



# Cecal Microbiota of Free-Range Hens Varied With Different Rearing Enrichments and Ranging Patterns

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Free-range pullets are reared indoors but the adult hens can go outside which is a mismatch that may reduce adaptation in the laying environment. Rearing enrichments might enhance pullet development and adaptations to subsequent free-range housing with impact on behavior and health measures including gut microbiota. Adult free-range hens vary in range use which may also be associated with microbiota composition. A total of 1,700 Hy-Line Brown® chicks were reared indoors across 16 weeks with three enrichment treatment groups: “control” with standard litter housing, “novelty” with weekly changed novel objects, and “structural” with custom-designed perching structures in the pens. At 15 weeks, 45 pullet cecal contents were sampled before moving 1,386 pullets to the free-range housing system. At 25 weeks, range access commenced, and movements were tracked via radio-frequency identification technology. At 65 weeks, 91 hens were selected based on range use patterns (“indoor”: no ranging; “high outdoor”: daily ranging) across all rearing enrichment groups and cecal contents were collected for microbiota analysis via 16S rRNA amplicon sequencing at V3-V4 regions. The most common bacteria in pullets were unclassified Barnesiellaceae, *Prevotella*, *Blautia* and *Clostridium* and in hens Unclassified, *Ruminococcus*, unclassified Lachnospiraceae, unclassified Bacteroidales, unclassified Paraprevotellaceae YRC22, and *Blautia*. The microbial alpha diversity was not significant within the enrichment/ranging groups (pullets:  $P \geq 0.17$ , hen rearing enrichment groups:  $P \geq 0.06$ , hen ranging groups:  $P \geq 0.54$ ), but beta diversity significantly varied between these groups (pullets:  $P \leq 0.002$ , hen rearing enrichment groups:  $P \leq 0.001$ , hen ranging groups:  $P \leq 0.008$ ). Among the short-chain fatty acids (SCFAs), the propionic acid content was higher ( $P = 0.03$ ) in the novelty group of pullets than the control group. There were no other significant differences in the SCFA contents between the rearing enrichment groups (all  $P \geq 0.10$ ), and the ranging groups (all  $P \geq 0.17$ ). Most of the genera identified were more abundant in the indoor than high outdoor hens. Overall, rearing enrichments affected the cecal microbiota diversity of both pullets and adult hens and was able to distinguish hens that remained inside compared with hens that ranging daily for several hours.

**Keywords:** bacteria, gut, indoor, outdoor, 16S rRNA, novel objects, perching structures, RFID

## INTRODUCTION

Numerous species of bacteria are sheltered in the gastro-intestinal (GI) tract of chickens, known as gut microbiota (Rychlik, 2020; Madlala et al., 2021), which play an important role in physiological processes including digestion, absorption, health, and production (Pourabedin and Zhao, 2015). Additionally, gut microbiota provide health benefits through inhibiting chronic diseases (Bavananthasivam et al., 2021), defending the host from various pathogens (Waite and Taylor, 2014), and enhancing gut health (Diaz Carrasco et al., 2019). Gut microbiota also contribute to pathogen expulsion from the host and development of the host immune system (Yeoman et al., 2012; Pan and Yu, 2014). Within chickens, distinct microbial communities are present across different areas of the GI tract with the complexity and absolute counts being comparatively high in the cecum (Rychlik, 2020). Fu et al. (2018) explored the gut microbiota diversity of laying hens based on 16S rRNA sequencing in China and found the Bacteroidetes, then Firmicutes phyla were the most predominant among the cecal microbes. Actinobacteria and Proteobacteria are also typically represented in the ceca of most adult chickens although there can be high variation in microbial composition between individuals (Rychlik, 2020). The most common genera of microbiota in the ceca of laying hens have been shown to be the *Bacteroides* and *Prevotella* (Yan et al., 2017). Identifying the composition of microbes within chickens is important for understanding their function and how they may be affected by internal and external environmental parameters.

Research to date in chickens (both meat birds and egg laying hens) has shown that gut microbiota varied across different ages and production stages of both broilers (Han et al., 2016; Jurburg et al., 2019) and laying hens (Videnska et al., 2014; Ngunjiri et al., 2019; Joat et al., 2021). Microbiota populations, their diversity, and composition varied in hens with different diet compositions (Borrelli et al., 2017; Zhou et al., 2021). There is also some evidence that microbiota composition can vary among groups of hens selected for behavioral differences in feather pecking (van der Eijk et al., 2019), although other studies have not found significant differences (Borda-Molina et al., 2021). Housing environments can affect microbial composition with differences found between broiler chickens that had fresh or reused litter on the floor (Wang et al., 2016), or different types of litter substrates (Torok et al., 2009). Several studies comparing different types of indoor cage and cage-free systems (Nordentoft et al., 2011; Adhikari et al., 2020) or caged indoor and outdoor systems in laying hens (Cui et al., 2017; Chen et al., 2019; Hubert et al., 2019; Schreuder et al., 2020, 2021; Seidlerova et al., 2020) have highlighted the impact that variation in the housing environment might have on the gut microbiota diversity and populations.

To date, some studies on housing system impacts in laying hens have made comparisons between hens that are housed inside (caged or cage-free systems), or housed cage-free with access to an outdoor area in both experimental (Chen et al., 2019; Hubert et al., 2019) and commercial settings (Cui et al., 2017; Schreuder et al., 2020, 2021; Seidlerova et al., 2020). Results

across these studies have shown that hens kept in backyard flocks outside or given outdoor range access had lower diversity of bacterial species than indoor-housed hens (cloacal microbiota: (Schreuder et al., 2020); cecal microbiota: Seidlerova et al. (2020). Alternatively, other studies have found greater intestinal/cecal microbial diversity in the hens with outdoor access (Cui et al., 2017; Chen et al., 2019; Hubert et al., 2019), and Schreuder et al. (2021) found no substantial differences in cloacal microbiota between hens with indoor only or outdoor access. Adult hens within a free-range system have access to an outdoor area, but not all hens will make use of this resource with some hens ranging infrequently or not at all while others range daily for several hours (Campbell et al., 2020a). Previous research has determined this variation in ranging patterns correlates with variation in fear behavior and other welfare measures (Campbell et al., 2020a). Thus, individual variation in gut microbiota may also be expected. To date, research investigating this is limited (Ruhnke et al., 2018).

Free-range egg production systems in Australia have been increasing to meet consumer preferences for both perceived better-quality eggs and perceived benefits to hen welfare (Bray and Ankeny, 2017; Scrinis et al., 2017). However, free-range housing can have both positive and negative impacts on hen welfare (Campbell et al., 2020a) which may in part be dependent on the discrepancies between rearing and adult housing (Janczak and Riber, 2015). In Australia and elsewhere internationally, free-range pullets are reared inside, but the adult hens have outdoor access. As outdoor access for pullets is challenging due to vaccination requirements and rearing shed construction, enriching the rearing environment instead may improve adaptation of the adult hens. These differences in rearing environments can affect bird behavioral, physical, and physiological development (Campbell et al., 2019) and may also affect the microbiota composition and diversity of the pullets as they grow.

