Community composition and development of the post-weaning piglet gut microbiome

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Research

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Abstract

Background

Early weaning and intensive farming practices predispose piglets to the development of infectious and often lethal diseases, against which antibiotics are used. Besides contributing to the build-up of antimicrobial resistance, antibiotics are known to modulate the gut microbial composition. Studies have previously investigated the effects of probiotics as alternatives to antibiotic treatment for the prevention of post-weaning diarrhea. In order to describe the post-weaning gut microbiota, and the effects of two probiotics formulations and of intramuscular antibiotic treatment on the gut microbiota, we processed over 800 faecal time-series samples from 126 piglets and 42 sows, generating over 8Tbp of metagenomic shotgun sequence data. Here we describe the animal trial procedures, the generation of our metagenomic dataset and the analysis of the microbial community composition using a phylogenetic framework.

Results

Factors such as age, litter effects, and breed, by significantly correlating with gut microbial community shifts, can be major confounding factors in the assessment of treatment effects. Intramuscular antibiotic treatment and probiotic treatments were found to correlate with alpha and beta diversity, as well as with a transient establishment of *Mollicutes* and *Lactobacillales*, respectively. We found the abundance of certain taxa to correlate with weight gain.

Conclusions

Our findings demonstrate that breed, litter, and age, are important contributors to variation in the community composition, and that treatment effects of the antibiotic and probiotic treatments were subtle, while host age was the dominant factor in shaping the gut microbiota of piglets after weaning. The current study shows, by means of a phylogenetic diversity framework, that the post-weaning pig gut microbiome appears to follow a highly structured developmental program with characteristic post-weaning changes that can distinguish hosts that were born as little as two days apart in the second month of life.

Background

As the world population grows, there is an accompanying demand for animal derived products. In a semi-natural environment pig weaning occurs between the 12th and the 17th weeks from birth, whereas in the farm this typically occurs at 4 weeks of age\(^1\). Intensive animal husbandry and early weaning practices are commonly used to maximise production rates while minimizing costs. Both practices increase the risk of infections with pathogenic organisms, and thereby the need for antimicrobial strategies, which has included the common use of antibiotics.\(^2\)\(^−\)\(^12\) Although antibiotics may kill some pathogens, the surviving bacteria can develop antimicrobial resistance (AMR) against the class of antibiotic used\(^13\)\(^−\)\(^20\), as well as
against other antibiotic classes. A retrospective U.S. livestock study found evidence of multidrug resistance (resistance to >3 antimicrobial drug classes) in *Escherichia coli* that increased from 7.2% to 63.6% between the 1950s and the 2000s. As the incidence of multidrug resistant (MDR) pathogens expands, the World Health Organization (WHO) has recognized AMR as one of the top health challenges of the current century.

Besides leading to AMR development, antibiotic usage is known to cause dysbiosis, the disruption of a balanced state within a gut microbial community. A balanced state decreases the chance of pathogens gaining a foothold, as there is niche and nutrient competition in play. In livestock production, intramuscular (IM) antibiotic treatment is preferred over oral antibiotic treatment, as it is thought to contribute less to AMR than oral antibiotic treatment. Extensive evidence exists on dysbiosis as a consequence of oral antibiotic treatment on the gut microbiome, whereas only a handful of studies report the effects of intramuscular (IM) antibiotic treatment on the gut microbiota.

A number of non-antibiotic strategies to increase resistance to disease have been studied. A number of strategies for which beneficial effects have been reported consist of hydrolases, fiber intake, β-lactamase enzymes, non-resistance inducing antimicrobial drugs, vaccination, phage therapy, in-feed organic acids, starches, and liquid feed. Extensive evidence exists on the use of probiotics. Probiotics are reported to promote intestinal health in multiple ways: improvement of mucosal integrity, competitive exclusion against pathogenic species, reduction of intestinal inflammation and pathogen translocation. Their efficacy has been determined, among others, in cell cultures, mice, and swine. However, while most evidence of beneficial effects from probiotic treatment is derived from host immunity responses, host physiology, and pathogen relocation, fewer studies exist on the effects of probiotics on the whole gut microbial community.

This study was conducted to: 1) generate a public metagenomic databank of the gut microbiome of weaner piglets aged between 3 and 9 weeks old; 2) assess the effects of IM neomycin antibiotic use on the gut community; and 3) assess the effects of two probiotic formulations on the gut community. The data we present here is analysed from the perspective of the phylogenetic diversity of the microbial communities.

In the first results section we describe the phylogenetic diversity of the intestinal microbiome of the piglets and how it compares to the known composition of positive controls. Then, we describe evidence that highlights the importance of age, breed and litter as strongly associated factors with changes in community composition. Lastly, we describe the strong effect of time on community composition and the milder effect of antibiotic and probiotic treatments on community composition.
PhyloSift 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 at the start of the trial. To this end, analysis of the unrooted phylogenetic diversity (PD), the balance weighted phylogenetic diversity (BWPD) and principal component analysis (PCA) of the Kantorovitch-Rubinstein distances (beta diversity analysis) were performed.

**Comparison of the expected and the observed taxonomic profile of the positive controls**

All the mock community members, in seven of the eight technical replicates, were detected by MetaPhlAn2 (S8). One sample failed to sequence, reporting zero counts for any species. The observed mean relative abundances were as follows: *B. subtilis* (mean ± SD: 2.92 ± 0.994), *E. cloacae* (mean ± SD: 38.0 ± 6.404), *E. faecium* (mean ± SD: 0.97 ± 0.081), *E. coli* (mean ± SD: 10.12 ± 1.480), *E. coli* unclassified (mean ± SD: 7.83 ± 1.755), *P. aeruginosa* (mean ± SD: 26.72 ± 3.026), *S. aureus* (mean ± SD: 9.90 ± 3.613), *S. epidermidis* (mean ± SD: 3.54 ± 1.435). Isolate *E. cloacae* C15117, used in this study for the make-up of the mock community, was recently found to be most closely related to the *Enterobacter hormaechei* phylogenomic group C type strain DSM 16687 and therefore re-identified as *Enterobacter hormaechei subsp. oharae*90. For this reason, taxonomic assignment by MetaPhlAn2 attributed the reads to *E. cloacae* instead. The expected proportions of the mock community members were derived from the CFU by the genome size. Based on the expected and the observed relative abundance, we found, with the exception of *S. aureus* (exp: 8.7% obs: 9.9%), three Gram positive members to be under-represented (*B. subtilis*: exp: 13.0% obs: 2.9%; *E. faecium*: exp: 7.7% obs: 1.0%; *S. epidermidis*: exp: 16.7% obs: 3.5%) and, with the exception of *P. aeruginosa* (exp: 38.9% obs: 26.8%), two Gram negative members to be over-represented (*E. cloacae*: exp: 14.5% obs: 38.0%; *E. coli*: exp: 0.4% obs: 7.8–10.1%) (S9). Taxonomic assignment of the mock community samples reported one contaminating species in one of the eight replicates: *Lactobacillus salivarius* (mean: 0.008) (S10).

