A new technique for use in the study of the microbiome: An evaluation of a three-dimensional cell culture technique in maintaining the gastrointestinal microbiome of four Balb/c female mice and implications for future studies

Maintaining the integrity of an explant microbiome and implications for future studies

Everest Uriel Castaneda\textsuperscript{1,2}, Jeff Brady\textsuperscript{1}, Janice Speshock\textsuperscript{2}

\textsuperscript{1}Texas A&M AgriLife Research and Extension Center, Stephenville, Texas, United States of America

\textsuperscript{2}Department of Biological Sciences, Tarleton State University, Stephenville, Texas, United States of America

*Corresponding author

Email: Speshock@Tarleton.edu
Abstract (Level 1)

Fluctuations in oxygen, pH, nutrients, or other factors such as food or pharmaceuticals, may perturb the microbiota of the gastrointestinal (GI) tract. This environmental variation is a cause for concern given dysbiosis of the microbiome is correlated with disease states; thereby, model organisms are utilized to study microbial communities during, after, or before shifts in microbes since intact ex vivo microbiomes have historically been challenging to utilize. The objective of this study is to culture an explant microbiome of 4 Balb/c, laboratory bred mice to develop an ex vivo tool for future microbiome studies. We cultured homogenates of the distal colon of 4 mice in three dimensional, 24 well plate culture dishes. These dishes were incubated for 24 hours in two different oxygen concentration levels, 0% and 20%. The pH of the plate was tested before and after incubation. To analyze the integrity of the microbiome, we utilized 16S sequencing. Further, we utilized 16S metagenomics to characterize fecal samples and colon samples to speculate whether future studies may utilize feces in constructing an explant microbiome to spare animal lives. We found that pH and familial relationship had a profound impact on community structure while oxygen did not have a significant influence. The feces and the colon were similar in community profiles, which lends credence to utilizing feces in future studies. In addition, our efforts successfully cultured archaea, which included difficult to culture strains such as Miscellaneous Crenarchaeota group (MCG) and Methanobacteria. Ultimately, further attempts to culture and preserve an animal’s microbiome needs to control for and maintain stable pH.
The microbiome forms a symbiotic relationship with its host [1]. Essentially, microbes have a cooperative role in the GI tract and contribute to a host’s immune system and metabolism [2–5]. Although the natural relationship between the microbiome and the host is essential, overpopulation by an undesirable species, or dysbiosis, has been linked to particular diseases and phenomena such as autism spectrum disorder [6], cancer [3], and obesity [7]. Historically, researchers have utilized animal models such as gnotobiotic mice to study microbiome-animal interactions but experimentation with such mice is expensive [8–10]. Thus, an *ex-vivo* model of the microbiome that provides cost-effective, reproducible, and reliable results is highly desirable in studying this dynamic [11].

Mouse microbiome studies have expanded our views on the impact of prokaryotes on digestion, disease, and even behavior, but many microbial species cannot persist in culture [12,13]; therefore, most research currently relies on germ-free mice for microbiome studies, which can be cost-prohibitive for many laboratories [7,8,14]. To further assess this transient mixture of microbiota [15], scientists have been utilizing culture independent, next-generation sequencing to inquire about shifts within the microbiome and what stimuli affect these changes in composition [16]. With the decreasing cost of next-generation sequencing, an influx of research has been possible in this area [17]. It is worth noting the financial and ethical burden of raising, sacrificing, and housing model organisms [9,10]; therefore, it would be beneficial to develop techniques to save organisms and further decrease costs.

In this study, we cultured and maintained the GI microbiomes of 4 laboratory-bred female Balb/c mice in three-dimensional (3D) well plates, partitioned into 2 oxygen levels. Due to the variable
nature of the oxygen levels of the GI tract [18], we cultured 3D plates in both a conventional incubator and an anaerobic chamber, both at 37 degrees Celsius. Additionally, we determined the microbial composition of the mouse stool and the distal colon to observe if future studies may utilize feces and avoid sacrificing organisms altogether. We utilized sequencing methods to verify final proportional community composition of each sample.

