Phylogenetic diversity analysis of shotgun

² metagenomic reads describes gut microbiome

3 development and treatment effects in the

⁴ post-weaned pig

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19 Abstract

20 **Background**: Intensive farming practices can increase exposure of animals to infectious agents

21 against which antibiotics are used. Besides leading to antimicrobial resistance (AMR), orally

22 administered antibiotics are well known to cause dysbiosis. To counteract dysbiotic effects,

23 numerous studies in the past two decades sought to understand whether probiotics are a valid

24 tool to help re-establish a healthy gut microbial community after antibiotic treatment. However,

25 although dysbiotic effects of antibiotics are well investigated, little is known about the effects of

26 intramuscular antibiotic treatment on the gut microbiome and a few studies attempted to study

27 treatment effects using phylogenetic diversity analysis techniques. In this study we sought to

- 28 determine the effects of two probiotic- and one intramuscularly administered antibiotic treatment
- 29 on the developing gut microbiome of post-weaning piglets between their 3rd and 9th week of life.

30 Methods: Shotgun metagenomic sequences from over 800 faecal time-series samples derived 31 from 126 piglets and 42 sows were analysed in a phylogenetic framework to characterise the 32 developing gut microbial community composition of post-weaning piglets. We assessed the 33 effects of intramuscular antibiotic treatment and probiotic oral treatment on the diversity of these 34 gut microbial communities using alpha and beta diversity measures. 35 **Results**: Differences between individual hosts such as breed, litter, and age, were found to be 36 important contributors to variation in the community composition. Host age was the dominant 37 factor in shaping the gut microbiota of piglets after weaning. The post-weaning pig gut 38 microbiome appeared to follow a highly structured developmental program with characteristic 39 post-weaning changes that can distinguish hosts that were born as little as two days apart in the 40 second month of life. Treatment effects of the antibiotic and probiotic treatments were found but 41 were subtle and included a higher representation of Mollicutes associated with intramuscular 42 antibiotic treatment, and an increase of Lactobacillus associated with probiotic treatment. 43 **Discussion**: The discovery of correlations between experimental factors and microbial 44 community composition is more commonly addressed with OTU-based methods and rarely 45 analysed via phylogenetic diversity measures. The latter method, although less intuitive than the 46 former, suffers less from library size normalization biases, and it proved to be instrumental in 47 this study for the discovery of correlations between microbiome composition and host-, and 48 treatment factors.

49

50 Introduction

51 As the world population grows, there is an accompanying demand for animal-derived products. Intensive animal husbandry and early weaning practices are commonly used to maximize 52 53 production rates while minimizing costs. In a semi-natural environment, pig weaning occurs 54 between the 12th and the 17th week following birth whereas in intensively farmed pigs it typically 55 occurs at 3-4 weeks of age ^{1,2}. In intensive pig production early weaning practices increase the risk of enteric infections $^{3-6}$, and thereby the need for antimicrobial strategies, which has 56 57 included the common metaphylactic use of antibiotics ^{7,8}. Besides contributing to the concerning 58 issue of antimicrobial resistance (AMR) build up, antibiotic usage causes dysbiosis ^{9–11}, a 59 disruption of a balanced state within a gut microbial community, which increases the chance of

60 pathogens gaining a foothold (*i.e.*: colonization of the host or overgrowth) 12,13 . Researchers sought to determine the gut microbial modulatory effects of oral and in-feed antibiotic treatment 61 62 in the pig ^{10,11}, mouse ^{14–16}, and human ¹⁷, and only a few studies report the effects of intramuscular (IM) antibiotic treatment on the gut microbial community of the pig ^{18,19}. 63 Dysbiosis following antibiotic treatment of the young pig via intramuscular (IM)²⁰ and 64 subcutaneous ⁹ routes has been documented. It has been reported that a single dose of 65 intramuscularly administered amoxicillin in pigs, immediately after birth, causes long lasting 66 dysbiosis²¹, but no studies, to our knowledge, assessed the effects of IM neomycin treatment on 67 68 the post-weaning pig gut microbiota. IM antibiotic use has a lower risk for inducing AMR than 69 oral antibiotics in livestock production ^{22–25}, and according to a national survey of antibiotic use 70 in 197 large Australian commercial pig herds, 9.6% reported the use of injectable apramycin/neomycin⁷. For these reasons it is relevant to the industry to assess if, and to what 71

72 extent, IM neomycin causes dysbiosis. 73 Probiotics may play a role in counteracting the dysbiotic effects of antibiotics. Probiotic 74 treatment has been reported to affect the physiology of the host by improving mucosal integrity $^{26-29}$, inducing competitive exclusion against pathogenic species $^{30-35}$, reducing intestinal 75 inflammation ^{26,36,37}, and pathogen translocation ^{26,38,39}. In terms of microbial community 76 77 modulating potential of probiotics, studies in swine using culture based methods reported the reduction of pathogenic species ^{32,34,40–48} and an increase of *Lactobacillus* species ^{43,47–49} 78 79 following the administration of probiotic formulas. However, only in the past decade have 80 advances in and higher accessibility to high-throughput sequencing techniques enabled 81 researchers to better characterize microbial dynamics following probiotic treatment. Several of 82 these studies used the amplification of the bacterial 16S rDNA V4 region, determining the effects of probiotics on the gut microbial community of swine ^{50–53}. The choice of shotgun 83 84 metagenomic sequencing over amplicon sequencing, in this study, was dictated by the use of this 85 dataset outside of the scope described in this manuscript. In this study we assessed the effects of 86 two probiotic formulas on the gut microbial community of post-weaning piglets undergoing a 87 two-weeks probiotic treatment immediately after weaning. To assess the effects on the microbial 88 community of the administration of probiotic following antibiotic treatment, two treatment 89 groups were administered IM neomycin treatment followed by two weeks of either of the two 90 probiotic formulae.

The data was analysed from the perspective of the phylogenetic diversity of microbial communities. Firstly, we describe the phylogenetic diversity of microbial communities, and secondly, the role of age, breed and litter in microbial community composition. Finally, we describe the temporal effects on microbial communities, and the associated effects of IM

- 95 neomycin and probiotic treatments.
- 96

97 Materials & Methods

98

99 Animal trial

Metagenomic samples were derived from a pig study conducted at the Elizabeth Macarthur
 Agricultural Institute (EMAI) NSW, Australia, and approved by the EMAI Ethics Committee

(Approval M16/04). Below, we briefly summarise the origin of the samples, with comprehensive
 details on the animal trial and sample workflow being described previously ⁵⁴.

104 Post-weaning piglets (*n*=126) derived from a large commercial pig herd were transferred to

105 the study facility in January 2017. Piglets, aged 22.5±2.5 days at the start of the trial, consisted of

106 4 main cross-breed types: "Duroc × Landrace" (*n*=46), "Duroc × Large White" (*n*=59),

107 "Landrace × cross bred (LW×D)" (*n*=9), "Large White × Duroc" (*n*=12); and three pig lines

108 (line 319: n=9; line 316: n=46; line 326: n=71)⁵⁴. The study facility consisted of 4

109 environmentally controlled rooms (Rooms 1–4) with air conditioning, concrete slatted block

110 flooring with underground drainage, and open rung steel pens. The floor was swept daily and the

111 under-floor drainage was flushed twice weekly. A rubber ball was added to each pen for

environmental enrichment. Piglets were fed *ad libitum*, a commercial pig grower mix of 17.95%

113 protein, free of antibiotics.