The probiotic D-Scour™ is expected to contain, per gram, a total of 180 million CFU of *Lactobacillus acidophilus*, *Lactobacillus delbrueckii* subspecies *bulgaricus*, *Lactobacillus plantarum*, *Lactobacillus rhamnosus*, *Bifidobacterium bifidum*, *Enterococcus faecium*, *Streptococcus salivarius* subspecies *thermophilus* in unknown proportions. From taxonomic analysis with MetaPhlAn2, we can conclude that 6 of the 7 expected species were determined to be present in the replicates in the following mean relative abundances: *Bifidobacterium bifidum*: mean ± SD: 40.01 ± 12.558; *Enterococcus faecium*: mean ± SD: 30.98 ± 13.472; *Lactobacillus delbrueckii*: mean ± SD: 11.56 ± 7.148; *Lactobacillus plantarum*: mean ± SD: 6.23 ± 7.863; *Lactobacillus rhamnosus*: mean ± SD: 2.08 ± 1.226; *Streptococcus thermophilus*: mean ± SD: 4.28 ± 1.523. *Lactobacillus acidophilus* was not detected and *Lactobacillus helveticus* was detected instead (*Lactobacillus helveticus*: mean ± SD: 4.75 ± 2.431) (S8). An additional 25 taxa were detected, of which 18 and 7 were identified at the species and at the genus level, respectively. Contaminants were present at a higher concentration in three technical replicates (R3, R7, R8) with the most frequent contaminant (*Methanobrevibacter* spp.) being present in 5 of the 8 replicates (S11).
Taxonomic analysis of the technical replicates of the probiotic ColiGuard® also showed a species profile consistent with the expected profile, with *Lactobacillus salivarius* and *Lactobacillus plantarum* in a 9:1 ratio (*Lactobacillus salivarius*: mean ± SD: 93.52 ± 1.617; *Lactobacillus plantarum*: mean ± SD: 6.10 ± 1.134) across the replicates (S8). ColiGuard® contained a total of 20 contaminants, of which 16 and 4 were identified at the species and the genus level, respectively. Contaminants were present at a higher level in two technical replicates (R5, R7), with R7 displaying the most diverse and highest contamination rate (R7: 14 taxa; total contaminating reads: 2.67%; R5: 9 taxa; total contaminating reads: 0.30%). (S12).

**Phylogenetic diversity of positive controls and how it compares to the taxonomic profile**

Alpha diversity of the positive controls reflects the expected alpha diversity, with the mock community, ColiGuard® and D-Scour™ positive controls, displaying a progressively higher unrooted PD (mean ± SD: Mock community: 31.53 ± 29.50; ColiGuard®: 58.52 ± 21.70; D-Scour™: 64.84 ± 21.30). BWPD for the positive control ColiGuard® appears on the far left of the plot (mean ± SD: 0.82 ± 0.15), distant from the mock community (mean ± SD: 1.58 ± 0.12) and the positive control D-Scour™ (mean ± SD: 1.89 ± 0.26), which contains 2, 7 and 8 main species, respectively (S13; Supplementary file 1). The unrooted PD of the positive control ColiGuard® (mean ± SD: 58.52 ± 21.70) is closer to the unrooted PD of the Mock Community (mean ± SD: 31.53 ± 29.50) and to that of the positive control D-Scour™ (mean ± SD: 64.84 ± 21.30), than the BWPD.

The contribution to total diversity of phylogenetic tree edges with uneven quantities of reads placed on either side is down-weighted in BWPD, which is reflected in the low reported BWPD of the positive control ColiGuard®. ColiGuard® is mainly composed of two species in an uneven ratio (1:9) as by taxonomic analysis with MetaPhlAn2 (S8), in contrast with the mock community (0.01:1:3:4:8:10:10:27:38) and the positive control D-Scour™ (2.08:4.28:4.75:6.23:11.56:30.98:40.01) (S8). On the other hand, the unrooted PD reflects the absolute diversity, independently of the 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-Scour™: 25 taxa) (S11-13) contributes toward the absolute diversity in the unrooted PD, erroneously inflating this value (mean ± SD: Mock Community: 31.53 ± 29.50; ColiGuard®: 58.52 ± 21.70; D-Scour™: 64.84 ± 21.30) (Fig. 2).

**Phylogenetic diversity of piglet gut communities immediately after weaning**

The 126 piglets used in this study were derived from 4 main breed cross types “Duroc × Landrace” (n = 46), “Duroc × Large White” (n = 59), “Landrace × cross bred (LWxD)” (n = 9), “Large White × Duroc” (n = 12), and three pig lines (line 319: n = 9; line 316: n = 46; line 326: n = 71). The piglets also differed slightly by age, being born between 1 and 6 days apart.

Based on Kruskal-Wallis one-way analysis of variance (Hommel adjusted \( p \) values to correct for multiple testing), phylogenetic diversity of the piglet samples did not cluster significantly by breed cross type (\( p > \)
0.05) or by pig line ($p > 0.05$) at the start of the trial, but a significant difference was found with breed and line in the fourth week of the trial (breed: $p = 0.04$; pig line: $p = 0.02$) (Fig. 2). Additionally a correlation of breed cross type was found with beta diversity principal components (PC) in the second week of the trial (PC3: $p = 0.0357$) and in the fifth week of the trial (PC1: $p = 0.027$). Alpha diversity significantly correlated with pig line in the fourth week of the trial (BWPD: $p = 0.017$). (Fig. 2; Supplementary file 5)

Notably, we found a significant correlation between alpha diversity and the age of the piglets at the first sampling time point (unrooted PD: $p = 0.0016$; BWPD: $p = 0.0355$) (Fig. 2; Supplementary file 5). While a correlation with age was found for alpha diversity only at the start of the trial, age of the piglet was observed to be weakly associated with differences in community composition in week two (PC3: $p = 0.0507$) (S14; Supplementary file 5).