The data presented in this study were from birds that were part of a larger study on the effects of rearing enrichments and ranging variation on the behavior, health, welfare, and production of free-range laying hens (Bari et al., 2020a,b,c, 2021; Campbell et al., 2020c,b). The results across the wider project showed that rearing enrichments had few effects on the behavior and welfare of pullets (Campbell et al., 2020c), but there were impacts of rearing enrichments across the flock cycle in the adult birds. For example, the enriched-reared hens had better plumage coverage (Bari et al., 2020b,c), increased ranging (Campbell et al., 2020b), and hens that ranged for longer had lower body weight but better plumage (Bari et al., 2020c). Thus, if enrichment in rearing had multiple effects on the development of the pullets, including long-term impacts, enrichment might also affect the microbiota of both pullets and adult hens. Individual variation in ranging behavior has previously been correlated with other physical and behavioral differences and, thus, may also be correlated with differences in microbiota composition in adult hens. This study was conducted to evaluate the gut microbiota composition and diversity of the pullets from different rearing environments, and the microbiota composition and diversity of hens from

different rearing enrichment groups and ranging patterns. This study represents a first step toward understanding the impacts that rearing environments and individual variation in ranging can have on internal bacterial communities. Short-chain fatty acids were also assessed to support any differences in microbiota although the functional link between microbiota and bird health, welfare, or production were not assessed. A more diverse microbiota population was predicted in the enriched pullets and adult hens than the control groups, but the direction of differences in cecal microbiota diversity between the indoor hens and outdoor rangers was uncertain.

## MATERIALS AND METHODS

### Ethical Approval

The University of New England Animal Ethics Committee approved all experimental procedures (AEC17-092).

### Pullet Housing

A flock of 1,700 commercially supplied Hy-Line® Brown chicks were reared indoors at the Rob Cumming Poultry Innovation Centre at the University of New England, Armidale, NSW, Australia. The day-old chicks were reared across 16 weeks within 9 pens (6.2 m<sup>2</sup> × 3.2 mW) distributed across three separate rooms with 3 separate rearing enrichment treatments as previously described (Bari et al., 2020a,b,c). The rearing treatments included a control group with rice hulls as floor litter only, a novelty group where novel objects were also added and changed at weekly intervals (e.g., balls, bottles, bricks, brooms, brushes, buckets, containers, pet toys, plastic pipes etc.) and a structural group where five custom-designed H-shaped perching structures (L, W, H = 0.60 m) with two solid panels and one open-framed side were placed for the whole period of rearing. The pullets were isolated visually but not acoustically from the pullets of other pens via shade cloth on the wire pen dividers. At 16 weeks of age, bird density was approximately 15 kg/m<sup>2</sup> or 9 pullets/m<sup>2</sup> (average 174–190 pullets/pen).

The pullets were provided with *ad libitum* access to feed (commercially formulated mash) and water using round feeders and nipple drinker lines. The resources provided met or exceeded the current Australian Model Code of Practice for the Welfare of Animals Domestic Poultry (Primary Industries Standing Committee, 2002). Artificial lighting and temperature schedules followed the recommended Hy-Line® Brown alternative management guidelines (Hy-Line, 2016) but the LED lighting was maintained at 100 lux as the pullets were destined for outdoor access (no natural light was present during rearing). Mechanical ventilation and heating systems were used as needed. Chicks were beak-trimmed using infra-red at the hatchery with a vaccination schedule throughout rearing as per regulatory requirements and standard recommendations including vaccination against Newcastle disease, Marek's disease, fowl pox, fowl cholera, egg drop syndrome, *Mycoplasma gallisepticum*, *Mycoplasma synoviae*, infectious bronchitis, infectious laryngotracheitis, and avian encephalomyelitis.

### Pullet Sampling

At 15 weeks of age, 45 pullets ( $n = 15$  per treatment group) were selected from a random sample of 90 pullets [dissected in Campbell et al. (2020c)] based on balanced live weights across pen replicates. The live weights between the selected pullets varied from 1,250 to 1,370 g (mean  $\pm$  SE =  $1312 \pm 4.68$  g). The selected hens were killed with CO<sub>2</sub> and then opened when the movements of the birds had completely ceased. Across a single sampling day, the contents from both ceca were collected, mixed well, transferred to Eppendorf tubes and placed in liquid nitrogen until storage at  $-80^{\circ}\text{C}$  at the conclusion of the sampling day.

### Adult Housing

A total of 1,386 pullets were transferred to the free-range facility at the Laureldale farm of the University of New England, Armidale, NSW, Australia at 16 weeks of age and remixed within pen replicates. The hens were housed within the three rearing treatments across 9 adjacent pens (4.8 m<sup>2</sup> × 3.6 mW) located in a single shed with an indoor density of approximately 9 hens/m<sup>2</sup> (154 hens/pen). Shade cloth on wire pen dividers was used to visually separate the hens from other pens. The indoor pens contained nest boxes (2 small and 1 large tiered nest box), perches, round hanging feeders and water nipples to fulfill the requirements of the Australian Model Code of Practice for the Welfare of Animals: Domestic Poultry (Primary Industries Standing Committee, 2002). Rice hulls were used as floor litter material with one complete litter replacement mid-way through the flock cycle. The LED lighting gradually increased to 16 h light and 8 h dark by 30 weeks of age with an average pen intensity of 10.0 (0.84 SE) lux (Lutron Light Meter, LX-112850; Lutron Electronic Enterprise CO., Ltd., Taipei, Taiwan) as measured at birds' eye height from 3 pen locations (front, middle, back) when the pop-holes were closed. The shed was mechanically ventilated with no automated cooling system.

Each of the 9 pens was connected to an outdoor range area (31 m<sup>2</sup> × 3.6 mW for each pen with a density of approximately 1.4 hens/m<sup>2</sup>). The range was accessed via two pop-hole openings (18 cm W × 36 cm H) per pen. The range area immediately after the pop-holes was 1.1 m length of concrete path, then 1.6 m length of river rock followed by a grassed area with no additional trees or shelter. Each range was visually divided by shade cloth hung along the wire fences. Hens were provided access to the outdoor area from 25 weeks of age (May 2018) for most of the daytime via automatic opening and closing of the pop-holes. The pop-holes opened at 9:15 am and closed after sunset daily (approximately 9 h of ranging time across winter and 11 h ranging after daylight saving time started in October 2018).