114 The piglets were distributed over 6 treatment cohorts: a placebo group (Control *n*=29); two

115 probiotic groups (D-Scour n=18; ColiGuard n=18); one antibiotic group (Neomycin n=24) and

116 two antibiotic-then-probiotic treatment groups (Neomycin+D-Scour *n*=18;

117 Neomycin+ColiGuard n=18). The commercial probiotic paste preparation D-ScourTM was

sourced from International Animal Health, Australasia, and is composed of 1.8 10⁸ CFU/g of

119 each of the following: Lactobacillus acidophilus, Lactobacillus

120 delbrueckii subspecies bulgaricus, Lactobacillus plantarum, Lactobacillus rhamnosus,

121 Bifidobacterium bifidum, E. faecium, and Streptococcus salivarius subspecies thermophilus, with

122 an additional 20mg/g of garlic extract (*Allium sativum*). ColiGuard® is a probiotic formulation

123 developed for the treatment of entero-toxigenic *Escherichia coli* (ETEC) in weaner pigs,

124 developed in collaboration between the NSW DPI and International Animal Health Product, and

125 contains undefined concentrations of: L. plantarum and Lactobacillus salivarius. In addition, an

126 in-house developed mock community was used as a positive control, composed of *Bacillus*

127 subtilis strain 168, Enterococcus faecium, Staphylococcus aureus ATCC25923, Staphylococcus

128 epidermidis ATCC35983, Enterobacter hormaechei CP_032842, Escherichia coli K-12

129 MG1655, and *Pseudomonas aeruginosa* PAO1, in the following proportions:

130 8.7:13.0:7.7:16.7:38.9:14.5:0.4.

131 Groups of piglets were euthanized at the start (t0: n=6), a week after (t2: n=12), two weeks

132 after the start of the trial (t4: n=36) and at the end of the trial (t10: n=72) to obtain biopsy

133 samples that were used in another study. All surviving animals were euthanized at the end of the

animal trial. Euthanasia occurred as follows: the piglets were restrained in dorsal recumbency in

a bleeding cradle and pentobartitone euthanasia solution (Lethabarb, Virbac Australia, 325 mg

136 pentobarbitone sodium/mL) diluted 1:1 in sterile normal saline was administered via the precava

using a 20-21G needle depending on pig weight. The piglets received a dose of approximately 30

138 mg pentobarbitone/Kg to achieve deep anaesthesia, and then immediately exsanguinated.

139 Exsanguination after a non-lethal dose of barbiturate was undertaken to reduce excessive

140 congestion of the visceral blood vessels at the time of intestinal specimen collection (which is

141 likely to occur if euthanasia is undertaken using an overdose of barbiturate alone).

Faecal sampling occurred twice weekly for a subset of the piglets (8 per cohort; n=48), while it occurred weekly for all the living piglets (n=126; see euthanasia groups described above). In addition, 42 faecal samples derived from the piglets' mothers (n=42), 18 samples derived from three distinct positive controls (D-ScourTM, ColiGuard®, and mock community), and 20 negative controls were included.

A thorough description of the animal trial, and the metadata containing behaviour, weight,
and faecal consistency scores recorded over the 6-week period of the trial, is available in our
previous work ⁵⁴.

150

151 Metagenomic samples processing

- 152 Samples underwent homogenization and storage at -80 °C. After sample thawing, DNA
- 153 extraction was performed with the PowerMicrobiome DNA/RNA EP kit (Qiagen), and libraries
- 154 were prepared using the Hackflex method ⁵⁵. Sequencing was performed on three Illumina
- 155 NovaSeq S4 flow cells, after library normalization and pooling. The data is deposited to the
- 156 NCBI Short Read Archive under project PRJNA526405 and <u>http://dx.doi.org/10.5524/100890</u>⁵⁴.
- 157 Samples were assessed for quality using FASTQC
- 158 (http://www.bioinformatics.babraham.ac.uk/projects/fastqc/) and a combined report of all
- 159 samples was obtained with MULTIQC ⁵⁶ (available in our GitHub repository
- 160 <u>https://github.com/GaioTransposon/metapigs_phylodiv</u>).
- 161

162 **Determination of microbial diversity among samples**

Phylogenetic diversity of all samples was assessed with PhyloSift ⁵⁷ using the first 1M read 163 pairs of each sample (parameters: --chunk-size 1000000 --paired) (script: 164 *phylosift.nf*). In addition, a separate analysis with PhyloSift ⁵⁷ was performed using a smaller 165 166 downsampling, by including only the first 100K read pairs of each sample (parameters: --167 chunk-size 100000 --paired). In order to test for associations of phylogenetic 168 diversity with treatment, time of sampling, and differences among hosts at the start of the trial 169 (first post-weaning sample collection time point), analysis of the unrooted phylogenetic diversity (PD) ⁵⁸, the balance weighted phylogenetic diversity (BWPD) ⁵⁹ and principal component 170 analysis (PCA) of the Kantorovich-Rubinstein distances ⁶⁰ (beta diversity analysis) were 171 172 performed. Collections of phylogenetic placements produced by PhyloSift are compared using guppy ⁶⁰ to produce the Kantorovich-Rubenstein distances output. The collections of 173 174 phylogenetic placements are grouped based on the variable of interest (*i.e.*: treatment, time of 175 sampling, and differences among hosts at the start of the trial), and consequently the selection is 176 fed to guppy. Guppy is used in two modalities: 1. guppy epca to obtain edge principal 177 components; 2. guppy fat to annotate the edges of the phylogenetic tree using the relative 178 abundance of reads placed on each lineage. Alpha-diversity and beta-diversity were analysed and the results were visualized with R⁶¹ and R packages ^{62–76}. Scripts for the analysis are available at 179

180 <u>https://github.com/GaioTransposon/metapigs_phylodiv</u>. The data analysis workflow is

181 schematically represented in **Figure 1**.

182	All samples were included in the analysis, except for the analysis of treatment effect, where,
183	in order to minimize the effect of age as a confounding factor, we excluded animals born on
184	2017-01-06 and on 2017-01-07. In this manner the largest age gap between animals was 3 days,
185	with animals born on 2017-01-08 (n=24), 2017-01-09 (n=24), 2017-01-10 (n=23), and 2017-01-
186	11 (n=39). As a result, sample sizes at each time point were the following: Control (t0: 26; t2:
187	20; t4: 15; t6: 12; t8: 12; t10: 12), D-Scour [™] (t0: 15; t2: 15; t4: 15; t6: 10; t8: 12; t10: 10),
188	ColiGuard® (t0: 16; t2: 16; t4: 16; t6: 12; t8: 12; t10: 12), neomycin (t0: 22; t2: 22; t4: 17; t6: 12;
189	t8: 12; t10: 12), neomycin+D-Scour [™] (t0: 17; t2: 17; t4: 17; t6: 12; t8: 12; t10: 12), and
190	neomycin+ColiGuard® (t0: 14; t2: 13; t4: 14; t6: 10; t8: 10; t10: 10).
191	Analysis of raw reads with SortMeRNA ⁷⁷ (version 4.0.0) was performed as described in
191 192	Analysis of raw reads with SortMeRNA ⁷⁷ (version 4.0.0) was performed as described in Gaio <i>et al</i> (2021) ⁵⁴ . Briefly, raw reads were mapped against the rRNA reference database silva-
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192 193 194	Gaio <i>et al</i> (2021) ⁵⁴ . Briefly, raw reads were mapped against the rRNA reference database silvabac-16s-id90.fasta (parameters: $fastxblast 1num_alignments 1$) (script: <i>sortmerna.sh</i>) and filtered based on e-value (e-value <= 1e-30), identity (identity \geq 80%), and
192 193 194 195	Gaio <i>et al</i> (2021) ⁵⁴ . Briefly, raw reads were mapped against the rRNA reference database silvabac-16s-id90.fasta (parameters: $fastxblast 1num_alignments 1$) (script: <i>sortmerna.sh</i>) and filtered based on e-value (e-value <= 1e-30), identity (identity \ge 80%), and alignment length (length \ge 100 bp) (script: <i>sortmerna_filter.sh</i>). ⁵⁴ The output from
192 193 194 195 196	Gaio <i>et al</i> (2021) ⁵⁴ . Briefly, raw reads were mapped against the rRNA reference database silvabac-16s-id90.fasta (parameters: $fastxblast 1num_alignments 1$) (script: <i>sortmerna.sh</i>) and filtered based on e-value (e-value <= 1e-30), identity (identity \ge 80%), and alignment length (length \ge 100 bp) (script: <i>sortmerna_filter.sh</i>). ⁵⁴ The output from SortMeRNA ⁷⁷ was used to compute Principal Component Analysis (PCA) with R ⁶¹ (script:
192 193 194 195 196 197	Gaio <i>et al</i> (2021) ⁵⁴ . Briefly, raw reads were mapped against the rRNA reference database silvabac-16s-id90.fasta (parameters: $fastxblast 1num_alignments 1$) (script: <i>sortmerna.sh</i>) and filtered based on e-value (e-value <= 1e-30), identity (identity \ge 80%), and alignment length (length \ge 100 bp) (script: <i>sortmerna_filter.sh</i>). ⁵⁴ The output from SortMeRNA ⁷⁷ was used to compute Principal Component Analysis (PCA) with R ⁶¹ (script: <i>08_sortmerna.R</i>). Sample counts were normalized for library size by proportions and were tested