Materials and Methods (Level 1)

Subjects (Level 2)

The study was performed under a protocol approved by the Tarleton State University Institutional Animal Care and Usage Committee (Animal Use Protocol 12-009-2016-A1). 4 Balb/c females 8 weeks in age were utilized in this experiment. Females were housed together and raised on identical chow diets and similarly weaned. Mice 1 and 2 were siblings while mice 3 and 4 were siblings. The siblings were born from different dams and sires. All mice were held in sterile containers and euthanized with 150 µL of sodium pentobarbital delivered intraperitoneally. Post injection, mice shed two to three samples of stool which were recovered utilizing sterile forceps and immediately frozen. Once deceased, 2.5 cm of the large distal colon were removed. After, two to three small additional 0.5 cm samples of the large distal colon were excised from the specimen and immediately frozen. Colon tissue extractions were added to a sterile tissue grinder along with 5 mL of Dulbecco’s Modified Eagle Medium (DMEM). The sample was manually homogenized into a liquid solution. Homogenate was checked for pH by applying a small droplet with a mechanical pipette onto litmus paper.

Culture method (Level 2)
Cultures were established in sterile 24 well plates with multiwell tissue culture inserts, 8 µm pore size (Corning Incorporated, New York). Corning PuraMatrix Peptide Hydrogel was prepared using 8 mL of molecular grade water and 20 ml of hydrogel to create a 0.25% solution. 150 mL of the prepared solution were added to each of the 12 well inserts. In addition, 500 mL of supplementary DMEM was added under each well insert. This technique is a modified version of 3D tissue culture repurposed for culturing our ex-vivo microbiome. Once the 3D culture plates were prepared, 250 mL of the homogenized colon were added to each of the prepared 12 well inserts. Plates were checked for baseline pH by transferring a small drop of medium with a mechanical pipette onto litmus paper. The plates were then added to a single incubator, but to create an anoxic environment, half of the plates were incubated in an anaerobic chamber. Plates were incubated for 24 hours. Each insert’s medium was then tested for pH again and transferred into sterile 2.5 mL storage tubes and frozen for future DNA extraction.

DNA extraction and library production (Level 2)

DNA was extracted from each sample using a solid phase extraction protocol from Brady et al. [19]. After extraction, DNA was amplified utilizing prokaryote-specific primers, 519F 5’-CAGCMGCCGCGGTAA-3’) and 785R (5’- TACNVGGGTATCTAATCC-3’), that target the V4 region of the 16Ss rRNA [20,21]. PCR amplification was accomplished through denaturation at 95°C for 3 minutes, followed by 35 cycles of 95°C for 10 seconds, 55°C for 30 seconds, and 72°C for 30 seconds. Dual 6 bp DNA barcodes were added to sequencing libraries using the same PCR protocol and Illumina P5 and P7 flowcell binding adapters [22]. Sequencing libraries were size-selected with a Pippin Prep instrument (Sage Science, Beverly, MA) to a length of 300-600 base pairs. Sequencing was conducted on a MiSeq instrument using 600 cycle paired end v3 sequencing kits at the Texas A&M University Genomics Core Facility. Raw sequences
were processed with QIIME [23] and USEARCH [24]. Taxonomy was assigned using the
Greengenes 13.8 database [25] as a reference with UCLUST [24], and reference-based
Operational Taxonomic Unit (OTU) picking was conducted at 97% sequence similarity using the
RDP [26] method in QIIME.