### Radio-Frequency Identification of Ranging

All the hens were banded with microchips (Trovan® Unique ID 100 (FDX-A): operating frequency 128 kHz, Microchips Australia Pty Ltd., Keysborough, VIC, Australia) glued into adjustable leg bands (Roxan Developments Ltd., Selkirk, Scotland) to track their movement in and out of the range pop-holes until 64 weeks via radio-frequency identification (RFID) systems.

The RFID systems were designed and supported by Microchips Australia Pty Ltd. with equipment developed and manufactured by Dorset Identification B.V. (Aalten, the Netherlands) using Trovan<sup>®</sup> technology (RFID Systems Ltd., North Ferriby, United Kingdom). The date and time of each tagged bird passing through and in which direction (onto the range, or into the pen) were recorded with a precision of 0.024 s (maximum detection velocity 9.3 m/s). A custom-designed software program written in the “Delphi” language (Bryce Little, Agriculture and Food, CSIRO, St Lucia, QLD, Australia) filtered out unpaired or “false” readings from the RFID data from 56 until 64 weeks of age (54 days of data). The same program summarized the daily data to provide the mean number of hours outside per hen across the sampling period.

## Hen Selection and Sample Collection

At 65 weeks of age,  $n = 91$  hens were selected across all pens from all rearing treatments with specific ranging patterns as identified from the RFID data. The hens were categorized as “high outdoor” which were hens that accessed the outdoor range on all the selected days for 5 h 12 min to 9 h on average daily, and “indoor” which were hens that accessed the range on one or zero of the 54 days. Of the selected 91 hens, 44 were indoor hens (control  $n = 14$ , novelty  $n = 14$ , structural,  $n = 16$ ), and 47 were high outdoor hens (control  $n = 15$ , novelty  $n = 17$ , structural,  $n = 15$ ) totaling  $n = 29$  control,  $n = 31$  novelty, and  $n = 31$  structural hens. Hen selection was balanced across all treatments as best as possible, but some pens within treatments had higher numbers of hens with extreme ranging patterns. The live weight of the selected hens ranged from 1,800 to 2,110 g (mean  $\pm$  SE =  $1966.81 \pm 7.39$  g). The body weight was not completely balanced as the hens were primarily selected based on their ranging patterns and different rearing enrichment groups; closer body weight between them was prioritized where possible during selection. The same hens were also reported on in Bari et al. (2020c) as they, along with a further 216 hens had post-mortem assessments conducted including internal organ weights, presence of disease/infections, and carcass composition.

At 65 weeks, the selected hens were transported in carrier crates from the free-range facility to a post-mortem facility at the University of New England (~5.5 km distance) just before (~2 h) the post-mortem. The hens were killed with CO<sub>2</sub>. Immediately after the cessation of all movements, the hens were opened, and the cecal contents were collected and mixed together from both ceca with approximately 200 mg of contents transferred to an Eppendorf tube (2 ml size). These samples were placed on ice immediately after sampling, then transferred to a  $-20^{\circ}\text{C}$  freezer half-way through the day. At the end of the sampling day all samples were stored in a  $-80^{\circ}\text{C}$  freezer until laboratory analysis.

## DNA Extraction

The DNA of pooled cecal content of both ceca collected at 65 weeks of age was extracted using DNeasy 96 PowerSoil Pro QIAcube HT Kit, (Qiagen, Inc., Doncaster, VIC, Australia) with slight modification. Approximately 65 mg of frozen cecal contents were weighed in a 2 ml Eppendorf tube containing 300 mg of glass beads. Then, 800  $\mu\text{l}$  of solution CD1 was

added and vortexed for 5 s. The tubes were then placed in the TissueLyser II for 5 min at a frequency of 30 Hz to disrupt bacterial cells. The tubes were spun briefly and placed in the heat block at  $90^{\circ}\text{C}$  for 10 min. The tubes were vortexed for 5 s followed by centrifuging at  $15,000 \times g$  for 1 min. Then, the supernatants (approximately 500–650  $\mu\text{l}$ ) were transferred to a new 2 ml tube. Later, 250  $\mu\text{l}$  of CD2 were added and the tubes were inverted 3 times followed by centrifuging again at  $4,500 \times g$  for 5 min. The supernatants were transferred to a new S-Block. The S-Block was placed in the correct position in QIAcube HT. The reactions were loaded into the dedicated cassette with written names. Then the extraction was performed using the QIAcube HT following the manufacturer’s instruction. The quality and quantity of extracted DNA was determined using a NanoDrop spectrophotometer (Nanodrop 8000; Thermo Scientific, Wilmington, DE, United States). The ratios A260/A280 being  $> 1.8$  were considered as of high quality and the extracted DNA were kept at  $-20^{\circ}\text{C}$  until required.

## 16S rRNA Gene Sequencing

The V3–V4 region of 16S rRNA genes were amplified using forward primer 16S\_341f (TCGTCGGCA GCGTCAGATGTGTATAAGAGACAGCCTACGGGNGGCWGCAG) and reverse primer 16S-805r (GTCTCGTGGGCTCGGAGATGTGTATAAGAGACAGGACTACHVGGGTATCTAA TCC) (Klindworth et al., 2013). The sequencing was performed on an Illumina MiSeq system ( $2 \times 300$  bp) at the Ramaciotti Centre for Genomics (UNSW, Sydney, NSW, Australia).

## Short-Chain Fatty Acids Analysis

The cecal short-chain fatty acids (SCFAs) were analyzed according to the method described by Bach Knudsen et al. (1991) and Richardson et al. (1989) with slight modifications. Briefly, frozen cecal samples were defrosted and homogenized keeping them at  $4^{\circ}\text{C}$  overnight. Approximately 0.8 g of the homogenized cecal sample was weighed into a centrifuge tube and 1 mL of internal standard solution (0.01M ethylbutyric acid) was added. The sample with solution was vortexed for 1 min and centrifuged at 15,000 rpm for 20 min at  $5^{\circ}\text{C}$ , with approximately 1 mL supernatant transferred into an 8 mL vial. Using the same method, a blank and an internal standard solution (0.1 mL of 0.1M ethyl butyric acid) were also prepared into 8 mL vials by replacing 1 mL of the supernatant with the same amount of water and standard acid mixture, respectively. Then, 2.5 mL of diethylether and 0.5 mL of concentrated HCl (36%) were added to 8 mL vials containing the supernatant, blank and standard solution, and thoroughly mixed by using a vortex mixer. The mixture was vortexed, and centrifuged at 3,000 rpm at  $5^{\circ}\text{C}$  for 15 min. An aliquot of 400  $\mu\text{L}$  of the supernatant was transferred to a gas chromatograph vial (2 mL) and mixed with 40  $\mu\text{L}$  of *N*-tert-butyltrimethylsilyl-*N*-methyltrifluoroacetamide (MTBSTFA) and incubated at  $80^{\circ}\text{C}$  for 20 min. The GC vials were tightened appropriately and left at room temperature for at least 48 h. The cecal SCFAs were measured using a Varian CP3400 CX gas Chromatograph (Varian Analytical Instruments, Palo Alto, CA, United States) after adding approximately 0.5 mL

ether into each GC vial. The concentrations of cecal SCFAs were expressed as  $\mu\text{mol/g}$  cecal samples.