200

201 Batch effects

A randomized block design was adopted to mitigate batch effects. Because samples were
distributed across ten 96-well plates during DNA extraction and library preparation, plate effects
were expected. Although samples did not visibly cluster by DNA extraction plate across the first
five principal components, a batch effect was found by multiple comparison analysis with
ANOVA (alpha diversity: *p* range=0.0001-1; beta diversity: *p* value<0.0001) (Supplementary
Figure 1). Significance of pairwise comparisons was obtained by running Tukey's *post-hoc*analysis (Supplementary Table 1). Batch effects were removed with ComBat ⁷⁸ (script:

- 209 02_phylodiv.R). ComBat relies on robust empirical Bayes regression to remove heterogeneity
- 210 due to the use of batches (in this study a batch is a 96-well plate) while maintaining the
- biological variance among samples.⁷⁸ The input data used for ComBat in this study is not subject
- to normalization as it consists of phylogenetic diversity values obtained from a defined number
- 213 of read-pairs per sample. A parametric adjustment was used in the batch effect correction
- 214 (par.prior=TRUE) and both means and variances were adjusted (mean.only=FALSE).
- 215

216 Data availability

- 217 The raw sequencing data has been submitted to the NCBI Short Read Archive under project
- 218 PRJNA526405 and is also available through GigaDB at <u>http://dx.doi.org/10.5524/100890</u>. A
- thorough description of this dataset and the metadata has been reported previously ⁵⁴. Scripts
- 220 used for the data analysis in this study are available in our GitHub repository
- 221 <u>https://github.com/GaioTransposon/metapigs_phylodiv</u>.
- 222

223 **Results**

PhyloSift ⁵⁷ was employed as a means to study microbial community diversity among the
samples, and to test for associations with treatment, time of sampling, and differences among
hosts during the first week post-weaning. To this end, analysis of the unrooted phylogenetic
diversity (PD) ⁵⁸, the balance weighted phylogenetic diversity (BWPD) ⁵⁹, and principal
component analysis (PCA) of the Kantorovic-Rubinstein distances ⁶⁰ (beta diversity analysis)
were performed.

230

231 Phylogenetic diversity of positive controls

Unrooted PD was highest for positive control D-Scour[™] (mean±SD=162.2±28.3), slightly lower

- for positive control ColiGuard® (mean±SD=129.9±50.0) and lowest for the mock community
- 234 (mean±SD=123.6±28.4). BWPD was highest for positive control D-Scour[™] (mean±SD:
- 1.9±0.2), lower for the mock community (mean±SD: 1.6±0.1), and lowest for positive control
- 236 ColiGuard® (mean±SD: 0.9±0.1). (Supplementary Table 1; Supplementary Figure 2).

237 Principal component analysis (PCA) of the Kantorovich-Rubenstein distances (beta 238 diversity analysis) was performed on positive control samples. Samples clearly separated in PC1 239 (accounting for 83.62% of the variation), where mock community samples clustered in the lower 240 end of principal component 1 (PC1) showing a higher representation of Enterobacteriaceae and 241 Pseudomonadaceae, while ColiGuard[®] and D-Scour[™] samples clustered in the higher end of 242 PC1, showing a higher representation of *Lactobacillales*. D-Scour[™] samples clustered in the 243 higher end of principal component 2 (PC2) (accounting for 14.52% of the variation), showing a 244 higher representation of Lactobacillus delbrueckii, Enterococcus faecium, and Bifidobacterium, 245 while in the lower end of PC2, mock community samples and ColiGuard® samples, forming two 246 separate clusters, showed a higher representation of *Bacilli*, *Enterobacteriaceae*, and 247 Gammaproteobacteria. (Supplementary Figure 3).

248

249 *Phylogenetic diversity of piglet gut communities: host factors*

250 Based on Kruskal-Wallis one-way analysis of variance (Hommel adjusted p values to correct for

251 multiple hypotheses testing), alpha diversity of the piglet samples did not cluster significantly by

cross-breed type (p > 0.05) or by pig line (p > 0.05) in the first week post-weaning (Figure 2;

253 Supplementary Table 1). A correlation of cross-breed type was found with beta diversity

254 (principal component 3) at t2 (*p*=0.024). (Figure 2; Supplementary Table 1)

255 The piglets differed slightly by age, being born between 1 and 5 days apart. Notably, we 256 found a significant correlation between alpha diversity and the age of the piglets at the first 257 sampling time point (unrooted PD: p < 0.0001; BWPD: p=0.011) (Figure 2; Supplementary 258 Table 1) and between age and beta diversity at t2 (PC3: p=0.047) and t7 (PC3: p=0.018) (Figure 259 2; Supplementary Table 1). As age groups were confounded with cross-breed types (*i.e.* not all 260 age groups are represented by each of the four cross-breed types), we compared the phylogenetic 261 diversity of age groups within each breed. As cross-breed types "Landrace \times cross bred (LW \times 262 D)" and "Large White \times Duroc" had only a small number of piglets in each age group (n=9 and 263 n=12, respectively), we tested for an association between phylogenetic diversity and age in cross-breeds "Duroc \times Landrace" and "Duroc \times Large White" (*n*=46 and *n*=59, respectively). 264 265 Among these cross-breeds, age in the "Duroc \times Landrace" piglets (n=46) correlates with alpha diversity during the first week post-weaning (unrooted PD: p=0.006; BWPD: p=0.047) and with 266

beta diversity at t2 (PC2: *p*=0.048) (Figure 2; Supplementary Figure 4; Supplementary Table
1).

269 As piglets were derived from 42 distinct sows (maternal sows) and nursed by either the 270 same or a different sow (a nurse sow), a litter effect was expected and determined. Based on 271 Hommel adjusted p values, a similarity of alpha phylogenetic diversity can be seen among 272 piglets from the same maternal sow (unrooted PD: p=0.001; BWPD: p=0.017) and in piglets 273 from the same nurse sow (unrooted PD: p=0.002; BWPD: p=0.027) (Figure 2; Supplementary 274 **Table 1**). The litter effect described was found significant only in samples from the first week 275 post-weaning as the significance of the correlations did not persist thereafter (Figure 2; 276 Supplementary Table 1).

277

278 A strong effect of aging on phylogenetic diversity

279 Beta diversity analysis of all samples revealed a distinct and consistent change of the microbial 280 community over time in all piglets, regardless of the treatment. Beta diversity analysis was 281 performed from all reads and from the analysis of 16S rDNA V4 region-containing reads. 282 Samples collected immediately after weaning (t0) were characterized by a higher representation 283 of Bacteroidetes chlorobi group and Clostridia (PC1, 47.68%), particularly of Sedimibacterium 284 and *Desulfosporosinus* (PC2 17.2%), respectively. Between day 0 (t0; immediately after 285 weaning) and day 14 (t4), samples shifted towards a higher representation of *Bifidobacterium* 286 and Lactobacillus, as measured from beta diversity analysis (PC1; 47.68% var. explained) and 287 from analysis of 16S rRNA reads (PC1 and PC2; 23.9% and 17.2% var. explained, respectively). 288 During the last two weeks (t6-t10), corresponding to day 21 and 35 of post-weaning, samples 289 shifted towards a higher representation of Actinobacteria, (PC2 21.79% var. explained), 290 particularly of Collinsella (PC1 23.9%, PC2 17.2%). Lineages of the Erysipelotrichales 291 *Catenibacterium* and *Solobacterium* were also found to be representative of samples of this time 292 interval (t6-t10). (Figure 3).