**Statistical analysis (Level 2)**

Cumulative Sum Scaling [27] was utilized to normalize the data. Biom files were constructed
with QIIME [28] and transferred into R [29] and Microsoft Excel for further statistical analysis.
Phyloseq [30], ggplot2 [31], and vegan [32] packages were utilized to evaluate alpha and beta
diversity with seed set at 1400. Alpha diversity was assessed using the Shannon diversity index.
Variation in alpha diversity for oxygen, pH, mouse, feces, and colon comparisons were first
checked for normality using the Shapiro-Wilk test for normality [33]. The data was non-normal
in distribution (Shapiro-Wilk test, w=0.9506, p<0.01); therefore, comparisons were made with
non-parametric tests. All multivariate tests were corrected using false discovery rate (FDR) [29].
Comparisons of alpha diversity were conducted using Kruskal Wallis one-way analysis of
variance (KW ANOVA) and Wilcoxon rank sums test (Wilcoxon test) while comparisons of beta
diversity were assessed with unweighted unifrac distance metrics at 1000 permutations using
permutational multivariate analysis of variance (PERMANOVA). Dunn’s test, post-hoc analysis
was conducted using the dunn.test package in R [34]. In addition, non-parametric t-tests were
used for comparisons of mean abundance in individual bacterial strains between samples.
Principle coordinate analysis (PCoA) and canonical correspondence analysis (CCA) were
performed at the level of OTU using unweighted unifrac distance metrics. PRIMER 7 [35] was
used for the hypothesis testing utilizing square root transformed Bray-Curtis ordination data at
9999 permutations. A microbial network was constructed using the Co-occurrence Network
Interferences (CoNet) [36] application for Cytoscape [37]. Feces and colon data were removed before CoNet analysis. CoNet has been utilized in previous studies to investigate defined interactions between microbes [38–40]. Spearman correlation coefficient with a cutoff ratio of 0.6 was utilized, and to focus the network, only microbes with sequence counts greater or equal to 20 were included. 1,000 permutations were accomplished through a bootstrapping method with an FDR correction [39].

Results (Level 1)

**pH and oxygen (Level 2)**

pH readings of each plate were taken before and after incubation. As shown in Table 1, pH fluctuated from the original homogenate and baseline (before 24-hour incubation). In addition, mouse samples maintained varying levels of pH which correlated with differences in oxygen concentration (Table 1).

Table 1. Sample size, pH, and oxygen level.

| Mouse | Homogenate pH | Sample Size | Oxygen Level | Plate Baseline pH | Plate Final pH |
|-------|---------------|-------------|--------------|-------------------|---------------|
| M1    | 7.5           | 12          | 20%          | 8                 | 10            |
|       |                | 11          | 0%           | 8                 | 9             |
| M2    | 7.5           | 11          | 20%          | 8                 | 10            |
|       |                | 12          | 0%           | 8                 | 9             |
| M3    | 7.5           | 12          | 20%          | 8                 | 7             |
|       |                | 11          | 0%           | 8                 | 6             |
| M4    | 7.5           | 12          | 20%          | 8                 | 7             |
|       |                | 12          | 0%           | 8                 | 6             |

16s rRNA sequencing results (Level 2)
We analyzed microbiomes from the 4 mice at varying oxygen levels after culturing them *in vitro*.

In the 24 well plate system, we utilized 12 membrane inserts to culture the microbiomes. After mice succumbed to euthanasia, colon samples were harvested to complement fecal shedding. Mouse 1, 2, and 4 shed two fecal samples each while mouse 3 shed three fecal samples; therefore, we had a total of 9 colon and 9 fecal samples across 4 mice. After sequence quality filtering we had a total sample size of 111 samples and 3,133,666 sequences total (Table 2).

| Mouse | Sample Size | Total Sequence Count | Average Sequence Count |
|-------|-------------|-----------------------|------------------------|
| M1    | 27          | 234051                | 8669                   |
| M2    | 27          | 384215                | 14230                  |
| M3    | 29          | 1243981               | 40128                  |
| M4    | 28          | 1194017               | 45924                  |