## Data and Statistical Analyses

A total of 45 cecal content samples of pullets at 15 weeks of age from different rearing enrichments including control, novelty and structural groups were analyzed. A total of 91 cecal content samples of free-range hens at 65 weeks of age from different rearing enrichments including control, novelty and structural, and different ranging patterns (indoor, high outdoor) were analyzed.

The sequence reads' quality was checked using fastQC v0.11.9 (Babraham Institute, Cambridge, United Kingdom) (Andrews et al., 2014). Quantitative Insights Into Microbial Ecology (QIIME2) (Bolyen et al., 2019) was used for upstream analysis of the sequence employing DADA2 plugin (Callahan et al., 2016) for error correction, filtering, merging pair ends and removing chimeras. The taxonomy was assigned against the Greengenes database (v13\_8) using a Naïve Bayes classifier trained against the primers used in this study. The amplicon sequence variants (ASV) table obtained from QIIME2 was then subjected to downstream analysis and visualization with Calypso (Zakrzewski et al., 2017).

The ASV data were normalized by total sum normalization of square-root transformed data in Calypso and the rare taxa were not excluded. The ordination of the groups was done with redundancy analysis (RDA) while a non-parametric multivariate test ANOSIM (Bray-Curtis) was used to test the differences between the groups. Shannon Index, Richness, Chao1, and Simpson's Index were calculated to analyze alpha diversity among the groups. Linear discriminant analysis Effect Size (LEfSe) was used to identify the differentially abundant representative taxa of each group while differences in individual taxon among different groups was also tested with ANOVA followed by *post hoc* comparisons where significant differences were present. All *p*-values were adjusted for multiple testing, which is a feature of Calypso (Zakrzewski et al., 2017).

For SCFA analysis, a total of 90 samples were used, of which 30 were from pullets (control  $n = 10$ , novelty  $n = 10$ , and structural  $n = 10$ ), and 60 were from adult hens [control hens:  $n = 20$  (9 indoor, 11 outdoor); novelty hens:  $n = 20$  (8 indoor, 12 outdoor); structural hens:  $n = 20$  (12 indoor, 8 outdoor)]. A general linear mixed model (GLMM) was applied to the pullet cecal SCFA data with rearing enrichments as a fixed effect and Bird ID nested within Pen nested within rearing enrichments as a random effect. For the adult hens, cecal SCFA data, a GLMM was applied with rearing enrichments and ranging patterns as fixed effects and Bird ID nested within Pen and Pen nested within rearing enrichments and ranging patterns as random effects. The interaction term for the adult data was not included to better match the microbiota analyses from these birds. Data were  $\text{Log}_{10}$  transformed for formic acid and lactic acid content in the pullet SCFAs. For the adult hen SCFAs, the formic acid values were  $\text{Log}_{10}$  transformed and the lactic acid values were square-root transformed. Where significant differences were present, *post hoc* Student's *t*-tests were applied to the least squares means. All these statistical analyses were

conducted in JMP 14.0® (SAS Institute, Cary, NC, United States) with  $\alpha$  set at 0.05.

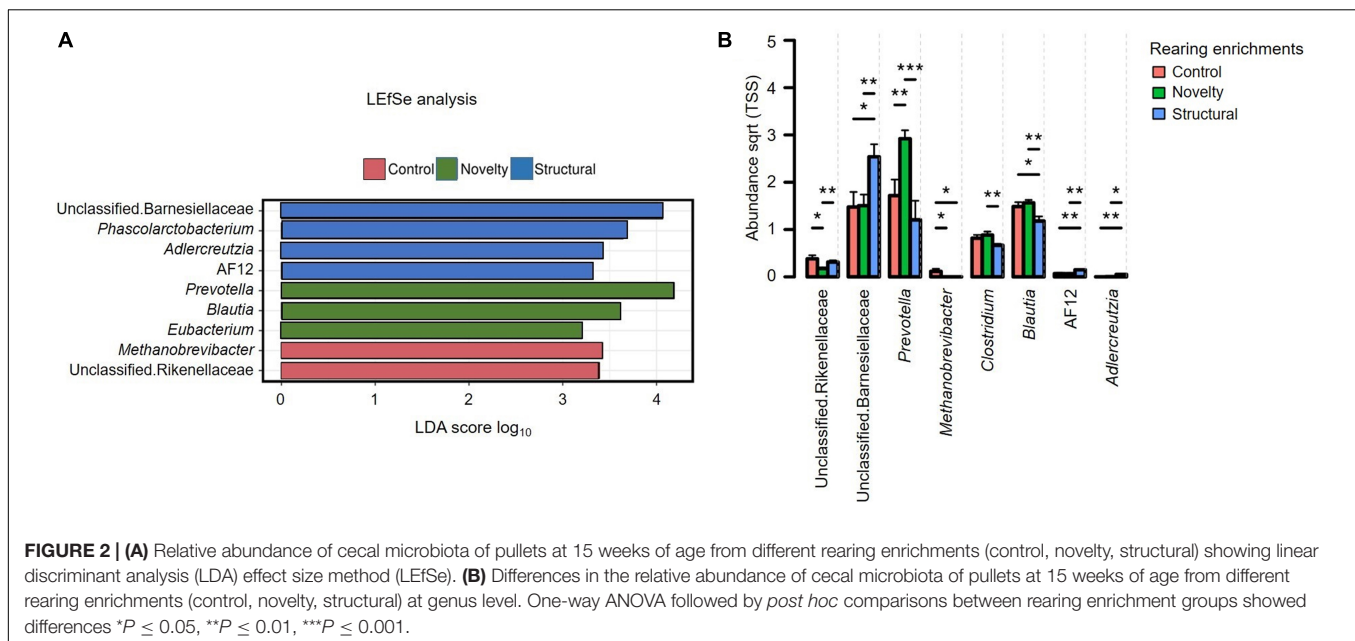
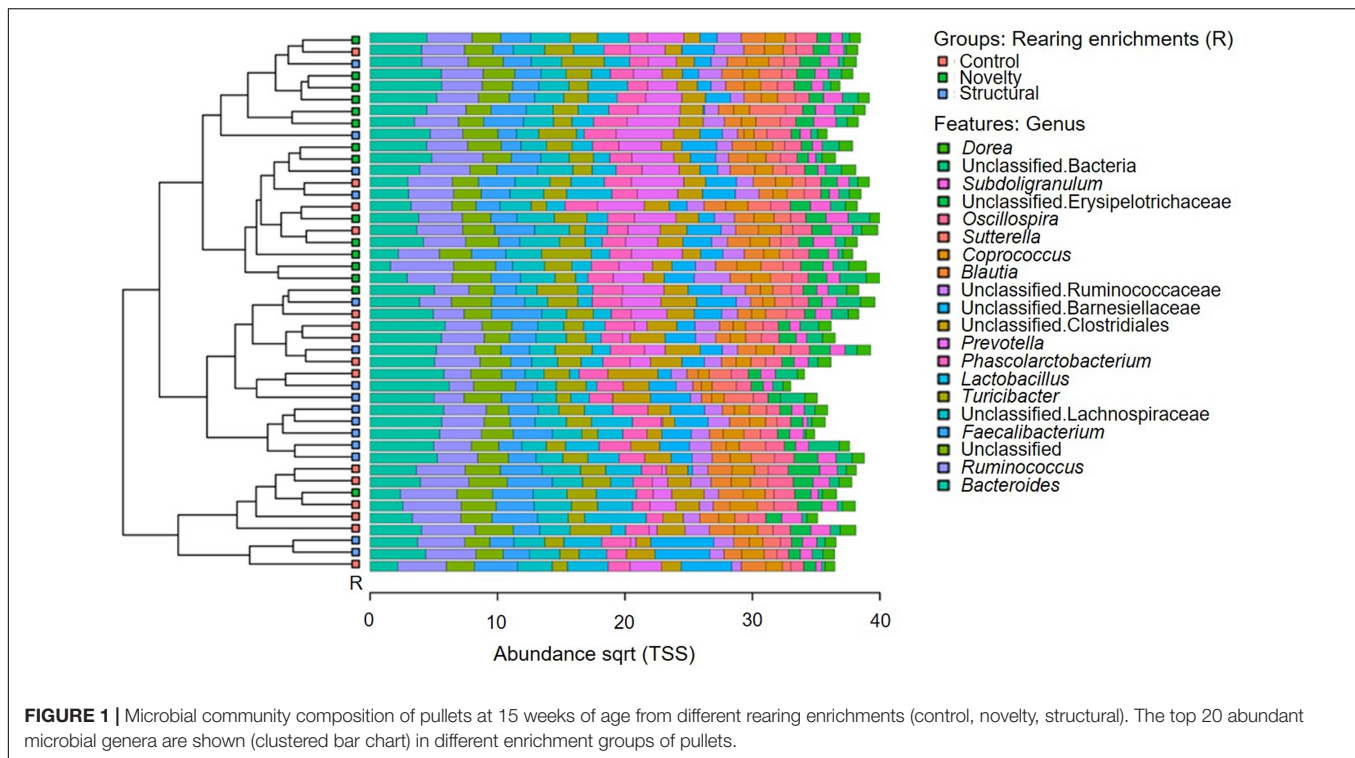
## RESULTS

