc: variation in Isc (following glucose or carbachol or monochloramine addition).

534 **Table 3.** Influence of site and dietary treatment on gut para- and trans-cellular permeabilities determined ex vivo in Ussing chambers (log-
 535 transformed data; n=6-8 per treatment).

	<i>Site</i> <i>Treatments</i> ²	Jejunum		Ileum		Colon		SEM	Statistics (P=) ¹		
		C	SB	C	SB	C	SB		Tr.	Site	Tr.*Site
Basal condition ³											
FD4 (ng/cm ² /h, log)		2.9	2.9	3.1	2.8	3.0	2.9	0.10	0.15	0.73	0.15
HRP (ng/cm ² /h, log)		2.0	2.0	2.3	1.9	2.3	2.0	0.15	0.09	0.61	0.24
Under oxidant stress (monochloramine)											
FD4 (ng/cm ² /h, log)		2.8	2.9	3.1	2.9	3.1	2.9	0.09	0.49	0.34	0.49
HRP (ng/cm ² /h, log)		1.9	2.0	2.2	2.1	2.4	2.0	0.17	0.70	0.52	0.22

536 ¹ Tr.: Treatment; Tr.*Site: treatment by site interaction.

537 ² Treatments: C: control; SB: sodium butyrate.

538 ³ FD4: Fluorescein dextran 4000 (para-cellular permeability marker, 4 kDa); HRP: horseradish peroxidase (trans-cellular permeability marker, 40 kDa).

539

540

541 **Table 4.** Influence of dietary treatment on body weight, relative pancreas weight and protein concentration, and specific and total activities of the
 542 pancreatic enzymes (n=7-8 per treatment).
 543

Treatment	C	SB	SEM	Statistics (P=)
Body weight (Kg)	42.3	41.8	1.2	0.95
Fresh pancreas weight (mg/kg BW¹)	2.25	2.13	0.8	0.39
Pancreas protein				
mg/g fresh pancreas	207.7	201.5	12.6	0.77
mg/kg BW	468.2	431.8	37.4	0.88
Specific activities² (/g fresh pancreas)				
Trypsin (IU)	444.3	404.2	35.6	0.89
Amylase (10 ⁻³ IU)	7.2	7.5	0.8	0.37
Lipase (10 ⁻³ IU/)	79.8	81.1	9.5	0.45
Relative activities² (/kg BW¹)				
Trypsin (10 ³ IU)	208.7	175.3	21.8	0.44
Amylase (IU)	32.8	33.1	4.4	0.52
Lipase (IU)	38.0	34.2	5.8	0.69

544 ¹ Body weight.

545 ² IU = mmol substrate hydrolyzed/min.

546

547 **Table 5.** Influence of site and dietary treatment on enzyme activities along the gut (n=7-8 per treatment).

<i>Site Treatments</i> ²	Duodenum		Jejunum		Ileum		Colon		SEM	Statistics (P =) ¹		
	C	SB	C	SB	C	SB	C	SB		Tr.	Site	Tr.*Site
Peptidases												
APN (SA) ^{3,4}	55	48	201	159	136	125	/	/	16	0.17	<0.0001	0.59
APN (TA) ^{3,4}	5.7	5.4	19.5 ^a	14.8 ^b	13.4	13.8	/	/	1.1	0.09	<0.0001	0.06
DPP4 (SA)	26	25	26	21	12	15	/	/	2.5	0.65	0.0005	0.07
DPP4 (TA)	2.9	2.9	2.9	2.4	1.4	1.7	/	/	0.3	0.87	0.0005	0.11
Disaccharidases												
Sucrase (SA)	15	11	115	90	35	27	/	/	7.6	0.14	<0.0001	0.43
Sucrase (TA)	1.5	1.3	11.1	8.9	3.3	2.9	/	/	0.6	0.20	<0.0001	0.35
Maltase (SA)	84	78	334	282	129	91	/	/	22	0.15	<0.0001	0.31
Maltase (TA)	9.0	8.9	32.6	27.6	12.1	10.0	/	/	1.8	0.22	<0.0001	0.32
Alkaline Phosphatase												
IAP (SAC) ^{3,4}	1.5	2.1	3.4	2.9	3.5	3.6	1.8	2.0	0.4	0.86	<0.0001	0.69
IAP (TAC) ^{3,4}	105	143	229	169	210	214	70	71	26	0.81	<0.0001	0.32

548 ¹ Tr.: Treatment; Tr.*Site: treatment by site interaction.

549 ² Treatments: C: control; SB: sodium butyrate.

550 ³ APN: amino-peptidase N; DPP4: dipeptidyl-peptidase 4; IAP: intestinal alkaline phosphatase.

551 ⁴ SA: Specific activity (μmoles/min/mg protein; nmoles/min/mg protein for DPP4); SAC: Specific activity concentration for IAP (μg IAP/ g protein). TA: total activity

552 (μmoles/min/g fresh mucosa; nmoles/min/g fresh mucosa for DPP4); TAC: Total activity concentration for IAP (μg IAP/ g fresh tissue).