As age groups were confounded with cross-breed types (*i.e.* not all age groups are represented by each of the four cross-breed types), we compared the phylogenetic diversity of age groups within each breed. As cross-breed types “Landrace × cross bred (LW × D)” and “Large White × Duroc” had only a small number of piglets in each age group, we tested for an association between phylogenetic diversity and age in cross-breeds “Duroc × Landrace” and “Duroc × Large White”. In these cross-breeds, alpha and beta diversity are correlated with age in the “Duroc × Landrace” piglets (unrooted PD: $p = 0.0059$; BWPD: $p = 0.0226$; PC2: $p = 0.0263$; PC5: $p = 0.0063$) and to a lesser extent in the “Duroc × Large White” piglets (unrooted PD: $p = 0.0529$; PC5: $p = 0.0310$) during the first week of the trial. In the “Duroc × Landrace” piglets, a correlation between age and beta diversity was detected in week two (PC2: $p = 0.0347$) (Fig. 2; S14; Supplementary file 5). Differences in beta diversity by date of birth, within the same cross-breed (Duroc × Large white), were found to be significant (TukeyHSD adjusted $p$ value: 1 day difference: 0.000782; 2 days difference: 0.018603) (Supplementary file 5).

**Maternal effect on phylogenetic diversity**

As piglets were derived from 42 distinct sows (maternal sows), and nursed by either the same or a different sow (a nurse sow) (Supplementary File 5), a litter effect was expected and determined. Based on Hommel adjusted $p$ values, a similarity of alpha and beta phylogenetic diversity can be seen among piglets from the same maternal sow (unrooted PD: $p = 0.0096$; BWPD: $p = 0.0467$) and in piglets from the same nurse sow (unrooted PD: $p = 0.0320$; PC1: $p = 0.0391$) (Fig. 2; S16-19; Supplementary file 5). The litter effect described was found in samples at the start of the trial. Significance of the correlations did not persist thereafter (Fig. 2; Supplementary file 5).

**A strong effect of aging on phylogenetic diversity**

Beta diversity analysis revealed a distinct and consistent change of the microbial community over time in all piglets, regardless of the treatment. Samples clustered in PC2 (accounting for 21.68% of the variation), showing a higher representation of *Bacteroidetes, Gammaproteobacteria* and *Prevotellaceae* from day 0–20, and a higher representation of *Firmicutes, Mollicutes* and *Ruminococcaceae* from day 20–40 (Fig. 3; S20). In PC1 (accounting for 46.99% of the variation) samples shifted towards a higher representation of *Lactobacilli* from day 0 to day 20, and towards a higher representation of *Actinobacteria, Clostridiales*
and Mollicutes from day 20 to day 40 (Fig. 3; Supplementary file 5). A temporary shift towards a higher representation of Bifidobacteriaceae (Bifidobacterium being a component of the probiotic D-Scour™) is seen in PC5 (3.04% of the variation explained) one and two weeks from the start of the trial (S21).

Beta diversity analysis was performed separately for samples within each time point in order to find taxa 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 (S22). The following taxa were responsible for variation only at the start of the trial: Gammaproteobacteria (t0 = 0.12), Enterobacteriaceae (t0 = 0.08) and Archaea (t0 = 0.03). The following taxa were responsible for variation throughout the trial: Clostridiales (min = 0.11; max = 0.25), Bacteroidetes (min = 0.06; max = 0.11), Bacteroidetes chlorobi group (min = 0.05; max = 0.10). Lactobacillus became variable after the first week of the trial and remained highly variable throughout the trial (min = 0.13; max = 0.22). Some taxa contributed less to the variability among the piglets’ faecal population, but they nonetheless consistently contributed to variability: Bacteroidales (min = 0.01; max = 0.04); Firmicutes (min = 0.05; max = 0.08); Bacilli (min = 0.06; max = 0.09); Lactobacillales (min = 0.04; max = 0.08); Actinobacteria (min = 0.02; max = 0.09); Proteobacteria (min = 0.02; max = 0.07). The variability of Prevotella (min = 0.01; max = 0.04) increased from the second week of the trial. (S22)

Taxonomic representation in terms of abundance was derived from analysis of samples with guppy fat, where abundance was derived from the branch width (Fig. 4) and combined with PhyloSift’s taxonomic annotation of the marker gene phylogeny. Overall, Clostridiales were the most abundant throughout the trial (t0 = 0.18; t2 = 0.20; t4 = 0.11; t6 = 0.09; t8 = 0.10; t10 = 0.08). Lactobacillus acidophilus increased at the start (t0 = 0.00; t2 = 0.05; t4 = 0.13) then decreased (t6 = 0.09; t8 = 0.05; t10 = 0.07). Subdoligranulum variabile decreased after the first week (t0 = 0.05; t2 = 0.02) then gradually increased (t4 = 0.04; t6 = 0.05; t8 = 0.06; t10 = 0.07). Among other prevalent taxa at the start of the trial up to the second week and dropping at later time points we found Metahobrevibacter smithii (t0 = 0.06; t2 = 0.03) and Bacteroidales (t0 = 0.02; t2 = 0.01). Following an opposite trend we found Ruminococcus sp. JC304 (t0 = 0.00; t2 = 0.00; t4 = 0.01; t6 = 0.02; t8 = 0.02; t10 = 0.03), Solobacterium moorei (t0 = 0.00; t2 = 0.01; t4 = 0.02; t6 = 0.02; t8 = 0.02; t10 = 0.02) and Prevotella copri (t0 = 0.00; t2 = 0.00; t4 = 0.04; t6 = 0.03; t8 = 0.05; t10 = 0.02). In modest and stable abundance across the trial were Mogibacterium sp. CM50 (t0 = 0.02; t2 = 0.03; t4 = 0.02; t6 = 0.03; t8 = 0.02; t10 = 0.04) and Oscillibacter (t0 = 0.06; t2 = 0.04; t4 = 0.03; t6 = 0.03; t8 = 0.03; t10 = 0.03). In transient abundance we found Bifidobacterium thermophilum RBL67 (t2 = 0.01; t4 = 0.02). Gradually increasing from the second week we found Eubacterium biforme DSM3989 (t2 = 0.01; t4 = 0.02; t6 = 0.02; t8 = 0.02; t10 = 0.02), Eubacterium rectale (t4 = 0.02; t6 = 0.02; t8 = 0.03; t10 = 0.01) and, after the third week, Faecalibacterium prausnitsii (t6 = 0.01; t8 = 0.02; t10 = 0.02).