Beta diversity analysis was performed separately for samples within each time point in order to find lineages associated with variation within each time point. Extent of variation was derived from the product of branch width by the variation explained by the principal component (**Figure** 4). The lineages *Enterobacteriaceae* (t0=0.05) and *Bacteroides* (t0=0.06) were responsible for variation only during the first week after weaning, and *Methanobrevibacter smithii* during the first and the second week (t0=0.05; t2=0.03). The *Bifidobacterium* lineage was responsible for variation in the second week (t2=0.09). The following lineages were responsible for variation throughout the 6 weeks after weaning: *Bacteroidales* (min=0.04; max=0.20), *Prevotellaceae* (min=0.02; max=0.13), *Coriobacteriaceae* (min=0.01; max=0.09). *Lactobacillus* became variable after the first week post-weaning and remained highly variable throughout the rest of the

303 trial (min=0.03; max=0.30) (**Figure 4**).

304 Taxonomic representation in terms of relative abundance was derived from the branch width

305 of the phylogenetic tree (**Supplementary Figure 5**) and combined with PhyloSift's taxonomic

306 annotation of the marker gene phylogeny. Lactobacillus acidophilus increased at the start

307 (t0=0.00; t2=0.07; t4=0.17) then decreased (t6=0.11; t8=0.06; t10=0.08). Among other prevalent

308 lineages during the week after weaning, up to the next week and dropping at later time points, we

found *Methanobrevibacter smithii* (t0=0.05; t2=0.03) and *Bacteroidales* (t0=0.01; t2=0.01).

Following an opposite trend we found *Ruminococcus* sp. JC304 (t0=0.00; t2=0.00; t4=0.01;

311 t6=0.03; t8=0.03; t10=0.03), *Solobacterium moorei* (t0=0.00; t2=0.01; t4=0.02; t6=0.03;

312 t8=0.02; t10=0.03) and *Prevotella copri* (t0=0.00; t2=0.00; t4=0.05; t6=0.03; t8=0.06; t10=0.03).

- 313 In modest and stable abundance across the post-weaning period were the following lineages of
- the order of the *Clostridiales: Mogibacterium* sp. CM50 (t0=0.03; t2=0.05; t4=0.03; t6=0.04;

315 t8=0.03; t10=0.04), *Oscillibacter* (t0=0.08; t2=0.06; t4=0.04; t6=0.03; t8=0.04; t10=0.03),

316 Subdoligranulum variabile (t0=0.07; t2=0.03; t4=0.05; t6=0.07; t8=0.07; t10=0.08), and

317 *Ruminococcus bromii* (t0=0.03; t2=0.02; t4=0.01; t6=0.01; t8=0.1; t10=0.01). In transient

318 abundance we found *Bifidobacterium thermophilum* RBL67 (t2=0.02; t4=0.03). Gradually

increasing from the second week we found *Eubacterium biforme* DSM3989 (t2=0.02; t4=0.02;

320 t6=0.03; t8=0.03; t10=0.02), *Eubacterium rectale* (t4=0.03; t6=0.03; t8=0.03; t10=0.01) and,

- 321 after the third week, *Faecalibacterium prausnitsii* (t6=0.01; t8=0.01; t10=0.01).
- 322 (Supplementary Figure 5)

323 The effect of time was also clear in alpha diversity, where all the piglet cohorts followed a

- 324 similar trend over time, independent of the treatment (**Supplementary Figure 6**;
- 325 Supplementary Table 1). Immediately after weaning (t0), the unrooted PD was lower for the

326 piglets than the sows (sows: 328.5±24.0; piglets: 296.5±34.7) and reached a higher unrooted PD

- 327 to the sows' in the following week (t2) (piglets: 336.18±33.0) (Supplementary Table 1). In
- 328 comparing four timepoints at one-week intervals from the start of the trial, changes in alpha
- 329 diversity among all the piglets were tested for and significance was determined using the
- 330 Bonferroni correction. Unrooted phylogenetic diversity increased in the first week following
- 331 weaning (t0-t2: +9.15%; *p*<0.001) and decreased in the following week (t2-t4: -4.54%; *p*<0.001)
- amongst the piglets. In contrast, BWPD decreased in the week after weaning (t0-t2: -6.07%;
- p<0.001), to increase in the following week (t2-t4: +4.77%; p<0.001) and decrease in the fourth
- 334 week (t4-t6: -3.67%; *p*=0.002). (Supplementary Figure 6; Supplementary Table 1).
- 335

336 Effect of antibiotic and probiotic treatment on alpha diversity

We hypothesized that the probiotic treatments, whether alone (D-Scour[™] and ColiGuard[®]) or 337 338 administered after neomycin (neomycin+D-Scour[™] and neomycin+ColiGuard®) would cause a 339 change in the microbial community composition that would be measurable via phylogenetic 340 diversity. We tested whether the treatments correlated with a change in phylogenetic diversity 341 independently of the changes occurring with time. Given the differences in alpha and beta 342 diversity detected among the subjects after weaning, we analyzed the deltas of phylogenetic 343 diversity instead of relying on the absolute means, similar to the procedure applied by Kembel et 344 al (2012) ⁷⁹. Time-point measurements of alpha diversity were taken, and deltas were computed 345 for each piglet. Delta means were compared between cohorts, where the control cohort would 346 serve as a control group for neomycin, D-Scour[™] and ColiGuard[®] cohorts, whereas the 347 neomycin cohort would serve as a control group for the neomycin+D-Scour[™] and 348 neomycin+ColiGuard® cohorts.

One week after weaning, 90% of the piglets displayed an increase in unrooted PD and 71% displayed a decrease of BWPD. The following week the trend was opposite: 72% of the piglets displayed an increase of BWPD and 76% displayed a decrease of unrooted PD (**Supplementary Table 1**). However, the neomycin cohort displayed the smallest BWPD drop in the week following weaning, and the overall trend of neomycin in unrooted PD was the most different from the other cohorts (**Supplementary Figure 6**). Due to the lower drop in BWPD of neomycin, significance was found in BWPD during the first week between neomycin and

- 356 Neomycin+ColiGuard® (t0-t2; Tukey adjusted *p* value=0.031), and between neomycin and
- 357 ColiGuard® (t0-t2; Tukey adjusted *p* value=0.041). No other significant differences were found
- in changes in alpha diversity between time points among the cohorts. (Supplementary Table 1)
- 359

360 Effect of antibiotic and probiotic treatment on beta diversity

361 To investigate the treatment effect on beta diversity, principal component analysis (PCA) of the 362 Kantorovich-Rubenstein distances (beta diversity analysis) was performed on all samples and, 363 additionally, on samples within individual time points. This analysis is conceptually similar to 364 the weighted Unifrac approach for beta diversity analysis, but is designed to work with 365 phylogenetic placement data ⁶⁰. When examining all samples together, there was no clear 366 separation of cohorts on any of the five principal component axes. When individual time points 367 were analysed, some clustering by cohort was observed (Supplementary Figure 7). D-Scour[™] 368 separated from the control cohort in PC3 (10.97%) during the first week of probiotic treatment 369 (t2) (Supplementary Figure 7). ColiGuard® separated from control during the first week of 370 probiotic treatment (t2) in PC5 (2.82%), and two weeks after the end of probiotic treatment in 371 PC3 (7.87%), showing a higher representation of *Lactobacillus*.