### Cecal Microbiota of Pullets

The microbial community composition of the pullets is shown in **Figure 1**. The predominant microbiota genera included *Dorea*, unclassified bacteria, *Subdoligranulum*, unclassified Erysipelotrichaceae, *Oscillospira*, *Sutterella*, *Coproccoccus*, *Blautia*, unclassified Ruminococcaceae, unclassified Barnesiellaceae, unclassified Clostridiales, *Prevotella*, *Phascolarctobacterium*, *Lactobacillus*, *Turicibacter*, unclassified Lachnospiraceae *Faecalibacterium*, Unclassified, *Ruminococcus*, and *Bacteroides*.

The sequence reads per sample revealed no significant differences between the treatment groups ( $P = 0.35$ ) with a minimum read of 19,840 and a maximum read of 64,330 sequences. The Linear Discriminant (LDA) Effect Size (LEfSe) analysis (**Figure 2A**) showed the differentially abundant explanatory microbiota genera in the structural group included unclassified Barnesiellaceae, *Phascolarctobacterium*, *Adlercreutzia*, and AF12 with *Prevotella*, *Blautia* and *Eubacterium* predominant in the novelty group, and *Methanobrevibacter* and unclassified Rikenellaceae explanatory in the control group. Explanatory taxa (genera in our case) explain the differences between the groups. **Figure 2B** displays the relative abundance of cecal microbiota of pullets at 15 weeks of age from different rearing enrichments at the genera level with all bacterial groups showing a significant effect of rearing enrichment treatment (all  $P < 0.05$ ). The *Methanobrevibacter* genus was more abundant in the control group than the other two groups ( $P \leq 0.05$ ), and *Prevotella* was more abundant in the novelty group than the control ( $P \leq 0.01$ ) and structural groups ( $P \leq 0.001$ ) of pullets. The *Clostridium* genus was more abundant in the novelty group than the structural pullets only ( $P \leq 0.01$ ). The *Blautia* was less abundant in the structural group than in both novelty and control pullets ( $P \leq 0.01$ ), but there was no difference in the *Blautia* abundance between the control and novelty pullets. The *Adlercreutzia* genus was more abundant in the structural group of pullets than the control ( $P \leq 0.01$ ) and novelty ( $P \leq 0.05$ ) groups of pullets. The unclassified Rikenellaceae was more abundant in the control group than the novelty pullets ( $P \leq 0.05$ ), and more abundant in the structural group than the novelty pullets ( $P \leq 0.01$ ). The unclassified Barnesiellaceae was more abundant in the structural group than the novelty ( $P \leq 0.05$ ) and control ( $P \leq 0.01$ ) pullets. The genus AF12 was more abundant in the structural group than the novelty ( $P \leq 0.01$ ) and control groups ( $P \leq 0.01$ ) of pullets.

Two multivariate analyses of beta diversity at ASV level, redundancy analysis (RDA) (**Figure 3E**) ( $P = 0.002$ ) and ANOSIM (Bray-Curtis) ( $R = 0.112$ ,  $P = 0.004$ ) showed significant differences between the groups of rearing enrichment treatments (**Figures 3E,F**). However, there was no difference in the alpha diversity indices at ASV level among the rearing enrichment groups as assessed by Shannon index ( $P = 0.17$ ), Richness