The effect of time was also clear in alpha diversity, where all the piglet cohorts followed a similar trend over time, independent of the treatment (S23; Supplementary file 5). Upon arrival on the trial site, the piglets’ unrooted PD was lower than the sows (sows: 125.39; piglets range: 104.6-112.7) and reached a similar unrooted PD to the sows’ in the second week (range: 123.4-130.7) (Supplementary file 5).
In comparing four timepoints at one week intervals from the start of the trial (intervals labeled as A, B, C, D), changes in alpha diversity among all the piglets were tested for and significance was determined using the Bonferroni correction. Unrooted phylogenetic diversity increased after the start of the trial (A mean: 107.86; B mean: 127.21; $p < 0.001$), decreased in the following week (C mean: 122.92; $p = \text{ns}$) and the fourth week (D mean: 118.74; $p = \text{ns}$) amongst the piglets cohorts. In contrast, BWPD decreased after the start of the trial (A mean: 2.17; B mean: 2.05; $p < 0.0001$), to increase in the following week (C mean: 2.14; $p < 0.0001$) and decrease in the fourth week (D mean: 2.07; $p < 0.01$). (S23; Supplementary file 5).

The increase in unrooted PD in the first week was significant for all treatment cohorts ($p$ range: 0.0015-0.03) except ColiGuard® and neomycin ($p = \text{ns}$) cohorts. The decrease in BWPD in the first week was only significant for the control ($p = 0.04$), ColiGuard® ($p = 0.0015$) and neomycin + ColiGuard® ($p = 0.036$) cohorts. In week two only the increase of BWPD of the control cohort reached statistical significance ($p = 0.0121$). (S24; Supplementary file 5)

**Effect of antibiotic and probiotic treatment on alpha diversity**

We hypothesized that the probiotic treatments, whether alone (D-Scour™ and ColiGuard®) or administered after neomycin (neomycin + D-Scour™ and neomycin + ColiGuard®) would cause a change in the microbial community composition that would be measurable via phylogenetic diversity. We tested whether the treatments correlated with a change in phylogenetic diversity independently of the changes occurring with time. Given the differences in alpha and beta diversity detected among the subjects at the start of the trial, we analyzed the deltas of phylogenetic diversity instead of relying on the absolute means, similar to the procedure applied by Kembel *et al* (2012) 91. Time-point measurements of alpha diversity were taken and deltas were computed for each piglet. Delta means were compared between cohorts, where the control cohort would serve as a control group for neomycin, D-Scour™ and ColiGuard® cohorts, whereas the neomycin cohort would serve as a control group for the neomycin + D-Scour™ and neomycin + ColiGuard® cohorts.

After the first week of the trial, the majority of the piglets displayed an increase in unrooted PD (85%) and a decrease of BWPD (75%). The following week the trend was opposite: an increase of BWPD (78%) and a decrease of unrooted PD (60%) (Supplementary file 5). However, the neomycin cohort displayed the smallest BWPD drop in the week following the start of the trial and the overall trend of neomycin in unrooted PD was the most different from the other cohorts (S25). Due to the delayed rise in unrooted PD of neomycin, significance was found in unrooted PD between neomycin and neomycin + D-Scour™ during the first week of D-Scour™ treatment (neomycin mean: 1.59; neomycin + D-Scour™ mean: -11.79; $p = 0.012188$) (S25 B-C) and in unrooted PD between neomycin and neomycin + ColiGuard® (neomycin mean: -5.53; neomycin + ColiGuard®: -6.70; $p = 0.043138$) (S25 C-D). The change was opposite in BWPD ($p = 0.049263$) (S26 B-C) and persisted during the following week, up to 3 days post- D-Scour™ treatment (neomycin mean: -6.34; neomycin + D-Scour™ mean: 4.52; BH adjusted $p = 0.024092$) (S26 B-D).
The only significant difference between the neomycin and the control cohort was detected in BWPD in the second week of the trial (control mean: 7.10; neomycin mean: 0.28) \( p = 0.033751 \) (S26 B-C).

No significant differences were detected between the control and the D-Scour™ cohort \( (p<0.05) \) (Supplementary file 5). Instead, a significant decrease in BWPD was detected in the ColiGuard® cohort between the start of the trial and the end of the ColiGuard® treatment, where nearly all piglets (88%) in the ColiGuard® cohort displayed a decrease in BWPD, whereas the control cohort was split between piglets increasing (53%) and decreasing (47%) in BWPD (control mean: 0.85; ColiGuard® mean: -5.49; \( p = 0.040041 \) ) (S26 A-C).

Effects of antibiotic and probiotic treatment on beta diversity

To investigate the treatment effect on beta diversity, principal component analysis (PCA) of the Kantorovich-Rubenstein distances (beta diversity analysis) was performed on all samples and, additionally, on samples within individual time points. This analysis is conceptually similar to the weighted Unifrac approach for beta diversity analysis, but is designed to work with phylogenetic placement data. When examining all samples together, there was no clear separation of cohorts on any of the five principal component axes. When individual time points were analysed, some clustering by cohort was observed (S27). D-Scour™ separated from the control cohort in PC2 (17.95%) and in PC3 (8.06%) one week and two weeks after probiotic treatment, respectively (S27). ColiGuard® separated from control two weeks post probiotic treatment in PC3 (7.88%). Two weeks after the end of neomycin treatment, neomycin separated from control in PC4 (5.32%), with a smaller clustering and a higher representation of Mollicutes. Two and four weeks after probiotic treatment, neomycin + D-Scour™ separated from neomycin in PC2 (17.74%) and PC5 (3.81%), respectively. In both timepoints and principal components, neomycin + D-Scour™ showed a higher representation of Lactobacillales. (S27)

Association between weight and community composition

Weight correlated with the abundance of certain species \( (n=83) \) at each time point as it resulted from principal component analysis of 16S rRNA reads. Positively correlating with weight we found among others: Blautia (t0), Cetobacterium (t0), Ruminibacter (t0), Rickettsia (t2), Lactobacillus (t6), Mycoplasma (t6), Anaerostipes (t8), Ruminococcus (t8-t10), Cerasibacillus (t10). Negatively correlating with weight we found among others: Pyramidobacter (t0), Odoribacter (t2), Schwartzia (t6), Streptococcus (t6), Dokdonella (t8). (S28-S33) Significance values and correlation estimates are given in Supplementary file 5.