**Fecal and colon comparison (Level 2)**

The feces and the colon samples were characterized for microbial composition at the phylum and family level (Figs 1A and 1B, respectively). Community composition was dominated by the phyla Firmicutes and Bacteroidetes (Fig 1A), with means of 47% and 49%, respectively, and standard deviations (SD) of 23%. In addition, the *Bacteroidales* family S24-7 was highly abundant with a mean of 42% and a SD of 20% (Fig 1B). These results are consistent with recent microbiome studies of mice [41–43]. Beta diversity analysis for each of the feces and colon samples showed no difference in composition (Table 3), and alpha diversity analysis (Fig 1C) also revealed no difference (KW ANOVA, chi-squared = 6.64, df = 7, p = 0.47). Therefore, all colon samples were pooled together, and all feces samples were pooled together for a statistical comparison of feces and colon. The Shannon diversity index was utilized for comparison of the
bulk samples (Fig 1D). Results showed no difference between the feces and colon (Wilcoxon test, p = 0.44). In addition, beta diversity comparison showed no difference (PERMANOVA, Pseudo F = 1.06, p = 0.37). Since we found that feces and colon samples are similar, we pooled sequences from all feces and colon samples together into one bulk sample, named “microbiome,” for diversity comparisons with cultured microbiomes.

Table 3. Results of the pairwise PERMANOVA tests between feces and colon samples.

| Comparison          | Pseudo F | p-score |
|---------------------|----------|---------|
| M1 Feces vs M2 Feces| 3.16     | 0.32    |
| M1 Feces vs M3 Feces| 2.75     | 0.34    |
| M1 Feces vs M4 Feces| 2.60     | 0.31    |
| M2 Feces vs M3 Feces| 3.15     | 0.33    |
| M2 Feces vs M4 Feces| 2.8      | 0.34    |
| M3 Feces vs M4 Feces| 3.01     | 0.30    |
| M1 Colon vs M2 Colon | 1.47 | 0.34 |
| M1 Colon vs M3 Colon | 1.46 | 0.34 |
| M1 Colon vs M4 Colon | 1.40 | 0.33 |
| M2 Colon vs M3 Colon | 1.39 | 0.31 |
| M2 Colon vs M4 Colon | 1.29 | 0.33 |
| M3 Colon vs M4 Colon | 1.41 | 0.33 |

Fig 1. Community composition of fecal and colon samples. (A) Relative abundance of bacterial phyla. Phyla with observations less than 1% are pooled into “Other” category. The first two characters represent the mouse from which the organ and stool were dissected. “S” denotes the different samples acquired. (B) Relative abundance of bacterial families. Families with observations less than 1% are pooled into “Other” category (C) Comparison of Shannon diversity between feces and colon samples. “ns” means non-significant, p>0.05. (D) Shannon diversity comparison of pooled feces and colon samples.

Microbiome comparison (Level 2)

In the explanted microbiomes, Firmicutes and Bacteroidetes were the dominant phyla (Fig 2A). Firmicutes had the highest average relative abundance, 70% (SD 28%), with Bacteroidetes averaging 18% (SD 16%). Across all cultures, Enterococcus was highly abundant (Fig 2B) having a mean of 47% (SD 32%). The Shannon diversity index was different between the
explanted cultures and the microbiome of the mice (KW ANOVA, chi-squared = 73.58, df = 8, p < 0.01). Post-hoc analysis shows that, compared to the microbiome, mouse cultures 1 and 2 were the same while mouse cultures 3 and 4 differed (Fig 2C). The microbial profile of each sample revealed a difference in beta diversity between the plates and the microbiome (Table 4), which is also reflected in the PCoA plot (Fig 2D).