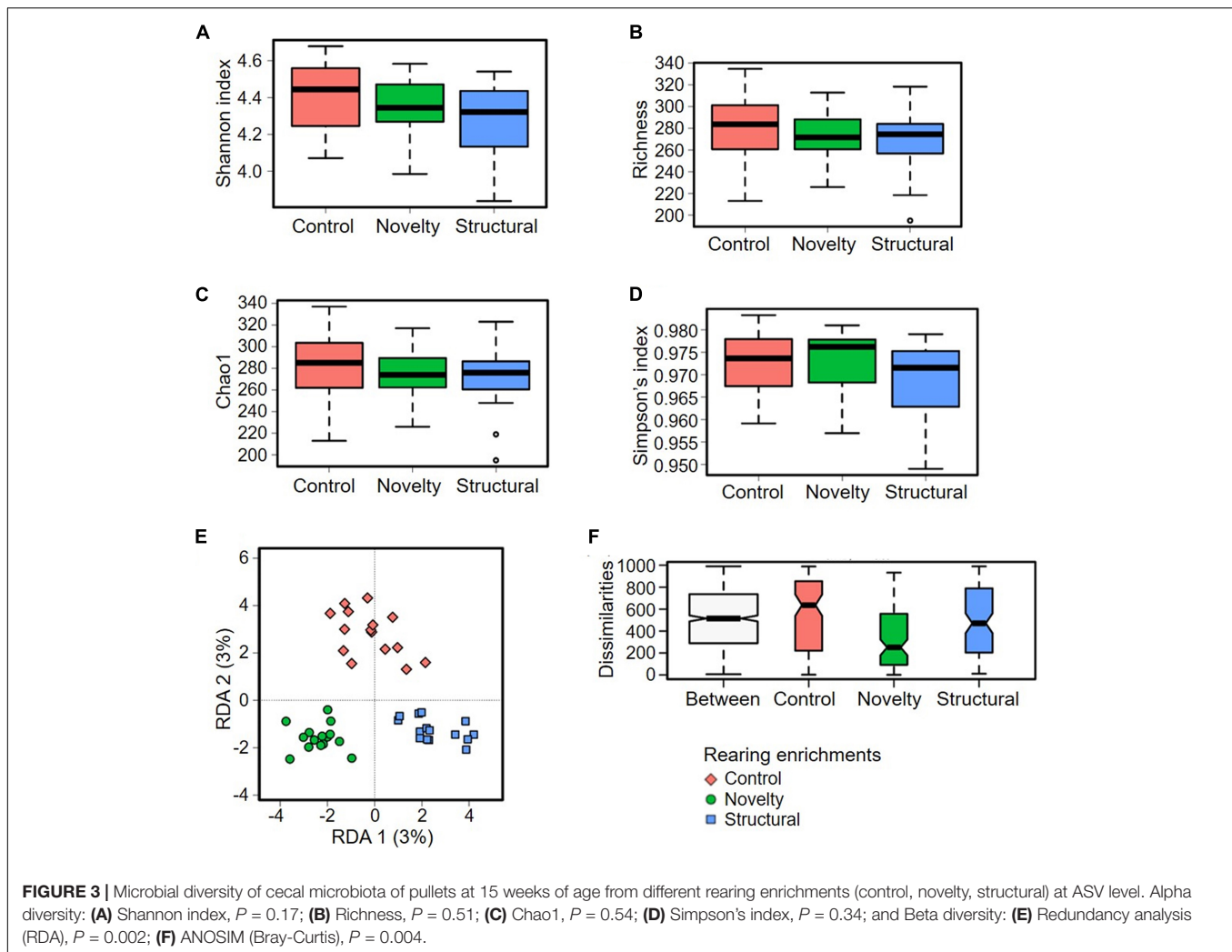
( $P = 0.51$ ), Chao1 ( $P = 0.54$ ) and Simpson's index ( $P = 0.34$ ) (Figures 3A–D).

## Cecal Microbiota of Free-Range Hens

The microbial community composition of the free-range hens from different rearing enrichments is shown in Figure 4; and from different ranging patterns in Figure 5. The predominant microbiota genera from the rearing enrichment groups and ranging patterns included *Bifidobacterium*, YRC22, *Oscillospira*,

unclassified Coriobacteriaceae, *Peptococcus*, *Megasphaera*, *Blautia*, unclassified S247, *Megamonas*, unclassified Paraprevotellaceae, unclassified Bacteroidales, *Turicibacter*, *Prevotella*, *Faecalibacterium*, *Phascolarctobacterium*, unclassified Lachnospiraceae, *Lactobacillus*, *Ruminococcus*, Unclassified, and *Bacteroides*.

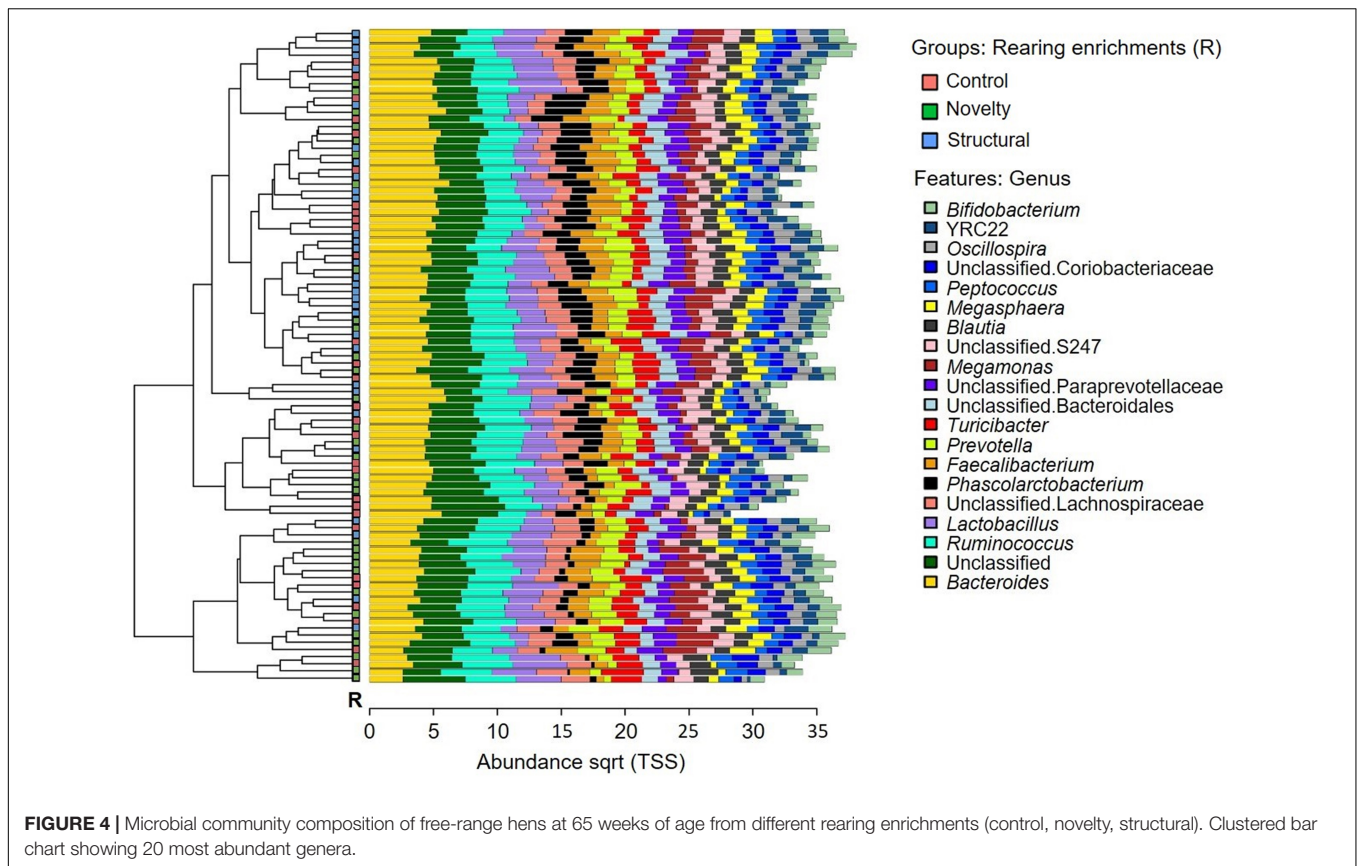
The sequence reads per sample revealed no significant differences between the rearing enrichment groups ( $P = 0.20$ ) with a minimum read of 14,055 and a maximum read of 52,056



sequences. The sequence reads per sample was also not significant between the ranging groups ( $P = 0.91$ ) with a minimum read of 14,055 and a maximum read of 52,056 sequences.