Effect of treatments on weight gain

Overall weight gain from initial to final weight (S34 A-E; Supplementary file 5) was not significantly affected by any treatment. However, the probiotic ColiGuard® was found to have a partial effect on piglet weight gain (S34; Supplementary file 5). Weight was measured weekly for a total of six measurements. Based on Tukey adjusted \( p \) values, a lower weight gain was detected in the ColiGuard® cohort in week 3.
compared to the control cohort ($p = 0.0008$) (S34 C-D; Supplementary file 5). Similarly, a lower weight gain was detected in the neomycin + ColiGuard® cohort in C-E compared to the neomycin cohort ($p = 0.0393$) (S34 C-E; Supplementary file 5). Breed and age differences among piglets were not associated with weight gain (Supplementary file 5).

**Discussion**

The consistent trend in community composition over time, across all the cohorts, indicates that an age-related process of ecological succession is the largest factor shaping the microbial community of post-weaning piglets, as found in this study where animals aged 20–63 days and were fed the same diet. A peak in unrooted phylogenetic diversity and drop in balance weighted phylogenetic diversity (BWPD) reflects the acquisition of new species with the loss of dominating species. This change, detected in the week following the piglets’ arrival at the trial site irrespective of the cohort, could be linked to the piglets being subjected to microbial interchange (e.g.: new pen mates \(^92,93\)) and/or to diet transition (peri-weaning transition to solid food \(^92,93\)) leading to the reshaping of the gut microbial community. The week following the drop of BWPD, a significant increase of BWPD was recorded, reflecting the acquisition of a larger proportion of the community by the newly introduced species. The strong changes in phylogenetic diversity detected in the first and the second week could as well be attributable to other post-weaning related physiological changes, as previous studies report \(^44,92−94\).

The highest inter-individual differences among piglets are seen in the first week of life, irrespective of maternal or environmental effects. The microbiota of 3 week old piglets is still very dynamic, but environmental factors become evident \(^93\). At 6 weeks of age, CD8 + T cells infiltrate the intestinal tissue and the mucosa and intestinal lining resemble that of an adult pig \(^115\). In this study, piglets reached a comparable alpha diversity to the sows after the first week of the trial, at which time the piglets were aged between 3.8 and 4.6 weeks. Unrooted PD did not reach higher levels at later sampling time points. The highest BWPD accompanied by a high unrooted PD was reached after the second week of the trial when piglets were aged between 4.9 and 5.6 weeks. Age-dependent physiological changes could explain i) the major shifts we detected in alpha diversity during the first two weeks of the trial and, ii) the distinct differences in community composition with age, even with a narrow age difference between piglets (1–6 days). We were able to appreciate a significant trend of increasing unrooted PD and decreasing BWPD with age in piglets that separated a 6 days maximum by day of birth from each other. Since age groups were confounded with breeds in our study, we attempted to determine the correlation within single 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.

Animal trials are often conducted in controlled environments so as to minimize environmental effects. However, individual variations such as breed and age are often unavoidable in large animal trials. Previously reported confounding factors include: individual variation \(^44\), cohabitation \(^92,93\), age \(^44,92,93\), maternal effects \(^92,93,95\), hormones \(^44\), behavioural differences between breeds (e.g. coprophagy, mouth to
mouth contact) and extent of long-term behavioural adaptation, which can differ between breeds for reasons not attributable to genetics. A litter effect was found in piglets at the start of the trial and was lost at later time points during the trial. This could be due to either of the aforementioned factors. Co-housing, aging and the splitting of the piglets in separate rooms to receive a different treatment, are possible causes for loss of the litter effect. In this study we confirm the importance of these factors in the contribution to inter-individual variability of gut microbial composition. Motta et al. (2019) report a correlation of beta diversity with age and no correlation of genotype and litter effect with either alpha or beta diversity. On the contrary, we found the piglet samples to significantly cluster by litter, breed and by age up to the second and the fourth week post weaning, in alpha diversity and beta diversity, respectively. We conclude that even small age differences among post-weaning piglets, down to the day, must be accounted for in an experimental set up.

Three groups of piglets (cohorts neomycin, neomycin + D-Scour™ and neomycin + ColiGuard®) underwent 5 days of treatment with the broad-spectrum antibiotic neomycin, via intramuscular administration. Intramuscular neomycin poorly diffuses (< 10%) into a healthy gastrointestinal tract, therefore a direct effect of neomycin on the gut microbiome may not be expected. However, neomycin showed a different trend in unrooted PD between the second and the third week of the trial, corresponding to the week following the neomycin treatment period for the neomycin cohort. Taking this time frame into consideration, the neomycin cohort did not increase in BWPD to the extent of the Control cohort. Although statistically significant differences between neomycin and Control in alpha diversity were not reached, the BWPD of the neomycin cohort appears to follow a different trend to the Control from the first week (during neomycin treatment) where neomycin treated piglets show the lowest decrease of BWPD compared to the control cohort and all other cohorts. While all cohorts show an increase in absolute phylogenetic diversity accompanied by a decrease of diversity evenness during this time frame, the neomycin cohort piglets show a lower drop in BWPD, suggesting an increase of species richness, without a corresponding loss of species evenness. Furthermore the neomycin cohort significantly separated from the control cohort in beta diversity in the third week of the trial, showing a higher representation of Mollicutes. Numerous studies report the link of oral antibiotic use with dysbiosis, as well as with host physiology changes. On the contrary, the effect of intramuscular antibiotic administration on the microbiome is less well investigated. Correlation between intramuscular antibiotic use and dysbiosis has been reported in fish, gorillas, humans, and pigs. In 1 day old piglets, a single IM injection of amoxicillin (penicillin class) is reported to have an effect on the intestinal microbiota, detectable days post treatment. Zeineldin et al. (2018) tested the effects of IM administration of several antibiotics of various classes (penicillin, macrolide, cephalosporin and tetracycline), in 8-week old piglets, reporting shifts of the Firmicutes/Bacteroidetes ratio following treatment (length of the treatment not reported). The effects of intramuscular administration of neomycin (aminoglycoside class) on the gut microbiota have to our knowledge not been investigated. Based on our results we conclude that a mild effect on phylogenetic diversity is appreciable post IM neomycin treatment, up to two weeks after termination of the treatment. Additional compositional and functional analysis is necessary to determine the source of this mild variation. Differences were not detected at later time points, based on our phylogenetic diversity.
analyses, suggesting a full recovery of the microbial communities after two weeks from the end of the treatment.