553 ^{a,b} P < 0.05

554

786 **Fig 1.** Changes in plasma concentrations of (A) lactate, (B) PYY and (C) GLP-1 before and
787 after drinking the beverages (both groups merged: pigs received either pure water (control) or
788 sodium butyrate-supplemented beverage). (D) Differences in plasma osmolality between
789 groups and sampling time. (E) Plasma lactate concentration a week later, just before brain
790 imaging.

791

792 **Fig 2.** Sagittal and axial magnetic resonance imaging models and sections showing the main
793 brain structures of differential glucose metabolism identified during the statistical parametric
794 mapping (SPM) analysis for the contrast butyrate vs. control. A) Three-dimensional brain
795 models showing the whole structures of interest. B) Magnetic resonance imaging sections
796 showing the significant clusters of brain activation or deactivation. The x or y coordinates in
797 the *commissura anterior-commissura posterior* (CA-CP) plane are indicated below the
798 images. The statistical threshold was set at $P < 0.01$ (uncorrected). Red spots indicate a
799 glucose metabolism higher in SB group compared to C group. Blue spots indicate a glucose
800 metabolism lower in SB group compared to C group. HIP: hippocampus; NAc: nucleus
801 accumbens; ISSC: primary somatosensory cortex.

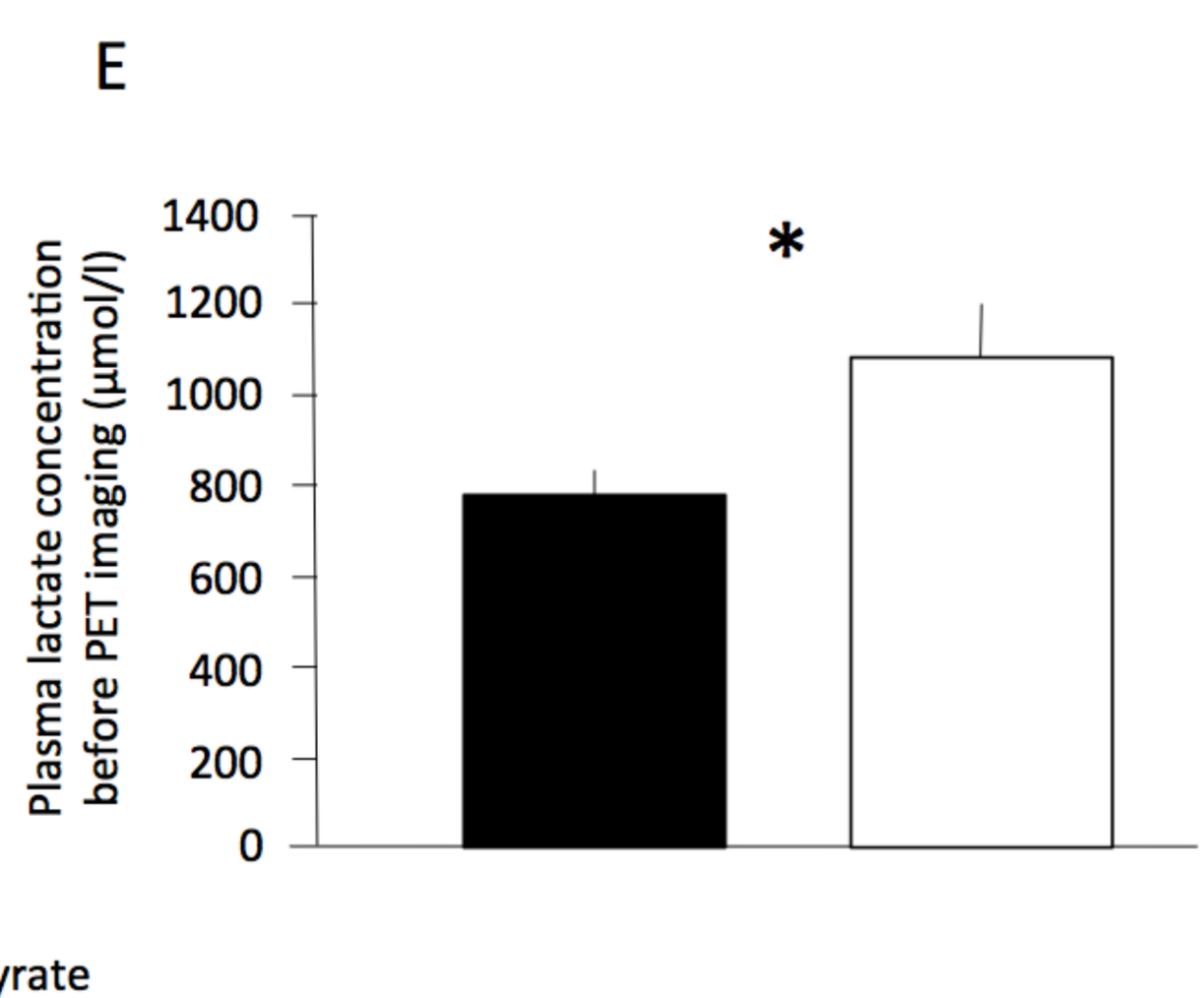
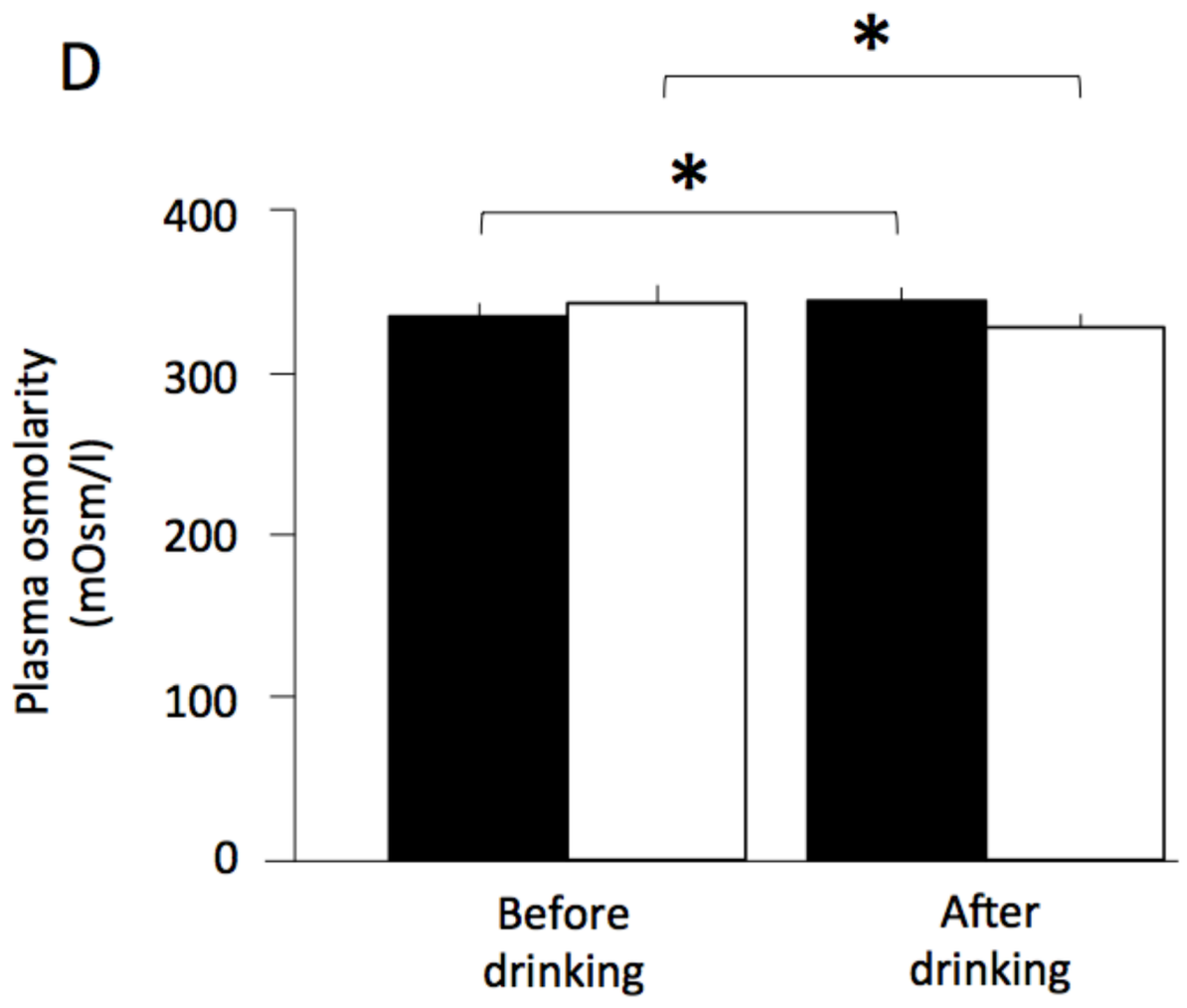
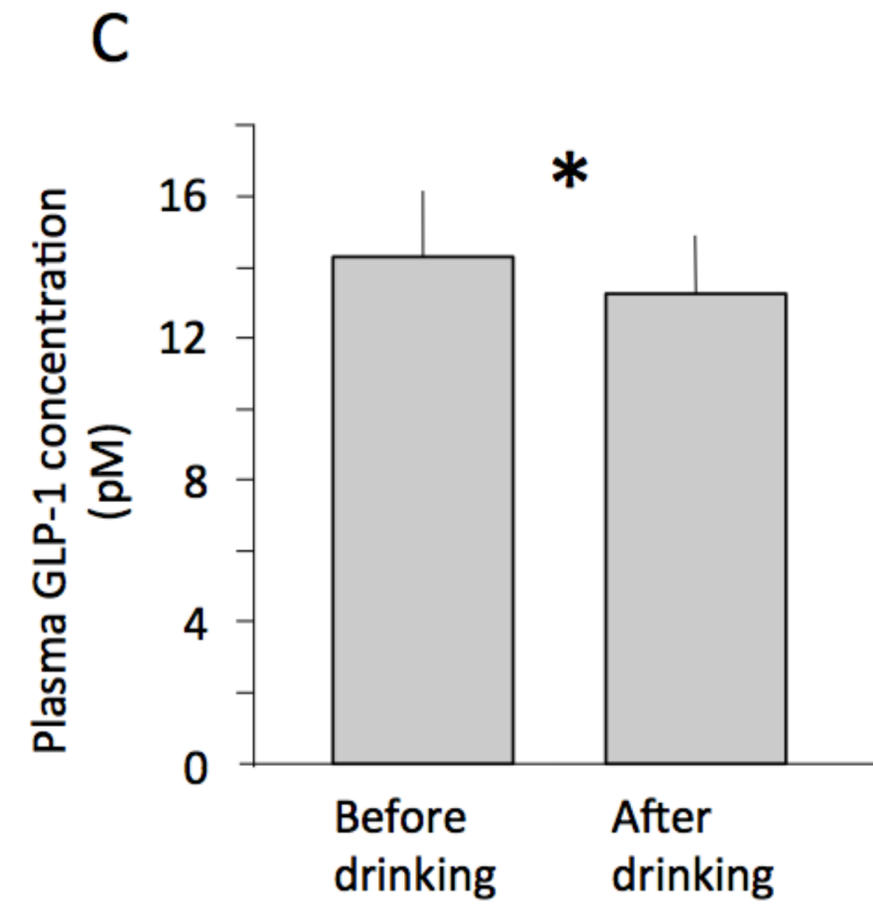
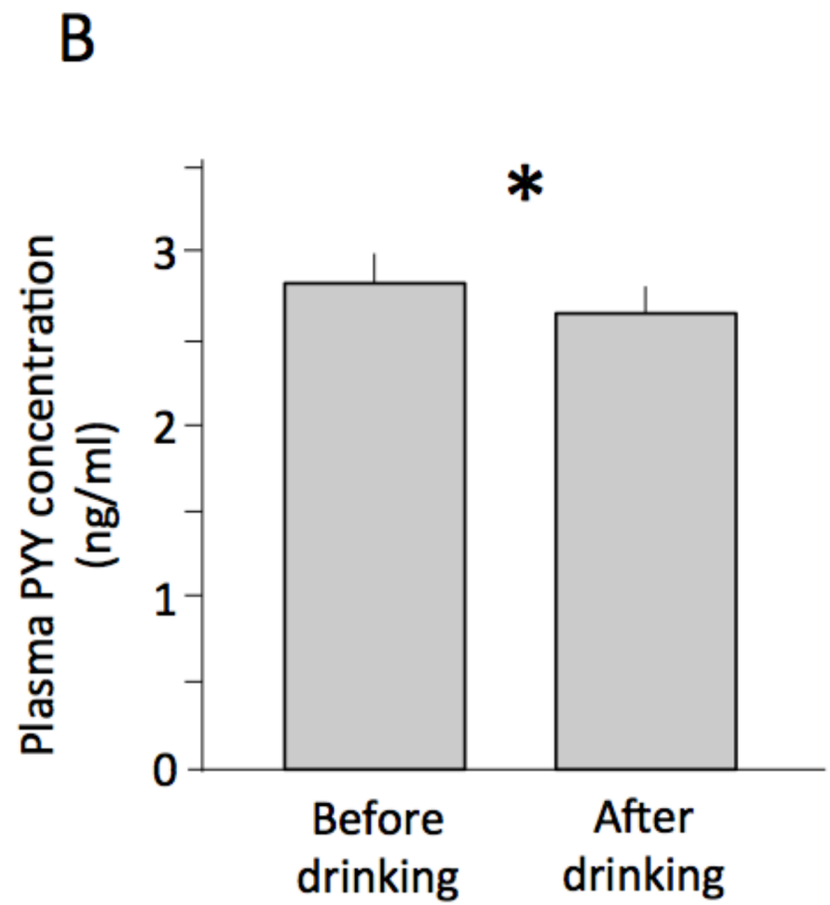
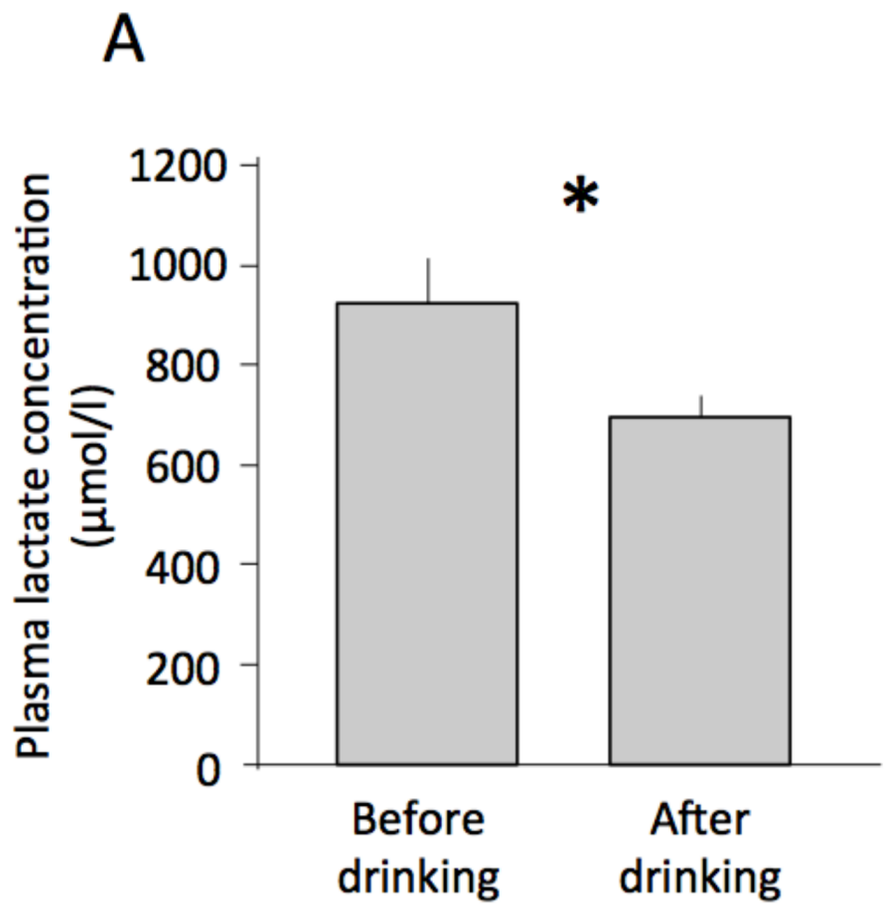
802