Table 4. Results of the pair-wise PERMANOVA tests between cultured plates and harvested samples.

| Comparison          | Pseudo F | p-score |
|---------------------|----------|---------|
| M1 0% vs Microbiome | 11.09    | p < 0.01|
| M1 20% vs Microbiome| 18       | p < 0.01|
| M2 0% vs Microbiome | 11.93    | p < 0.01|
| M2 20% vs Microbiome| 15.21    | p < 0.01|
| M3 0% vs Microbiome | 14.94    | p < 0.01|
| M3 20% vs Microbiome| 7.95     | p < 0.01|
| M4 0% vs Microbiome | 18.83    | p < 0.01|
| M4 20% vs Microbiome| 21.321   | p < 0.01|

Fig 2. Community composition of cultures and comparison of microbiome. (A) Relative abundance between bacteria Phyla. Phyla with observations less than 1% are pooled into “Other” category. The first two characters represent the mouse, and the percent oxygen used in culture conditions is noted. (B) Relative abundance of bacterial genera. Genera with observations less than 5% are pooled into “Other” category. (C) Results of the post-hoc, Shannon Diversity, pairwise comparisons between cultures and the microbiome. p<0.05 is noted by “*”, p<0.0001 is noted by “****”, and non-significance is noted by “ns.” (D) Unweighted unifrac PCoA plot for plates and microbiome comparison. 12 well plates are denoted by the mouse in which they were derived from, “M,” and the percent oxygen.

Environmental variables (Level 2)

A community profile of cultural composition due to varying levels of oxygen and pH was constructed at the phylum and genus levels (Fig 3A-D). No difference in alpha diversity existed between the two oxygen levels (Wilcoxon test, p = 0.34; Fig 3E). Results showed similarity in beta diversity (PERMANOVA, Pseudo F = 1.25, p = 0.21). Multivariate analysis shows a significant difference in alpha diversity associated with differences in pH (KW ANOVA, chi-
squared = 58.13, df = 3, p < 0.01). Post-hoc analysis revealed that plates reaching a pH of 6 and
7 were similar while all other comparisons differed (Fig 3F). Comparisons of microbial
communities also showed a marked difference between plates of varying pH levels (Table 5).
Additionally, CCA and PERMANOVA revealed that oxygen did not contribute to community
clustering (Table 6; Fig 4).

Table 5. Results of the pairwise PERMANOVA test between pH groups.

| Comparison   | Pseudo F | p-score |
|--------------|----------|---------|
| pH 6 vs pH 7 | 1.86     | 0.04    |
| pH 6 vs pH 9 | 17.15    | p < 0.01|
| pH 6 vs pH 10 | 25.46    | p < 0.01|
| pH 7 vs pH 9 | 15.27    | p < 0.01|
| pH 7 vs pH 10 | 22.31    | p < 0.01|
| pH 9 vs pH 10 | 1.62     | 0.03    |

Table 6. Hypothesis testing for sources of variation based on PERMANOVA analysis.

| Source                     | df | SS   | MS   | Pseudo F | P(perm) | Unique perms |
|----------------------------|----|------|------|----------|---------|--------------|
| Oxygen                     | 1  | 3432.6| 3432.6| 1.6677   | 0.0756  | 9903         |
| Sibling Relationship       | 1  | 52930 | 52930| 25.716   | 0.0001  | 9921         |
| Oxygen x Sibling Relationship | 1  | 2407.4| 2407.4| 1.1696   | 0.2437  | 9915         |
| Residuals                  | 89 | 1.8319E+05| 2058.3|         |         |              |
| Total                      | 92 | 2.4194E+05|      |         |         |              |

df, degrees of freedom; SS, sum of squares; MS, mean square.
The only significant effect fitted in the PERMANOVA is sibling relationship (fixed factor) while oxygen (fixed factor) is not significant. Sibling relationship explains the changes in community composition as time elapsed in the incubators.

Fig 3. Microbial structuring due to environmental factors. (A) Relative abundance between bacteria phyla. Phyla with observations less than 1% are pooled into “Other” category. (B) Relative abundance of bacterial genera. Genera with observations less than 1% are pooled into “Other” category. (C) Shannon diversity comparison between oxygen concentrations. Non-significance is shown by “ns.”
Fig 4. CCA for the effects of pH, oxygen, and sibling relationship on community structuring.