It is possible that the large shifts in phylogenetic diversity taking place in the first two weeks irrespective of the treatment (an increase, then decrease of unrooted PD, and an opposite trend of BWPD) have masked the milder effects of the treatment, despite our efforts to control for the effects of aging. This could be the reason why a significantly distinct alpha diversity trend was found in the neomycin + D-Scour™ cohort compared to the neomycin cohort, but not in the D-Scour™ cohort compared to the Control cohort. The neomycin + D-Scour™ cohort underwent 5 days of neomycin treatment followed by 2 weeks of D-Scour™ treatment. A significant increase of BWPD was detected in the two-week period of D-Scour™ treatment, indicating a possible enhancement of microbiome evenness following neomycin treatment. To our knowledge there are no studies reporting an increased evenness in piglet gut community composition following a specific probiotic treatment. There are instead multiple studies reporting beneficial effects of probiotic treatment in sucker and weaner piglets in terms of improved gut mucosal integrity, growth rate, digestibility of proteins and water absorption, reduction of pathogen invasion efficiency, and decreased mortality. Although the assessment of physiologic changes from probiotic treatments was outside the scope of this study, we found significant separation of neomycin + D-Scour™ cohort samples to neomycin cohort samples in beta diversity 3 and 10 days after D-Scour™ treatment, where neomycin + D-Scour™ samples showed a higher representation of Lactobacillales compared to neomycin samples, suggesting a 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 third principal component (explaining 7.88% of the variation) two weeks post 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) 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 taxa 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. For this reason it can be expected that some of the taxa we found to correlate with the weight of the piglets (eighty-three distinct species) could be spurious while other correlations may have been missed.

**Technical controls in metagenomic studies and methodological limitations**

Taxonomic assignment of the raw reads from the positive controls was performed with MetaPhlAn2 which relies on a ca. 1M unique clade-specific markers derived from 17,000 reference genomes. Such a database to map against the positive controls suffices as these organisms are cultivable, and for this
reason they are widely studied hence the sequences are known. This is not the case for real-world samples where mapping against a database (which completeness relies on studied and often cultivable organisms) would narrow the view on the true diversity within the sample.

Positive controls with well-studied members and known ratios within the samples, has proven to be a valuable tool to assess consistency among technical replicates across batches and to detect possible biases derived from the DNA extraction method.

Systematic taxonomic bias in microbiome studies, resulting from differences in cell wall structures between Gram positive and Gram negative bacteria, have previously been reported; sample treatment with enzymatic cocktails can modestly reduce this bias $^{102-104}$. Although we implemented this step in our workflow, it seems that, from the read abundance of our mock community, which contained three Gram negative and four Gram positive strains, a bias towards Gram negative taxa may still be present.

In terms of contamination we concluded that: a) contamination in our study was not batch specific; b) a problem of sample cross-contamination existed at the DNA extraction step between neighbouring wells. During the bead beating step of DNA extraction, the deep-well plate is sealed with a sealing mat, rotated and placed in a plate shaker for the bead beating to take place. We consider that sample cross-contamination is most likely to occur during this step.

**Conclusion**

Our study provides a publicly available databank of the pig gut metagenome. Our findings further stress the importance of confounding factors such as breed, age and maternal effects when assessing the effect of treatment on the gut microbiome. We found that age, even within a narrow age span (1–6 days) can have an impact on microbial shifts and should be accounted for in microbiome studies. Intramuscular neomycin treatment correlated with a clustering in alpha diversity and a higher representation of *Mollicutes* compared to control. D-Scour™ treated piglets displayed a mild shift in alpha diversity compared to control, and 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 taxa. Age was the strongest factor shaping phylogenetic diversity of the piglets.

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

**Methods**

**Pig trial**
Animal studies were conducted at the Elizabeth Macarthur Agricultural Institute (EMAI) NSW, Australia and were approved by the EMAI Ethics Committee (Approval M16/04). The trial animals comprised 4-week old male weaner pigs ($n = 126$) derived from a commercial swine farm and transferred to the study facility in January 2017. These were cross-bred animals of Landrace, Duroc and Large White breeds and had been weaned at approximately 3 weeks of age.

The pig facility consisted of four environmentally controlled rooms (Rooms 1–4) with air conditioning, concrete slatted block flooring with underground drainage and open rung steel pens ($\text{S1}$). Each room had nine pens, consisting of a set of six and a set of three pens, designated a-f and g-i respectively, with the two sets of pens being physically separate, i.e. animals could come in contact with each other through the pen's bars within each set of pens, but not between sets. The rooms were physically separated by concrete walls and contamination between rooms was minimized by using separate equipment (boots, gloves, coveralls) for each room. In addition, under-floor drainage was flushed twice weekly and the flushed faeces/urine was retained in under-floor channels that ran the length of the facility, so that Rooms 1, 2 were separate from Rooms 3, 4 and flushing was in the direction 1 to 2 and 3 to 4.

The pigs were fed ad libitum a commercial pig grower mix of 17.95% protein free of antibiotics, via self-feeders. On the day of arrival (day 1) 30, 18, 18, and 60 pigs were allocated randomly to Rooms 1, 2, 3 and 4 respectively in groups of 6, 6, 6 and 6–7 pigs per pen respectively ($\text{S1A}$). Pigs were initially weighed on day 2, and some pigs were moved between pens to achieve an initial mean pig weight per treatment of approximately 6.5 kg (range: 6.48–6.70; mean ± SD: 6.53 ± 0.08). Pigs were weighed weekly throughout the trial ($\text{Supplementary file 1}$). Behaviour and faecal consistency scores were taken daily over the 6-week period of the trial ($\text{Supplementary file 2}$). Developmental and commercial probiotic paste preparations ColiGuard® and D-Scour” from International Animal Health, Australasia, were used in some treatment groups.

The animals were acclimatised for 2 days before the following treatments were administered: Room 1 - oral 1 g/pig of placebo paste daily for 14 d; Room 2 - oral 1 g/pig of D-Scour™ paste daily for 14 d; Room 3 - oral 1 g/pig of ColiGuard® paste daily for 14 d; Room 4 - intramuscular (IM) injection of antibiotic administered at 0.1 mL per pig daily from a 200 mg/mL solution for a total treatment duration of 5 d.