803 **Fig 3.** Hippocampal and granular cell layer (GCL) volume determination along the
804 hippocampus sections. A) Representative view of 10 sections covering the first 7.5 mm of
805 hippocampus. B) Calculated block volume for each site. The presented values are considering
806 the entire block of 25 sections of 30 μm each, and give an insight of the overall volume of the
807 hippocampus and the GCL. Significant difference between treatments was found for GCL
808 volume only. Data are presented as mean \pm SEM.

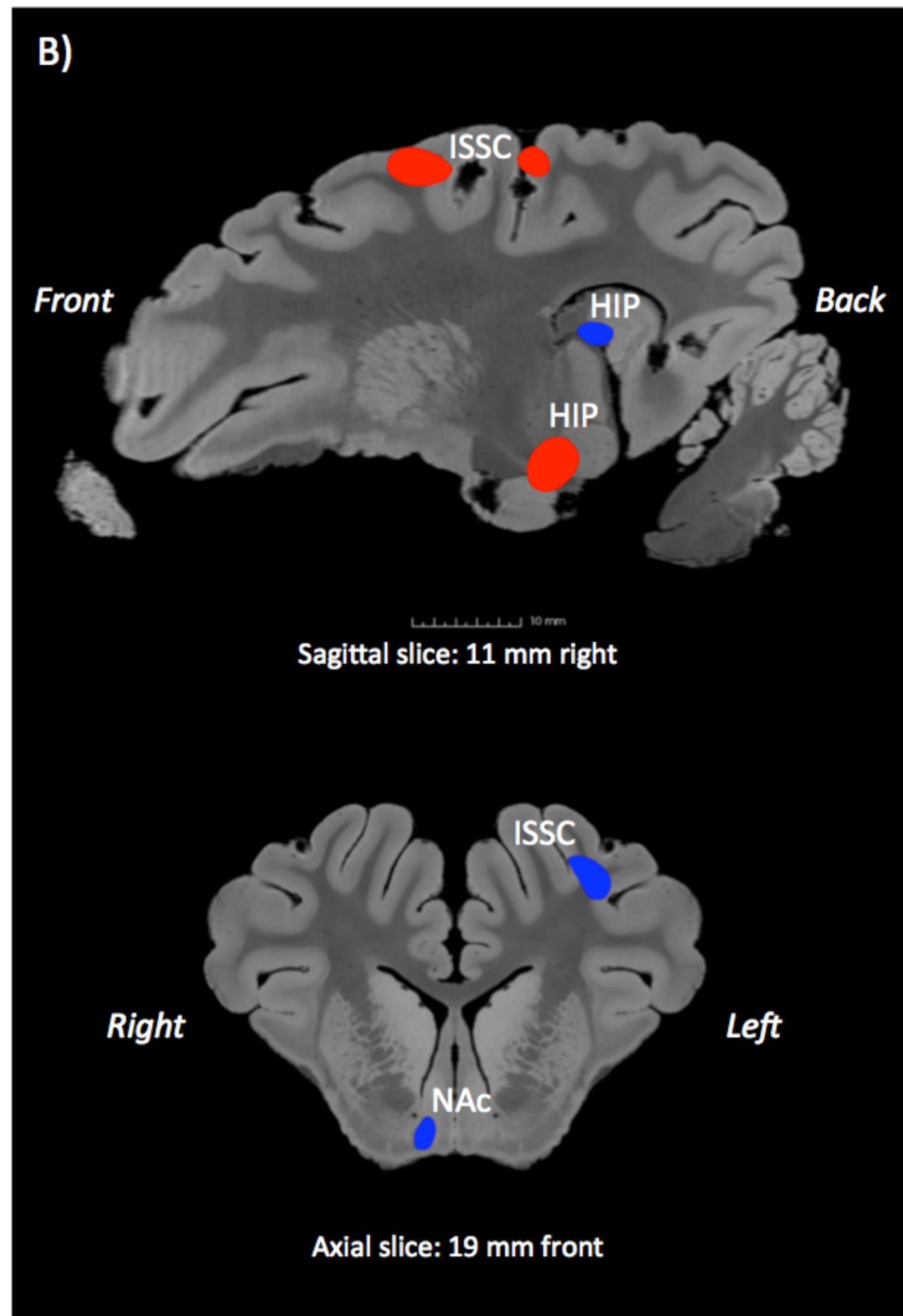
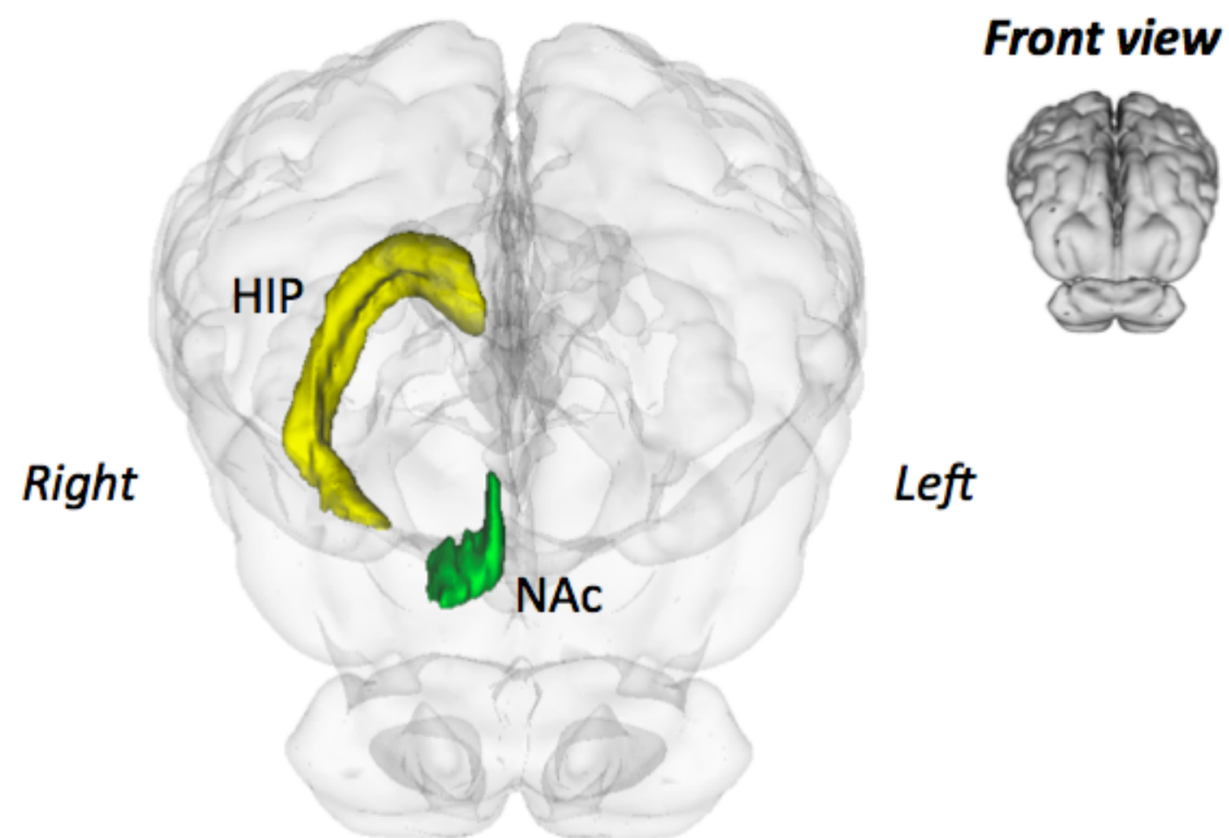
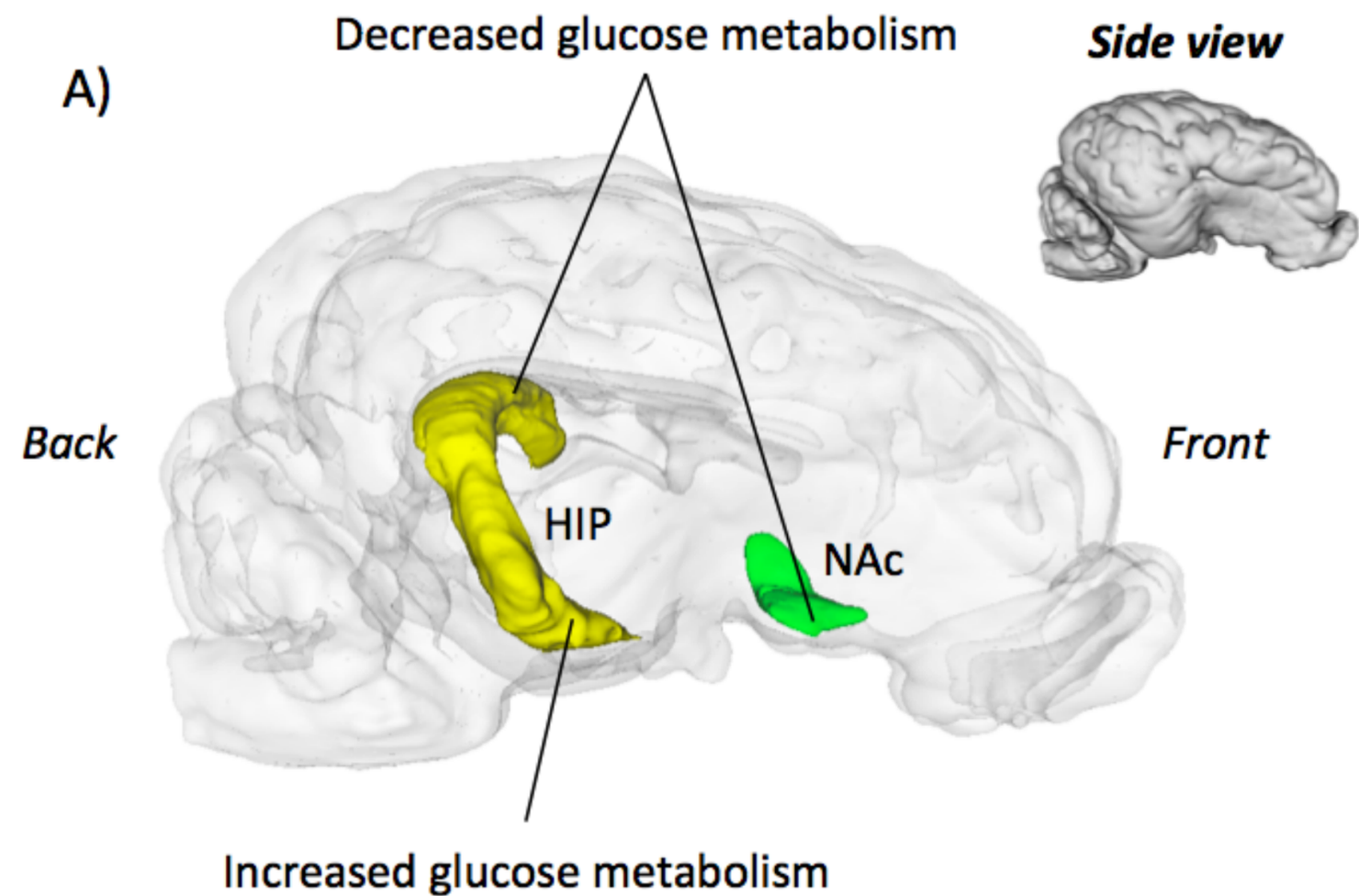
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810 **Fig 4.** Hippocampal structural plasticity. A) Immunostaining of brain sections showing Ki67-
811 positive nucleus distribution (left) and DCX-positive (doublecortin, right) cells within the
812 granular cell layer (GCL) of the hippocampus. B) Quantification of Ki67-positive nucleus in
813 the sub granular zone (SGZ, left) and DCX-positive cells in the GCL along the hippocampus.
814 The presented values are considering the entire block of 25 sections of 30 μm for each
815 histological slice. Significant differences were found between dietary treatments for Ki67 and
816 DCX. Data are presented as mean \pm SEM. An asterisk indicates a significant difference at
817 $P < 0.05$.