Two weeks after the end of neomycin treatment (t6), neomycin separated from control in PC4 (5.65%), with a smaller clustering and a higher representation of *Mollicutes*. Neomycin+D-ScourTM separated from neomycin in PC2 (17.80%) and PC4 (5.65%) at the end of probiotic treatment (t6), as well as a week later (t9) in PC5 (3.93%). In these instances, neomycin clustered towards a higher representation of *Mollicutes*, while Neomycin+D-ScourTM showed a higher representation of *Lactobacillus*. (**Supplementary Figure 7**)

378

379 Association between weight and community composition

- 380 Weight correlated with the abundance of certain taxa at each time point as it resulted from
- 381 principal component analysis of 16S rRNA reads. Positively correlating with weight we found
- 382 among others: *Blautia* (t0), *Cetobacterium* (t0), *Lactobacillus* (t6), *Mycoplasma* (t6),
- 383 Anaerostipes (t8), Ruminococcus (t8), Cerasibacillus (t10). Negatively correlating with weight
- 384 we found among others: *Pyramidobacter* (t0), *Odoribacter* (t2), *Schwartzia* (t6), *Streptococcus*

- 385 (t6), *Dokdonella* (t8). Correlations remained significant after p value correction for multiple
- 386 hypotheses testing (Hommel, and Benjamini & Yekutieli), but only one taxa
- 387 (Coprothermobacter; t0) remained significantly, positively correlated with host weight after
- applying the more stringent Bonferroni correction (**Supplementary Table 1**).
- 389

390 Effect of treatments on weight gain

- 391 Overall weight gain from initial to final weight (**Supplementary Table 1**) was not significantly
- 392 affected by any treatment. However, the probiotic ColiGuard® was found to have a partial effect
- 393 on piglet weight gain (Supplementary Figure 8; Supplementary Table 1). Weight was
- 394 measured weekly for a total of six measurements. To minimize age as a confounding factor from
- this analysis, we kept samples from piglets that were born a max of 3 days apart. Based on Tukey
- adjusted p values, a lower weight gain was detected in the ColiGuard® cohort compared to the
- 397 control cohort between the last day of probiotic treatment and a week after treatment (t4-t6)
- 398 (*p*=0.015) (**Supplementary Figure 8**; **Supplementary Table 1**). Similarly, a lower weight gain
- 399 was detected in the neomycin+ColiGuard® cohort between the 9th day of treatment and the 9th
- 400 day after probiotic treatment (t4-t8) compared to the neomycin cohort (p=0.011)
- 401 (Supplementary Figure 8; Supplementary Table 1). Age and cross-breed differences among
 402 piglets were not associated with weight gain (Supplementary Table 1).
- 403

404 Estimation of phylogenetic diversity using sample downsampling sizes: 100K 405 versus 1M

406 We compared unrooted PD and BWPD values obtained using either 100,000 or 1,000,000 reads

- 407 per sample (corresponding to 0.612% or 6.12% of the average sample). The median unrooted PD
- 408 was 2.6x higher, and variance improved when the analysis was run from 1M reads per sample,
- 409 compared to the analysis run using 100K reads per sample (100K: 121±18.67; 1M: 317±50.67;
- 410 Pearson's r=0.8965), while the median BWPD remained nearly unchanged (100K: 2.11±0.203;
- 411 1M: 2.11±0.193; Pearson's r=0.9747). Comparing beta diversity values obtained using either
- 412 100,000 or 1,000,000 reads per sample, we found a high correlation for all five principal
- 413 components (Pearson's r range=0.9809-0.9956). (Supplementary Figure 9)

414

415 **Discussion**

416 The microbial composition of positive control samples was analysed from a taxonomic 417 perspective in our previous study ⁵⁴, and it was here compared to the phylogenetic diversity 418 obtained for these samples. The unrooted PD reflects the absolute diversity, independently of the 419 relative abundance of each species, within a sample. In fact, low-level contamination (<0.1%) detected in each of the positive controls (mock community: 1 taxon; ColiGuard®: 20 taxa; D-420 421 Scour[™]: 25 taxa) ⁵⁴ contributes toward the absolute diversity in the unrooted PD, inflating this 422 value. Unrooted PD values obtained for the positive controls were directly proportional to their 423 respective count of taxa, which include contaminants (D-ScourTM: n=33; ColiGuard®: n=22; 424 mock community: n=8)⁵⁴. On the other hand, the contribution to total diversity of phylogenetic 425 tree edges with uneven quantities of reads placed on either side is down-weighted in BWPD. The 426 fact that ColiGuard[®], mock community, and D-Scour[™] are mainly composed of 2, 7, and 8 taxa 427 taking up even proportions in the samples, respectively (hence excluding contaminants)⁵⁴, was 428 reflected by their lowest, higher, and highest BWPD measured from these samples.

429 The consistent trend in community composition over time, across all the cohorts, 430 indicates that an age-related process of ecological succession is the largest factor shaping the 431 microbial community of post-weaning piglets, as found in this study where animals aged 20-63 432 days were fed the same diet. A peak in unrooted phylogenetic diversity and drop in balance 433 weighted phylogenetic diversity (BWPD) reflects the acquisition of new species with the loss of 434 dominating species. Similarly to Pollock *et al* (2018)⁸⁰, we found the relative abundance of 435 Clostridiales, and Lachnospiraceae to decrease from the start of weaning to 2 weeks into 436 weaning, while the relative abundance of Lactobacillus, Prevotella copri, Faecalibacterium prausnitsii and Erysipelotrichaceae increased. The increase of Prevotella with weaning 80-82 is 437 438 well documented and it is associated with the increased polysaccharide consumption associated 439 with the start of solid food consumption^{81,83}. The relative abundance of *Lactobacillus* goes up with weaning, as other studies suggested ^{80,81}. However, according to our analysis, this increase 440 441 concerns Lactobacillus acidophilus, while other Lactobacillus species, such as Lactobacillus 442 *vaginalis* ATCC follow the opposite trend. We also found a clear gradual increase in abundance

443 of Solobacterium moorei and Eubacterium biforme DSM3989, a transient increase of

444 Bifidobacterium thermophilum RBL67 during the second and the third week post weaning, and a

sharp decrease in the abundance of *Methanobrevibacter smithii* after the second week post-weaning.

447 The change in phylogenetic diversity detected in the week following the piglets' arrival at 448 the trial site irrespective of the cohort, could be linked to the piglets being subjected to microbial 449 interchange (e.g.: new pen mates) and/or to diet transition (peri-weaning transition) to solid food leading to the reshaping of the gut microbial community.^{84,85} The week following the drop of 450 451 BWPD, a significant increase of BWPD was recorded, reflecting the acquisition of a larger 452 proportion of the community by the newly introduced species. The strong changes in 453 phylogenetic diversity detected in the first and the second week could as well be attributable to 454 other post-weaning related physiological changes, as previous studies report $^{84-87}$.

The highest inter-individual differences among piglets are seen in the first week of life, irrespective of maternal or environmental effects. The microbiota of three week old piglets is still very dynamic, but environmental factors become evident ⁸⁵. At six weeks of age, CD8+ T cells infiltrate the intestinal tissue and the mucosa and intestinal lining resemble that of an adult pig ⁷⁹.

459 The increase of alpha diversity associated with weaning has been measured before ^{81,82,88}, but 460 rarely using metrics that allow the distinction of absolute diversity from evenness. In this study, 461 one week after weaning, piglets reached a comparable absolute diversity to the sows, at which 462 time the piglets were aged between 3.8 and 4.6 weeks. Unrooted PD did not reach higher levels 463 at later sampling time points. The highest BWPD accompanied by a high unrooted PD was 464 reached after the second week post-weaning when piglets were aged between 4.9 and 5.6 weeks. 465 Age-dependent physiological changes could explain i) the major shifts we detected in alpha 466 diversity during the first two weeks after weaning and, ii) the distinct differences in community 467 composition with age, even with a narrow age difference between piglets (1-5 days). We found a 468 significant difference in microbial composition in both absolute diversity as in balance-weighted 469 PD, only in the first week after weaning (piglets aged between 3.8 and 4.6 weeks), between 470 groups of piglets that were separated by up to five days maximum by day of birth. Since age 471 groups were confounded with breeds in our study, we attempted to determine the correlation

within single cross-breeds. Unfortunately, although the correlations with age could still be
detected, we could not determine the association at later time points due to the introduction of
treatment effects.