**Siblings (Level 2)**

A marked difference in cultural composition was noted by familial relationship (Table 6, Fig 4, Fig 5A, B). Not only was clustering associated with siblings, shown in Fig 4, but Shannon diversity significantly varied (KW ANOVA, chi-squared = 56.24, df = 3, p < 0.01). Post-hoc analysis showed mouse 1 and mouse 2 were similar and varied from mouse 3 and mouse 4, which were also similar (Fig 5C). Additionally, beta diversity varied according to familial relationship (PERMANOVA, Pseudo F = 12.82, p < 0.01).

**Fig 5. Microbial composition by mouse.** (A) Relative abundance between bacteria phyla. Phyla with observations less than 1% are pooled into “Other” category. (B) Relative abundance at the level of genus. Observations less than 0.03% are pooled into “Other” category. (C) Shannon diversity index comparison using mouse 2 as a reference group. “****” means significance p<0.0001 while “ns” means non-significant.

**Individual Strain Comparisons (Level 2)**

*Enterococcus* significantly increased between mice cultures and the microbiome (Fig 6A).

Additionally, Proteobacteria strains were more abundant in cultures reaching a high pH; although, cultures reaching pH 6 were equivalent in Proteobacteria compared to cultures reaching pH 9 (Fig 6B). *Enterococcus* strains also followed a similar dynamic in which they increased in cultures reaching a lower pH, 6 and 7 (Fig 6C). Subsequently, cultures with the lowest pH, 6, had a significantly high abundance of *Lactobacillus* (Fig 6D). Archaea were more abundant in plates with pH 9 and 10 (Fig 6E), and further, *Clostridium* strains were more likely to be present in mouse 1 and 2 cultures compared to mouse 3 and 4 (Fig 6F).

**Fig 6. Comparison barcharts of individual microbial taxa.** (A) Comparison of the mean abundance of *Enterococcus* in cultured plates compared to the microbiome. (B) Comparison of the mean abundance of Proteobacteria in samples with varying pH. (C) Comparison of mean
abundance of *Enterococcus* in samples with varying pH. (D) Comparison of mean abundance of *Lactobacillus* in samples with varying pH. (E) Comparison in mean abundance of archaea in samples with varying pH. (F) Comparison in mean abundance of *Clostridium* in cultures partitioned by which mouse, M, it originated. “ns”, non significant; “***”, p < 0.001; “**”, p < 0.01; “*”, p ≤ 0.05.

**Microbial network (Level 2)**

The OTUs in the microbial network represent 88% of the relative sequence count for the cultured well plates (Fig 7). Much of the interactions were positive in nature meaning copresence in a shared-niche is the most abundant interaction type. Negative, mutually exclusive interactions are only between OTUs from the genus *Enterococcus* and several OTUs from the order Bacteroidales (Fig 7). The interaction between the 4 mutually exclusive OTUs account for 50% of all sequences.

**Fig 7. Microbial network generated using Spearman’s rank correlation at the taxonomic level of genus.** Most of the edges, 54 of 58, are of a positive correlation. The rest are negative. Nodes sizes are configured based on abundance. Nodes greater than or equal 10% are the largest, intermediate sized nodes are greater than or equal to 1%, and the smallest nodes are less than 1%.

**Discussion (Level 1)**

Many studies of the microbiome utilize germ-free mice, which are expensive to house and breed and require sacrificing animals to study the GI microbiome [14,16]. In this study, we attempted to culture an *ex vivo* microbiome in 3D well plates to decrease the cost associated with studying animal microbiomes. We found that cultures for mouse 1 and 2 were comparable to the gut microbiome in Shannon diversity (Fig 2C), which is very promising. Ultimately, our explant microbiome was significantly different than the *in vivo* microbiome, but we were able to culture a diverse number of prokaryotic strains utilizing our method. Optimizing efforts in culture media, detection, and atmospheric gradients is extremely important in culturing desired microbes.
With very little optimization, we were able to culture many gut microbial species including difficult to culture strains such as Methanobacteria [45], mean of 0.57 (SD 4.31), and MCG [46], mean of 218.59 (SD 633.71), which included the fecal B10 strain [47], (Fig 6E).