On the day following the final neomycin treatment (day 8), 36 pigs were moved from Room 4 to Room 2 ($n = 18$, 6 in each pen, pens g-i), and to Room 3 ($n = 18$, 6 in each pen, pens g-i) ($\text{S1}$). The following day (day 9), oral administration of D-Scour™ (1 g/pig) and of ColiGuard® (1 g/pig) commenced for pigs in Room 2 pens g-i and in Room 3 pens g-i, respectively, and continued for a period of 14 days. Assignment of the 36 neomycin-treated pigs to the treatment groups neomycin + D-Scour™ ($n = 18$; Room 2 pens g-i) and neomycin + ColiGuard® ($n = 18$; Room 3 pens g-i), was carried out by distributing them so that the mean weight of the animals distributed across pens and rooms was similar ($\text{Supplementary file 1}$). By this time point, each occupied pen in the trial housed six pigs. ($\text{S1B}$) From that time, twelve piglets from the original 126 were no longer present, as they had been euthanised as pre-treatment controls at the start of the trial.
Faecal samples were collected from all piglets once per week and from a subset (n = 48 pigs; 8 from each of the six cohorts) twice per week over the 6-week study period (Fig. 1). From each piglet, faeces were collected per rectum with new disposable gloves; where minimal or no faeces could be collected on a collection day, sampling was performed the following morning. Samples were placed in 50 mL Falcon tubes and stored at 4°C within 30 min of sample collection for a minimum of 30 min and a maximum period of 6 h.

**Faecal sample processing**

Samples (3 g/pig) were mixed with 15 mL PBS (200 mg/mL), in sterile stomacher bags and homogenized with a Bio-Rad stomacher. The homogenised samples were divided in replicates: one replicate was stored directly at -80°C and one replicate was supplemented with glycerol (20% v/v) (Sigma-Aldrich) then stored at -80°C. In addition, single time-point faecal samples from the dams of the trial pigs (n = 42) were obtained from the commercial facility of origin and were pre-processed at EMAI as described above. Thus, a total of 911 unique samples, between one and ten samples per subject (mean: 4.8; median: 3) (S2), were obtained throughout this study. At the end of the trial period, all samples were transported from EMAI to the University of Technology Sydney (UTS) for further processing.

**Positive controls**

As a positive control “mock community” for this study, four Gram positive (*Bacillus subtilis* strain 168, *Enterococcus faecium*, *Staphylococcus aureus* ATCC25923, *Staphylococcus epidermidis* ATCC35983) and three Gram negative (*Enterobacter hormaechei* CP_032842, *Escherchia coli* K-12 MG1655, *Pseudomonas aeruginosa* PA01) bacterial strains from −80°C stocks were cultured at 37°C for 16 h in LB (Luria-Bertani) then centrifuged at 14,000 rpm for 10 min. From the resulting pellets, 1 g was transferred to 1 mL of LB and homogenised and a 1:10 dilution of this was made for each bacterial culture. Ten microliters of bacterial suspension from each of the cultures was used to determine the number of colony forming units (CFU) in the original suspension in the following manner: by further diluting tenfold in LB and by plating onto 1.6% LB agar plates and incubacted overnight. The remaining suspensions (990 µL from each bacterial culture) were pooled into a sterile tube, then aliquoted into Eppendorf tubes in 500 µL volumes/tube. As a washing step, Eppendorf tubes were centrifuged at 14,000 rpm for 10 min, 500 µL PBS was added to the pellet and subsequently resuspended. These tubes constituted the mock community samples and were stored at -80°C. Expected proportions of the mock community members were determined from the estimated colony forming units (CFU) multiplied by the genome size and were as follows: 8.7:13.0:7.7:16.7:38.9:14.5:0.4 for *S. aureus*, *B. subtilis*, *E. faecium*, *S. epidermidis*, *P. aeruginosa*, *E. cloacae*, and *E. coli* respectively.

The two probiotic formulations used in this study were used as two additional positive controls. D-Scour™ is a commercially available probiotic formulation for livestock, with each gram containing 180 million CFU of the following: *Lactobacillus acidophilus*, *Lactobacillus delbrueckii* subspecies *bulgaricus*, *Lactobacillus plantarum*, *Lactobacillus rhamnosus*, *Bifidobacterium bifidum*, *Enterococcus faecium*,
*Streptococcus salivarius* subspecies *thermophilus*, with an additional 20 mg of garlic extract (*Allium sativum*). The probiotic ColiGuard is a probiotic formulation developed for the treatment of enterotoxigenic *Escherichia coli* (ETEC) in weaner pigs, developed in collaboration between the NSW DPI and International Animal Health Products, containing undefined concentrations of *Lactobacillus plantarum* and *Lactobacillus salivarius*.

**DNA extraction**

Piglet and sow faecal samples, mock community samples, negative controls and probiotic samples (D-Scour™ and ColiGuard® paste) were allocated to a randomized block design to control for batch effects in DNA extraction and library preparation. The faecal samples were thawed on ice first, followed by the probiotics and mock community samples. MetaPolyzyme (Sigma-Aldrich) treatment was performed according to the manufacturer’s instructions except for the dilution factor, which we allowed to be 4.6 times higher. Immediately after incubation, DNA extraction was performed with the MagAttract PowerMicrobiome DNA/RNA EP kit (Qiagen) according to the manufacturer’s instructions. Quantification of DNA was performed using PicoGreen (Thermofisher) and measurements were performed with a plate reader (Tecan, Life Sciences) using 50 and 80 gain settings. All samples were diluted to 10 ng/µL.

**Library preparation**

Sample index barcode design using a previously introduced method resulted in a set of 96 × 8nt sequences with a 0.5 mean GC content and none of the barcodes containing 3 or more identical bases in a row. Nine hundred-sixty different combinations of i5 and i7 primers were used to create a uniquely barcoded library for each sample. The detailed sample-to-barcode assignment is given in Supplementary file 3. Library preparation was carried out using a modification of the Nextera Flex protocol to produce low bias, low cost shotgun libraries, as described in a previous manuscript. Following the amplification step, samples were centrifuged at 280 × g for 1 min and stored between 1 and 5 days at 4 °C.

**Size selection and purification**

Samples from the same 96-well plates were pooled into one tube by taking 5 µL from each library. This generated 10 pooled samples, one for each plate. A master pool was created by pooling 5 µL from the pool of each plate into a single pool. Forty microliters from each of the 10 plate pools and 40 µL from the master pool underwent library size selection and purification using equal volumes of SPRIselect beads (Beckman Coulter) and ultrapure water (Invitrogen). Sample cleaning with SPRI-beads was performed as described previously. A purified master pool comprising samples from all plates, and purified pools of individual plates to check for plate-specific anomalies, were diluted to 4 nM and fragment size distribution was assessed using the High Sensitivity DNA kit on the Bioanalyzer (Agilent Technologies, USA).

**Normalisation and sequencing**
The master pool was sequenced on an Illumina MiSeq v2 300 cycle nano flow cell (Illumina, USA). Read counts were obtained and used to normalise libraries. The liquid handling robot OT-One (Opentrons) was programmed to re-pool libraries based on read counts obtained from the previous MiSeq run. The script used to achieve the normalization is available through our Github repository.