475 Animal trials are often conducted in controlled environments to minimize environmental 476 effects. However, individual variations such as breed and age are often unavoidable in large 477 animal trials, especially involving animals derived from commercial herds. Previously reported 478 confounding factors include: individual variation, cohabitation, age, maternal effects, hormones, 479 behavioural differences between breeds (e.g. coprophagy, mouth to mouth contact) and extent of 480 long-term behavioural adaptation, which can differ between breeds for reasons not attributable to 481 genetics ^{84–86,89}. A litter effect was found in piglets at the start of the trial (piglets aged between 482 3.8 and 4.6 weeks; samples collected immediately after weaning) and was lost at later time 483 points during the trial. This could be due to either of the aforementioned factors. Co-housing, 484 aging and the splitting of the piglets in separate rooms to receive a different treatment, are 485 possible causes for loss of the litter effect. In this study we confirm the importance of these 486 factors in the contribution to inter-individual variability of gut microbial composition. Motta et al 487 (2019) report a correlation of beta diversity with age and no correlation of genotype and litter effect with either alpha or beta diversity ⁹⁰. On the contrary, we found the piglet samples to 488 489 significantly cluster by age and litter immediately after weaning in alpha diversity, and by age 490 and cross-breed in beta diversity during the second week after weaning. Samples also clustered 491 by age groups in beta diversity four weeks after weaning. The groups we tested, based on age, 492 cross-breed, and litter, differed in sample sizes, therefore a non-parametric test was used to test 493 for associations of host factors with phylogenetic diversity. Based on our results we conclude 494 that even small age differences among post- wearing piglets, down to the day, must be 495 accounted for in an experimental set up, but we acknowledge that the method we used may have 496 less power to detect small effects than methods which make assumptions of balance in group 497 sizes.

498 Three groups of piglets (cohorts neomycin, neomycin+D-Scour[™] and 499 neomycin+ColiGuard®) underwent five days of treatment with neomycin, via intramuscular 500 administration. Intramuscular neomycin poorly diffuses (<10%) into a healthy gastrointestinal 501 tract ⁹¹, therefore a direct effect of neomycin on the gut microbiome may not be expected.

502 However, neomycin showed a different trend in unrooted PD between the second and the third 503 week post-weaning, corresponding to the week following the neomycin treatment period for the 504 neomycin cohort. Taking this time frame into consideration, the neomycin cohort did not 505 increase in BWPD to the extent of the Control cohort. Although statistically significant 506 differences between neomycin and Control in alpha diversity were not reached, possibly due to the effect being too small to detect within a relatively small sample size ^{92–94}, BWPD of the 507 508 neomycin cohort appears to follow a different trend to the Control cohort from the first week 509 (during neomycin treatment) where neomycin treated piglets show the lowest decrease of BWPD 510 compared to the control cohort and all other cohorts. While all cohorts show an increase in 511 absolute phylogenetic diversity accompanied by a decrease of diversity evenness during this time 512 frame, the neomycin cohort piglets show a lower drop in BWPD, suggesting an increase of 513 species richness, without a corresponding loss of species evenness. Furthermore the neomycin 514 cohort significantly separated from the control cohort in beta diversity two weeks after neomycin 515 treatment, showing a higher representation of Mollicutes. Numerous studies report the link of oral antibiotic use with dysbiosis ^{10,14–17,86,95}, as well as with host physiology changes ¹⁵. On the 516 517 contrary, the effect of intramuscular antibiotic administration on the microbiome is less well 518 investigated. Correlation between intramuscular antibiotic use and dysbiosis has been reported in fish ⁹⁶, gorillas ¹⁹, and pigs ^{21,97}. In one day old piglets, a single IM injection of amoxicillin 519 520 (penicillin class) is reported to have an effect on the intestinal microbiota, detectable 40 days post treatment ²¹. Zeineldin et al (2018) tested the effects of IM administration of several 521 522 antibiotics of various classes (penicillin, macrolide, cephalosporin and tetracycline), in 8-week 523 old piglets, reporting shifts of the Firmicutes/Bacteroidetes ratio following treatment (length of 524 the treatment not reported) 97. The effects of intramuscular administration of neomycin 525 (aminoglycoside class) on the gut microbiota have to our knowledge not been investigated. 526 Based on our results we conclude that a mild effect on phylogenetic diversity is appreciable post 527 IM neomycin treatment, up to two weeks after termination of the treatment. Additional 528 compositional and functional analysis is necessary to determine the source of this mild variation. 529 Differences were not detected at later time points, based on our phylogenetic diversity analyses, 530 suggesting a full recovery of the microbial communities after two weeks from the end of the 531 treatment.

532 It is possible that the large shifts in phylogenetic diversity taking place in the first two weeks 533 irrespective of the treatment (an increase, then decrease of unrooted PD, and an opposite trend of 534 BWPD) have masked the milder effects of the treatment, despite our efforts to control for the 535 effects of aging. This could be the reason why a significantly distinct alpha diversity trend was 536 found in the neomycin+D-Scour[™] cohort compared to the neomycin cohort, but not in the D-537 Scour[™] cohort compared to the Control cohort. There are multiple studies reporting beneficial 538 effects of probiotic treatment in sucker and weaner piglets in terms of improved gut mucosal integrity^{27,98}, growth rate⁹⁸⁻¹⁰⁰, digestibility of proteins and water absorption ^{98,101}, reduction of 539 pathogen invasion efficiency ^{38,45,98}, and decreased mortality ^{98,100}. Although the assessment of 540 541 physiologic changes from probiotic treatments was outside the scope of this study, we found 542 significant separation of neomycin+D-Scour[™] cohort samples to neomycin cohort samples in 543 beta diversity 2 and 12 days after D-Scour[™] treatment, where neomycin+D-Scour[™] samples 544 showed a higher representation of Lactobacillales compared to neomycin samples, suggesting a 545 transient establishment of the probiotic strains in the piglet guts.

The second probiotic in this study, ColiGuard®, did not have an effect on alpha diversity, but clustering was detected in beta diversity, where ColiGuard® samples separated from Control cohort samples in the first week of probiotic treatment and two weeks after probiotic treatment. Additionally, the ColiGuard® treatment correlated with a lower weight gain, whether or not it was preceded by the antibiotic treatment. However, when comparing the overall weight gain (from the start to the end of the trial, corresponding to the six weeks after weaning) the weight gain in the cohorts receiving ColiGuard® did not differ from the other cohorts.

We extracted the 16S rRNA gene hypervariable regions from our dataset, obtained the counts, and ran a correlation analysis to discover lineages that correlated with the weight of the piglets. As a consequence of the library size normalization step, the use of correlation with compositional data can inflate the false discovery rate ^{102,103}. For this reason it can be expected that some of the lineages we found to correlate with the weight of the piglets (eighty-three distinct species) could be spurious while other correlations may have been missed.

As our initial analysis of phylogenetic diversity was obtained using a 100K reads downsampling size per sample ¹⁰⁴, we wanted to test to what extent the use of a larger sampling 561 size would affect the diversity obtained from these samples. A 100K downsampling size 562 corresponds to a 0.6% of reads of an average sample. Increasing the sampling size from 0.6% to 563 6%, affected BWPD only slightly, while it increased the unrooted PD and decreased its variance. 564 As expected, a larger sampling size enriches the absolute diversity by increasing the chance to 565 make new read placements, and it lessens the variance in estimates, but it does not affect the core 566 microbiome. However, the small difference between sampling sizes could 1. demonstrate how 567 powerful the method is in describing diversity using a small sampling size, or 2. reflect the limits 568 imposed by the phylogenetic markers database, as it was pointed out by Darling et al (2014)⁵⁷.

569

570 Conclusions

571

572 Our findings stress the importance of confounding factors such as breed, age and maternal 573 effects when assessing the effect of treatment on the gut microbiome. We found that age, even 574 within a narrow age span (1-5 days) can have an impact on microbial shifts and should be 575 accounted for in microbiome studies, either (i). by accounting for it as a confounding variable in 576 the hypothesis-testing model used, or (ii). by avoiding, where possible, the inclusion of subjects 577 of different ages, or (iii). by allowing a sufficiently long period of time prior to the start of the 578 treatment. The latter allows animals to become accustomed to the new environment and 579 researchers to perform additional sampling. Allowing animals a sufficiently long period of time 580 to become accustomed to the new environment (e.g.: temperature, humidity, new microbes, etc) 581 is meant to reduce noise derived from external factors, while repeated sampling, likewise, aims 582 to increase the confidence in the signal prior to treatment.