**Figure 6A** shows the LEfSe analysis results that indicated the differentially abundant explanatory microbiota genera from the rearing enrichment groups. LEfSe analysis showed *Mucispirillum*, *Sutterella*, *Helicobacter*, *Anaerobiospirillum*, unclassified Bacteria, and *Succinatimonas* were the characteristic genera from the structural group, *Ruminococcus*, *Blautia*, *Peptococcus*, *Dorea*, *Brachybacterium*, *Coprococcus*, *Jeotgalicoccus*, unclassified RF39, *Anaerofustis*, and unclassified Mollicutes from the novelty group, and Unclassified, unclassified Chlamydiaceae, *Butyricimonas*, and unclassified Barnesiellaceae from the control group of rearing treatments. **Figure 6B** shows the relative abundance of cecal microbiota of free-range hens at 65 weeks of age from different rearing enrichment treatments at the genera level where there were significant effects of rearing enrichment treatments across all bacterial groups (all  $P < 0.05$ ). The *Sutterella* genus was more abundant in the structural hens than in the novelty hens ( $P \leq 0.001$ ), but the control group did not differ. The *Succinatimonas* was more predominant in the

structural ( $P \leq 0.01$ ) and novelty groups ( $P \leq 0.05$ ) than the control hens. The *Sphaerochaeta* genus was more abundant in the control group than the novelty hens ( $P \leq 0.05$ ), but there was no difference between the novelty and structural hens. *Ruminococcus* was more abundant in the novelty hens ( $P \leq 0.01$ ) than the structural, but did not differ between control and structural hens. The *Peptococcus* genus was also more abundant in the novelty hens than the structural group ( $P \leq 0.01$ ). The *Mucispirillum* was more abundant in both the control and structural hens than the novelty group ( $P \leq 0.01$ ). The *Odoribacter* genus was more abundant in the control group than the novelty ( $P \leq 0.01$ ). The *Jeotgalicoccus* was more abundant in the novelty hens than the control hens ( $P \leq 0.01$ ), but did not differ between the novelty and structural hens. The *Corynebacterium* was more abundant in the novelty hens than both the control and structural hens ( $P \leq 0.05$ ). *Butyricimonas* was more abundant in the control hens than the novelty ( $P \leq 0.001$ ) and structural hens ( $P \leq 0.01$ ). The *Brachybacterium* and *Blautia* genera were more predominant in the novelty hens than the control ( $P \leq 0.01$ ) and structural hens ( $P \leq 0.05$ ), and the *Anaerofustis* was more predominant



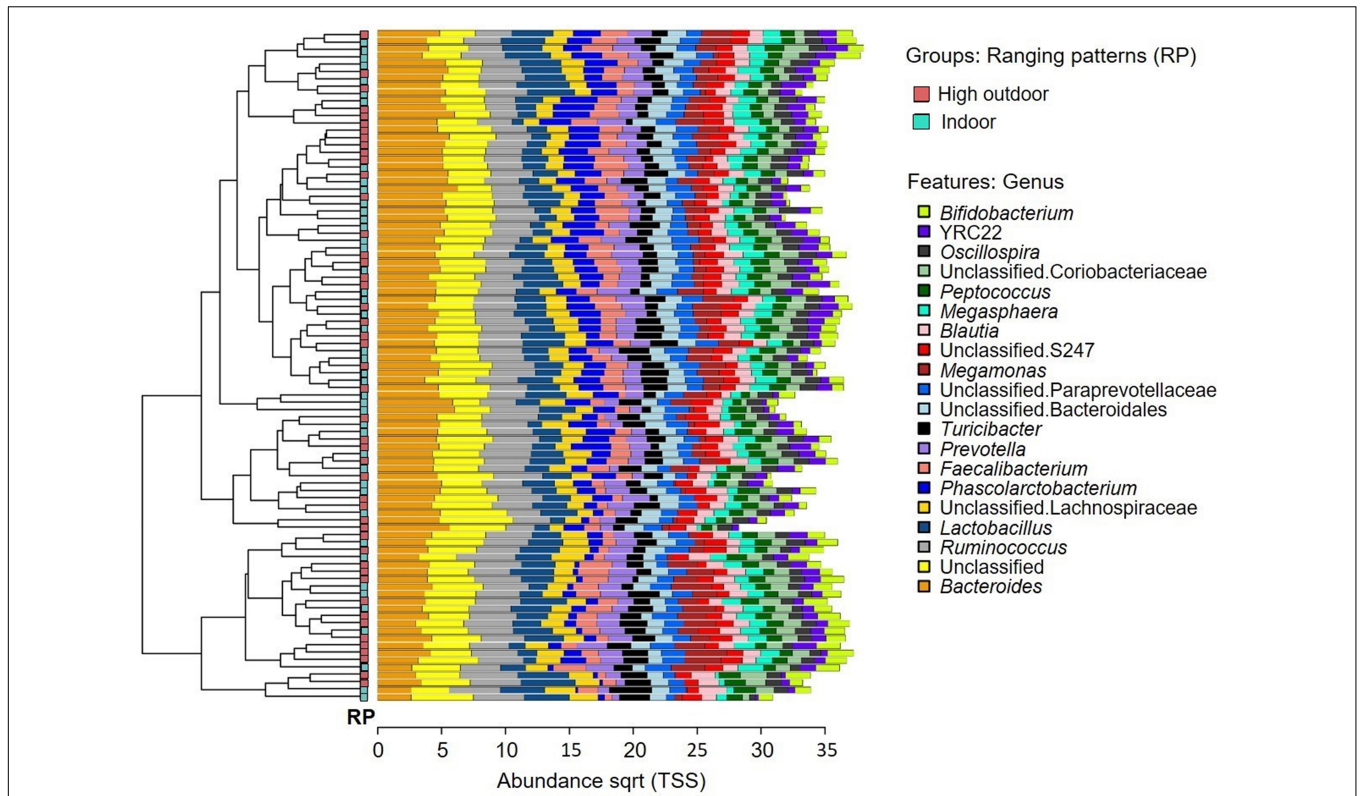
in the novelty hens than the control hens ( $P \leq 0.01$ ) but did not differ between the novelty and structural hens (**Figure 6B**). The unclassified RF39 had significantly lower abundance in the control hens than in both the novelty and structural groups ( $P \leq 0.05$ ), but there was no difference in the unclassified RF39 abundance between the novelty and structural hens. The control group had more abundant unclassified R441B bacteria than the control ( $P \leq 0.05$ ) hens. Overall, the significantly more abundant bacteria were YRC22 in structural, *Blautia*, *Brachyбактерium* and *Corynebacterium* in novelty and *Butyricimonas* in control hens; and the significantly less abundant bacteria were Unclassified in structural, *Succinatimonas* in control, and *Mucispirillum* and *Helicobacter* in novelty.