The read count distribution after normalisation is displayed in supplementary figure (S3). The normalized and purified pooled library was sequenced on an Illumina NovaSeq S4 flow cell at the Ramaciotti Centre for Genomics (Sydney, NSW, Australia), generating a total of 27 billion read pairs from 911 samples.

**Sequence data processing**

Adapter trimming (parameters: k = 23 hdist = 1 tpe tbo mink = 11), PhiX DNA removal (parameters: k = 31 hdist = 1), and quality filtering (parameters: ftm = 0 qtrim = r trimq = 20), were performed using bbdruk.sh (http://jgi.doe.gov/data-and-tools/bbtools). Quality assessment of raw reads was carried out using FASTQC (http://www.bioinformatics.babraham.ac.uk/projects/fastqc/) and a combined report of all samples was obtained with MULTIQC (Supplementary File 4). The presence of PCR duplicates was assessed by feeding read pairs to dedupe.sh (http://jgi.doe.gov/data-and-tools/bbtools) (parameters: ac = f). Nextflow was used to manage processing of the data on the HPC.

**Determination of microbial diversity among samples**

Phylogenetic diversity of all samples was assessed with PhyloSift using the first 100 k read pairs of each sample (parameters: --chunk-size 100000 --paired). In order to test for associations of phylogenetic diversity with treatment, time of sampling, and differences among hosts at the start of the trial, analysis of the unrooted phylogenetic diversity (PD), the balance weighted phylogenetic diversity (BWPD) and principal component analysis (PCA) of the Kantorovich-Rubinstein distances (beta diversity analysis) were performed. Alpha-diversity and beta-diversity were analyzed and the results were visualized with R and R packages. The data analysis is schematically represented in Supplementary Fig. 4 (S4).

Additionally, SortmeRNA (version 2.1) was applied to extract 16S ribosomal RNA genes from raw reads. The rRNA reference database silva-bac-16 s-id90 was used. Hits were filtered (e-value < = 1e-30) and PCA was computed with R. Sample counts were normalized for library size by proportions and were tested with the Spearman's Rank correlation coefficient method to find taxa correlating with the weight of the piglets across the trial.

**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 (S5), a batch effect was found by multiple comparison analysis with ANOVA and by applying Tukey post-hoc correction to pairwise comparisons. Batch effects were detected (ANOVA, alpha
diversity: \( p \) range = 4.8e-12 - 0.00011; beta diversity: \( p \) value < 0.0023) (S6; Supplementary file 5) and removed with COMBAT\(^7\) (ANOVA \( p \) value = 1) (S7; Supplementary file 5).

**Technical analysis of positive controls**

As a quality control step, taxonomic analysis of the positive control samples and their technical replicates was performed with MetaPhlAn2\(^10\). The positive controls included the mock community \((n = 8)\), D-Scour™ \((n = 8)\), and ColiGuard® \((n = 8)\) samples.

**Abbreviations**

| Abbreviation | Description |
|--------------|-------------|
| AMR          | Antimicrobial resistance |
| MDR          | Multidrug resistance |
| IM           | Intramuscular |
| CFU          | Colony forming units |
| PD           | Phylogenetic diversity |
| BWPD         | Balance-weighted phylogenetic diversity |
| PCA          | Principal component analysis |
| PC           | Principal component |
| ANOVA        | Analysis of variance |
| TukeyHSD     | Tukey’s honestly significant difference |

**Declarations**

**Ethics approval**

Animal studies were conducted at the Elizabeth Macarthur Agricultural Institute (EMAI) NSW, Australia and were approved by the EMAI Ethics Committee (Approval M16/04).

**Consent for publication**
Not applicable

**Availability of data and material**

The raw sequencing data has been submitted to the NCBI Short Read Archive under project PRJNA526405. Metadata and scripts are available in this article, its Supplementary material and in our Github repository https://github.com/GaioDany/metapigs_base

**Competing interests**

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

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**Contributions**

D.G is the main author of this paper. T.A.C., L.F., D.G., T.Z., G.J.E., collected and processed the samples. T.A.C., S.P.D., A.E.D. conceived and designed the project. D.G., M.L. performed the DNA extraction. D.G., M.L., K.A., A.E.D. contributed to the library preparation and the robot pooling. M.Z.D., D.G., A.E.D. processed the sequencing data. D.G., A.E.D. performed the data analysis. A.E.D., G.J.E., John Webster edited the manuscript. All authors have read, revised, and approved the manuscript.

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**Figures**

![Timeline](image)

**Figure 1**

Timeline. Timeline of the trial indicating the start and the length of the treatment for each cohort. Piglets arrived on the site of the trial on January 30th and were allowed 2 days of acclimatisation before the start of the treatments (pink: placebo paste; yellow: probiotic D-Scour™ formulation; green: probiotic ColiGuard® formulation; blue: antibiotic neomycin intramuscular injection). Vertical lines indicate main days of sampling where all piglets were sampled (n=126). Small arrows indicate days of sampling for a subset of the piglets (8 per cohort; n=48).
Figure 2

Significance of correlations between phylogenetic diversity and specified factors. Plots of p values derived from Kruskal-Wallis analysis of variance (black) and adjusted p values by Benjamini-Hochberg method (red). x-axis displays distinct sampling time points in chronological order. Correlation of specified factors of samples are shown with (A) alpha diversity and (B) beta diversity. Abbreviations: DxL = “Duroc x Landrace” cross breed; DxLW = “Duroc x Large White” cross breed.
Figure 3

Effect of time on beta diversity. Principal component analysis (PCA) of samples. PCA from edge component analysis with PhyloSift (top) and PCA from 20 most abundant 16S reads extracted with SortMeRNA (bottom). In the top, distribution of samples on either side of the plot (left versus right; top versus bottom) reflect the taxa that were found to explain the variation. Samples are colour coded by days into the trial. In the lower plot, arrows indicate which of the 20 taxa read contributed to the variation.
of samples, where arrows thickness represents a higher (thicker) or lower (thinner) contribution. Samples are colour coded by time point during the trial.

Figure 4

Abundance heatmap. Most abundant taxa within each time point are obtained from analysis with guppy fat. Abundance is derived from the branch width. The distance between each time point is of one week.
Supplementary Files

This is a list of supplementary files associated with this preprint. Click to download.

- metapigsbaseallfigures.pdf
- SupplementaryFile5.xlsx
- Supplementaryfile4.html
- Supplementaryfile3.csv
- Supplementaryfile2.xlsx
- Supplementaryfile1.hlsx.xlsx