Intramuscular neomycin treatment correlated with a clustering in alpha diversity and a higher representation of *Mollicutes* compared to control. D-ScourTM treated piglets displayed a transient establishment of *Lactobacillales*. ColiGuard® treated piglets displayed a clustering in beta diversity and a transient lower weight gain compared to control. Weight correlated with the abundance of a number of lineages. Age was the strongest factor shaping phylogenetic diversity of the piglets.

- 589 As previously mentioned, phylogenetic diversity is based on distinct lineages (richness) and
- 590 their collective structure (proportions reflected by BWPD) and not on a direct assessment of
- 591 composition and function. These types of analyses will be necessary to further describe the
- 592 effects of the treatments.
- 593

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- 601

602 **Conflicts of interest**

- 603 D-Scour[™] was sourced from International Animal Health Products (IAHP). ColiGuard[®] was
- 604 developed in a research project with NSW DPI, IAHP and AusIndustry Commonwealth
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606

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- 613 publication.

614

615 **Captions:**

616

617 **Figure 1**. Workflow.

618 Schematic workflow from sample collection to sequencing (orange) and data analysis (green).

619 Scripts (italic) are available in our GitHub repository.

620

621 Figure 2. Significant correlations of phylogenetic diversity with specified host factors.

622 The correlation of host factors birth day, cross-breed, birth day within the $D \times L$ cross-breed, birth 623 day within the D × LW cross-breed, line, litter (maternal sow and nurse sow) with alpha 624 phylogenetic diversity (\mathbf{A}) and beta phylogenetic diversity (\mathbf{B}) from each time point (x-axes) was 625 obtained. Shown in A and B are the significance values of these correlations derived from Kruskal-626 Wallis analysis, where the symbol shape indicates the phylogenetic diversity measure of alpha 627 (unrooted pd and BWPD) and beta (PC components 1 to 5), while the symbol color indicates the 628 significance before (black) and after (red) Hommel p value adjustment. C) For each red symbol in 629 A and B, describing a significant correlation after Hommel p value adjustment, a plot of the 630 phylogenetic diversity estimates is shown. Phylogenetic diversity estimates of the litter effect are 631 not shown. All p values are reported in Supplementary Table 1. Abbreviations: $D \times L = "Duroc \times$ 632 Landrace" cross breed; $D \times LW =$ "Duroc \times Large White" cross breed.

633

634 **Figure 3**. Effect of time on beta diversity.

635 Principal component analysis (PCA) of samples. PCA from edge component analysis with 636 PhyloSift (top) and PCA from 20 most abundant 16S reads extracted with SortMeRNA (bottom). 637 In the top, distribution of samples on either side of the plot (left versus right; top versus bottom) 638 reflect the lineages that were found to explain the variation. Samples are color coded by time post-639 weaning (days). In the lower plot, arrows indicate which of the 20 lineages contributed to the 640 variation of samples across time, where arrows thickness represents a higher (thicker) or lower 641 (thinner) contribution. Samples are color coded by time point during the trial (t0: start of the 642 trial/first day after weaning; t10: last day of the trial/40th day after weaning).

643

644 **Figure 4**. Lineages displaying the highest variation in beta diversity across time.

Heatmap of lineages explaining the community composition of samples from separate time points
of the trial (1 week interval between time points) derived from edge principal component analysis.
Intensity is derived from branch width by the percentage of variability explained by the principal
components.

649

650 **Supplementary Figure 1**. Batch effect on alpha and beta diversity before batch effect removal.

Batch effect by alpha (top two plots) and beta diversity (bottom five plots) before batch effect removal. Samples are grouped by DNA extraction plate. The *p* values are derived from multiple comparison analysis with ANOVA, indicating equality of the means. *Post hoc* corrected *p* values for pairwise comparisons are provided in Supplementary Table 1.

655

656 **Supplementary Figure 2**. Alpha phylogenetic diversity of cohorts.

657 Alpha phylogenetic diversity per cohort from samples across all time points. Balance-weighted 658 phylogenetic diversity (BWPD) (top) (mean±SD: Positive control Mock community: 1.59±0.07; 659 Positive control D-Scour[™]: 1.94±0.21; Positive control ColiGuard[®]: 0.86±0.14; Control: 660 2.13±0.13; D-Scour[™]: 2.16±0.12; ColiGuard[®]: 2.12±0.12; neomycin: 2.12±0.16; neomycin+D-661 Scour[™]: 2.13±0.13; neomycin+ColiGuard[®]: 2.10±0.14; sows: 2.12±0.15; all piglet cohorts: 662 2.13±0.13); Unrooted phylogenetic diversity (bottom) (mean±SD: Positive control Mock 663 community: 123.62±24.41; Positive control D-Scour[™]: 162.14±28.27; Positive control 664 ColiGuard®: 129.88±50.00; Control: 311.23±29.23; D-Scour™: 316.35±24.99; ColiGuard®: 665 311.98±42.51; neomycin: 314.81±44.01; neomycin+D-Scour[™]: 312.66±40.18; 666 neomycin+ColiGuard®: 316.29±31.20; sows: 328.51±24.00; all piglet cohorts: 313.86±36.00).

667

668 **Supplementary Figure 3**. Beta diversity of positive controls.

669 Principal component analysis (PCA) of positive control samples. PCA from edge component 670 analysis with PhyloSift. Distribution of samples on either side of the plot (left *versus* right; top 671 *versus* bottom) reflect the lineages that were found to explain the variation.

672

673 **Supplementary Figure 4**. Alpha phylogenetic diversity by age breed.

Alpha diversity of samples from the start of the trial (immediately after weaning) grouped by breed and by date of birth. Unrooted phylogenetic diversity (top) and balance-weighted phylogenetic diversity (bottom). P values are derived from Kruskal-Wallis analysis of variance. Piglets of the Duroc × Landrace breed (n=46) separated significantly by age in unrooted phylogenetic diversity and in BWPD at the start of the trial (t0) (Hommel adjusted p value: unrooted pd=0.006; BWPD=0.047). All post hoc corrected p values are provided in Supplementary Table 1.

680

681 **Supplementary Figure 5**. Relative abundance heatmap.

Most abundant lineages within each time point (columns) are obtained from analysis with guppy fat. Guppy fat outputs trees with fattened edges in proportion to the relative abundance of reads place in each lineage. The branch width of the trees, each corresponding to samples from distinct time points, are the entries for this heat map. The distance between each time point is one week.

686

687 **Supplementary Figure 6**. Time trend of alpha diversity by cohort.

688 Unrooted phylogenetic diversity (**A**) and balance-weighted phylogenetic diversity (**B**) describe 689 richness and evenness, respectively, of alpha phylogenetic diversity for all samples across time, 690 grouped and color coded by cohort. The *p* values derive from pairwise comparisons of time points 691 of all treatment cohorts. The *p* values and *post hoc* corrected *p* values of time points comparisons 692 for each separate treatment cohort are provided in Supplementary Table 1.

693

694 Supplementary Figure 7. Significant differences in beta diversity between cohorts at

695 specific time points.

696 Significance was determined by comparing groups by pairwise t-test and the resulting *p* values 697 were adjusted with the Bonferroni method. Significance values are provided in Supplementary 698 Table 1. The x-axes represent the principal component. As plots are derived from distinct guppy 699 runs, each principal component explains variation to a different extent (percentage specified in 700 parentheses). The number of samples is specified on the y-axis. Distribution of the samples on 691 either side of a plot (left *versus* right) reflects the lineages that were found to explain the variation. 702 Distributions are color coded by cohort.

703

704 Supplementary Figure 8. Change in weight gain of piglets by cohort across the trial.

On the y-axis of plots the change in weight gain between time points is provided in percentage. Letters on the top left of each plot indicate the time points compared with one week interval (t0t2, t2-t4, t6-t8, t8-t10) and with two weeks interval (t2-t6, t4-t8, t0-t8). Pairwise t-test comparisons between cohorts were computed. A significant difference was found between Control and ColiGuard (t4-t6, Tukey adjusted p value=0.0084), between neomycin and neomycin+ColiGuard (t4-t6, Tukey adjusted p value=0.0152) and between neomycin and neomycin+ColiGuard (t4-t8,

711 Tukey adjusted *p* value=0.011).

712

Supplementary Figure 9. Correlation of diversity indices between the analysis run using 100,000
(100K) and 1,000,000 (1M) reads per sample.