The upsurge of *Enterococcus* in cultured plates (Fig 6A) is explained by its competitiveness outlined in Fig 7. A likely scenario is that *Enterococcus* outcompeted strains within the order Bacteroidales, which makes up a high proportion of the gut microbiome [48,49]. *Enterococcus* is a facultative anaerobe [50] that may respire aerobically in the presence of hemin using cytochrome *bd* terminal oxidase, which reduces oxygen into water [51,52]. Since it is a known pioneer colonizer of the GI tract, its presence possibly established and maintained the anoxic environment by metabolically depleting the atmospheric oxygen, which allowed obligate anaerobes to thrive [53,54].

Post-incubation, a shift in pH was seen amongst the various 12 wells of the plates (Table 1). This shift in pH is accounted for by the increase in gram negative Proteobacteria in plates with a more basic pH (Fig 6B). Creation of the amine groups from this phylum perhaps led to the increase in pH [55–57]. Further, the decrease in pH may be related to the increase in the lactic acid fermenter *Enterococcus* in plates with a pH of 7 (Fig 6C) [52] while plates with a pH of 6 may be partially explained by the increase in both *Enterococcus* and *Lactobacillus* (Fig 6C, D) [52,58,59]. Additionally, pH was a strong influencer in the growth of archaea. Archaea grew more readily in plates with a higher pH (Fig 6E). Not only are archaea difficult to culture, but also their diversity is not well studied in regard to the gut microbiome [60]. Results are comparable to those of Ilhan et al. wherein pH had a strong influence on microbial composition in fecal anaerobic cultures [59]. There was a possible interaction between differences in the microbial communities seeding the culture plates and physio-chemical culture conditions causing
the pH in plates from different mice to swing in opposite directions. Additionally, culturing
plates in an anaerobic chamber instead of a CO₂ incubator may have exacerbated pH instability
[61]. Future efforts to stabilize pH may allow for additional growth of archaea and provide a
means to temporarily culture and study members that have in the past been recalcitrant to culture
methods.

The microbiome is passed on from mother to litter [8,62]. Our ex vivo microbiome was highly
impacted by familial relationship (Table 6, Fig 4). Further, mice differed in the amount of
Clostridium cultured. Mouse 1 and 2 had higher numbers of Clostridium than mouse 3 and 4 (Fig
7F). Not only were these mice siblings but also weaned by different mothers. The effects of
weaning are similar to Bian et al. wherein the abundance of an unclassified strain of
Clostridiaceae was affected by the nursing mother [63]. Our results reiterate the impact of the
mother on the microbiome, but also show this dynamic transfers even when explanted from the
source.

Essentially, our results indicate that the feces and large distal colon are highly similar; therefore,
it is reasonable to consider avoiding mouse sacrifice by culturing feces. Future experiments will
need to control pH shifts to avoid media-related population dynamics. Since none of the plates
maintained the original baseline pH or even homogenate pH, we assume that additional buffering
capacity or equilibrating media and culturing in a CO₂ incubator may create a more stable
explanted microbiome, possibly maintaining diversity more similar to in vivo microbiota. Even
with the pH swings seen here, we were able to culture bacteria that are difficult to routinely
culture. Ultimately, this study found that pH was a stronger influencer of community
composition than oxygen. pH has a strong influence on the establishment of the microbes that
will populate the explant culture. Future efforts at establishing an ex vivo mouse microbiome
should include additional measures geared towards stabilizing pH in order to avoid community
shifts related to physical changes in growth media.

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Barcharts of individual microbes
Plate microbiome inspection
Impact of environmental variables