818

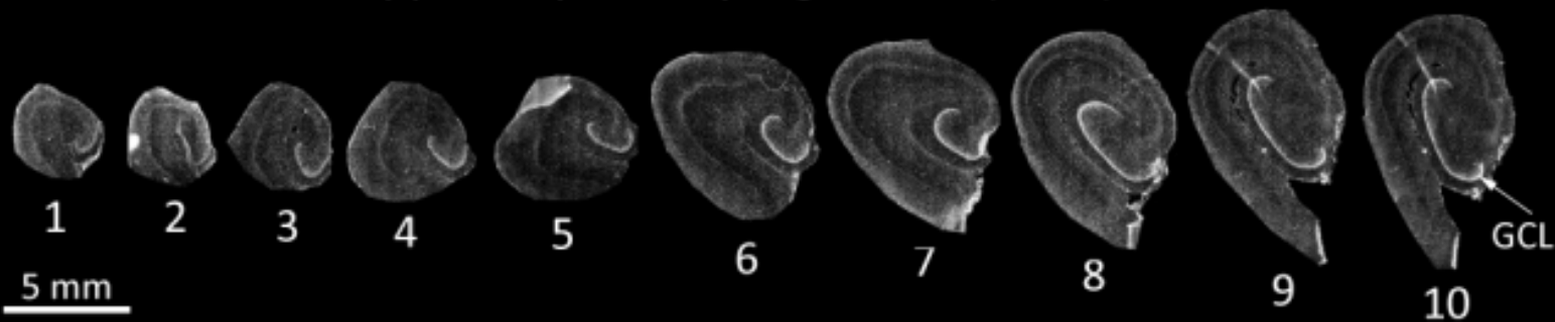
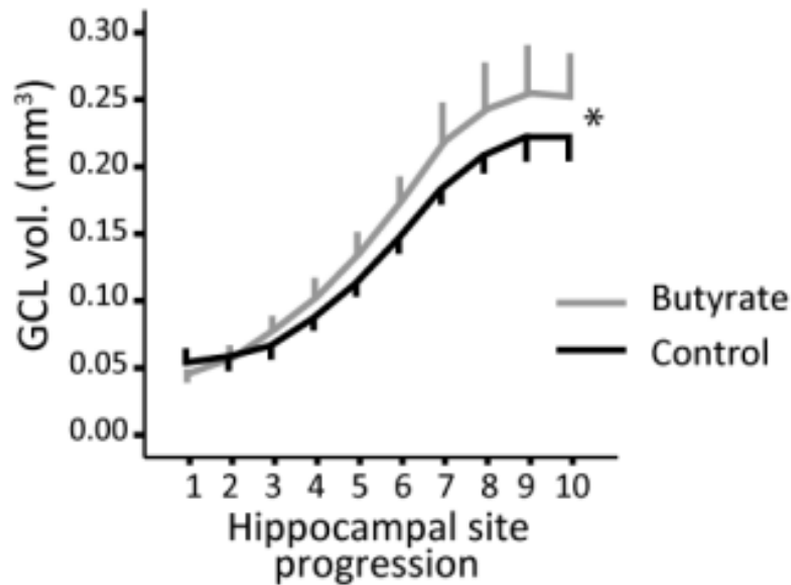
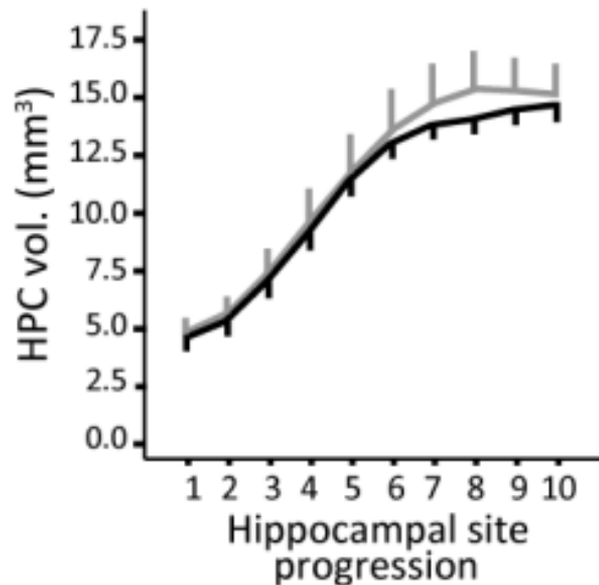


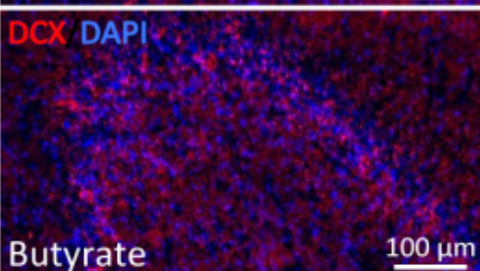
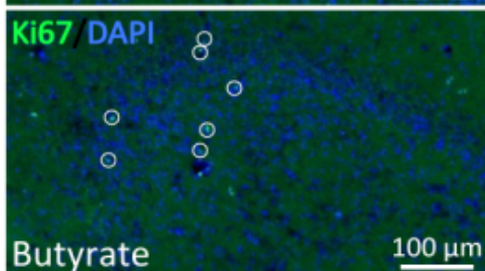
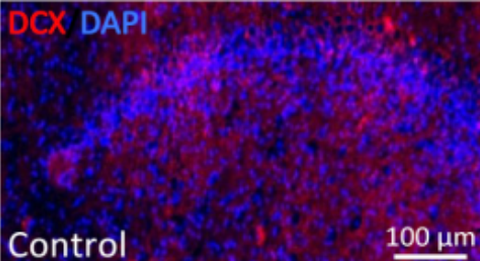
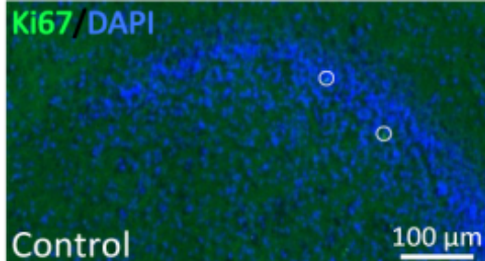
■ Butyrate
□ Control



A

Hippocampal site progression (DAPI)

**B**

A**B**