Phylogenetic diversity analysis (alpha and beta) was run using either 100K or 1M reads per sample, corresponding to a 0.6% or 6% of the average sample. We show the Pearson's correlation of these analyses run using distinct downsampling sizes. Significance of correlations is reported within each plot. All the diversity indices are provided in Supplementary Table 1.

719

720 Supplementary Table 1. Statistical analysis output.

721 Statistical analysis output is split across the following sheets. **all_padj_Hommel** : list of *p* values

722 obtained from running Kruskal-Wallis test, adjusted using the Hommel correction for

phylogenetic diversity values: unrooted pd, BWPD, and five principal components. Grouping of

724 samples is based on: cross-breed, line, date of birth, date of birth within distinct cross-breeds, 725 maternal sow and nurse sow. Sample size and sample collection date is reported. all pvalues : 726 reports the *p* values of "all_padj_Hommel" sheet, prior to Hommel *p* value adjustment. 727 alpha delta cohorts : deltas between time points are obtained per cohort and these deltas are 728 compared using ANOVA, adjusting p values using the TukeyHSD method. alpha_means : the 729 means and standard deviations of alpha diversity (unrooted pd and BWPD) obtained per time 730 point and cohort. **alpha time** : results of comparison between time points of alpha diversity 731 (unrooted pd and BWPD) using the t-test and Bonferroni p value correction. 732 **batch post process**: results of comparison of alpha diversity values by DNA extraction plate 733 (1-10) after batch effect removal, run using ANOVA and TukeyHSD p value adjustment method. 734 batch pre process : results of comparison of alpha diversity values by DNA extraction plate (1-735 10) before batch effect removal, run using ANOVA and TukeyHSD p value adjustment method. 736 **deltas_percent_change** : comparison of alpha diversity values (unrooted pd and BWPD) 737 between time points within distinct cohorts and within all cohorts. Sample size (n), deltas 738 (percentage) are shown. **guppy** padj : p values from "guppy pvalues" are adjusted using the 739 Bonferroni method and results are reported. guppy_pvalues : output of single guppy runs is 740 analyzed by comparing beta diversity values by cohort at each time point using the pairwise t 741 test; results of the tests are here shown. PD_100K_vs_1M_stats : alpha and beta diversity values 742 obtained with PhyloSift using either 100,000 reads or 1,000,000 reads downsampling size. 743 weight cohort stats : piglets were weighted at each time point. Deltas are obtained per piglet 744 and means were compared between cohorts using ANOVA and TukeyHSD p value correction, 745 results are reported. weight taxa: Spearman's rank correlation was assessed between pig weight 746 and abundance of lineages obtained from 16S rRNA containing reads. Significance values are 747 adjusted using the distinct methods listed. 748

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- 752

753 **References** 754

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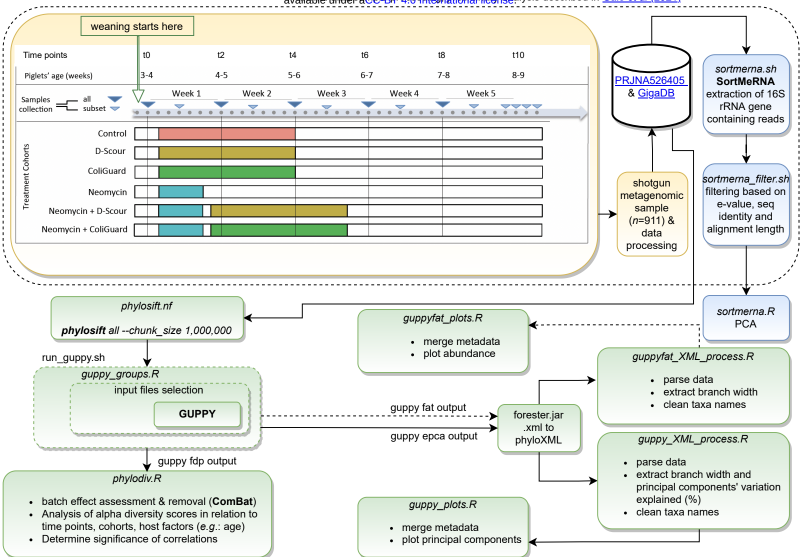
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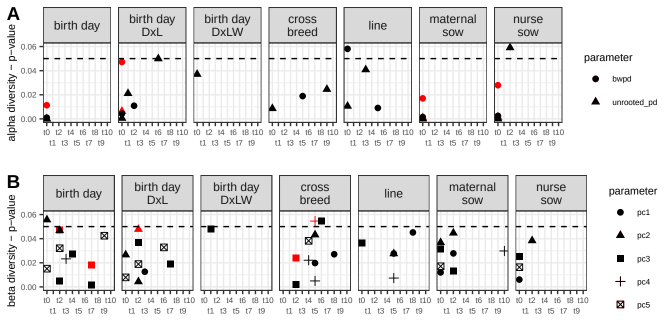
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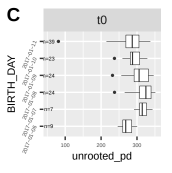
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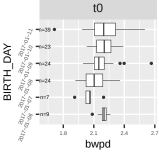
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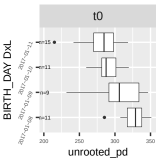
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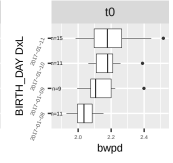


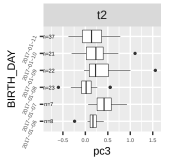


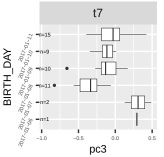


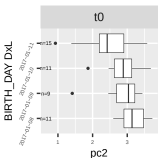


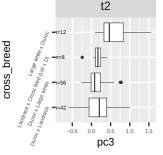




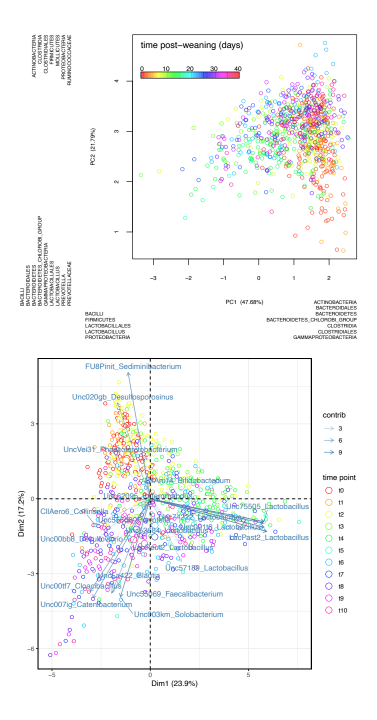








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Г	0.11	0.27	0.29	0.34	0.31	0.37	LA
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ЦЦ	0.07	0.00	0.00		0.00	0.00	BA
	0.05	0.00	0.00		0.00	0.00	ΕN
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	0.07	0.05	0.00		0.00	0.00	ME
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Ч Г	0.04	0.00	0.11	0.06	0.07	0.03	PR
1	0.09	0.02	0.13	0.07	0.08	0.04	PR
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ACTOBACILLALES	0.35
ACTOBACILLUS	0.3
BACTEROIDALES	0.25
BIFIDOBACTERIACEAE	0.2
BIFIDOBACTERIUM	0.15
RUMINOCOCCUS	0.1
PROPIONIBACTERIUM	0.05
SPIROCHAETALES	0
_ACTOBACILLUS_JOHNSONII	
CLOSTRIDIACEAE	
CLOSTRIDIUM	
ERYSIPELOTRICHACEAE	

ERYSIPELOTRICHACEAE FAECALIBACTERIUM_PRAUSNITZII CORIOBACTERIACEAE BACTEROIDES ENTEROBACTERIACEAE METHANOBREVIBACTER_SMITHII METHANOBREVIBACTER METHANOBACTERIACEAE METHANOBACTERIALES LACTOBACILLACEAE PREVOTELLA