**Figure 7A** shows the LEfSe analysis results that indicated the explanatory microbiota genera in the hens from different ranging patterns. The indoor hens had *Streptococcus*, *Clostridium*, *Enterococcus*, *Staphylococcus*, unclassified Ruminococcaceae, *Brachyбактерium*, unclassified Desulfobivriaceae, unclassified Erysipelotrichaceae, *Veillonella*, *Aeriscardovia*, *Brevibacterium*, *Roseburia*, cc.115, *Pseudoramibacter Eubacterium*, *Dietzia*, unclassified Bacillaceae and *Yaniella* as the explanatory genera whereas in the high outdoor group RFN20, *Sphaerochaeta*, unclassified Elusimicrobiaceae, and *Rickettsiella* were the explanatory bacteria. **Figure 7B** shows that the indoor hens had *Yaniella* ( $P \leq 0.05$ ), *Veillonella* ( $P \leq 0.01$ ), *Streptococcus* ( $P \leq 0.01$ ), *Staphylococcus* ( $P \leq 0.05$ ), *Roseburia* ( $P \leq 0.05$ ), *Gallibacterium* ( $P \leq 0.05$ ), *Enterococcus* ( $P \leq 0.001$ ), *Dietzia*

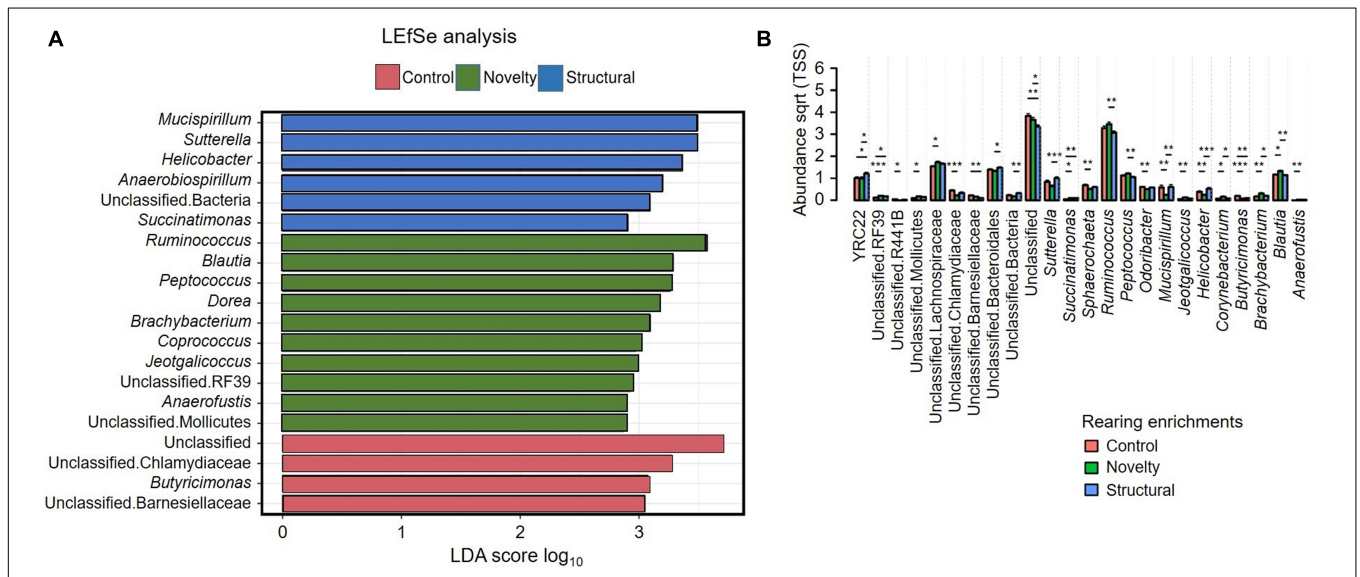
( $P \leq 0.05$ ), *Clostridium* ( $P \leq 0.001$ ), *Brevibacterium* ( $P \leq 0.05$ ), *Brachyspira* ( $P \leq 0.05$ ), *Brachyбактерium* ( $P \leq 0.01$ ), and *Aeriscardovia* ( $P \leq 0.05$ ) genera in greater abundance than the high outdoor hens, but the *Sphaerochaeta* were more predominant in the high outdoor hens than the indoor hens ( $P \leq 0.05$ ). Also, all of the unclassified genera including unclassified Ruminococcaceae ( $P \leq 0.01$ ), unclassified Erysipelotrichaceae ( $P \leq 0.01$ ), unclassified Desulfobivriaceae ( $P \leq 0.001$ ), unclassified Bacillaceae ( $P \leq 0.05$ ) and *Pseudoramibacter Eubacterium* ( $P \leq 0.01$ ) were more predominant in the indoor hens than the high outdoor hens. However, the RFN20 bacteria was more predominant ( $P \leq 0.05$ ) in the high outdoor hens than the indoor hens.

The cecal microbiota diversity of free-range hens at ASV level are shown in **Figures 8–10**. The alpha diversity at ASV level indicated no significant difference in microbiota populations within each of the rearing enrichment groups as assessed by Shannon index ( $P = 0.06$ ), Richness ( $P = 0.74$ ), Chao1 ( $P = 0.72$ ) and Simpson's index ( $P = 0.07$ ) (**Figures 8A–D**). The alpha diversity indicated no significant variation in the bacterial populations within each of the two range use groups of hens (**Figures 9A–D**) as assessed by Shannon index ( $P = 0.74$ ), Richness ( $P = 0.66$ ), Chao1 ( $P = 0.66$ ) and Simpson's index ( $P = 0.54$ ). The differences in beta diversity for rearing enrichments (**Figures 10A,B**) indicated by Redundancy analysis (RDA) ( $P = 0.001$ ) and ANOSIM (Bray–Curtis) ( $R = 0.06$ ,  $P = 0.001$ ) showed significant variation

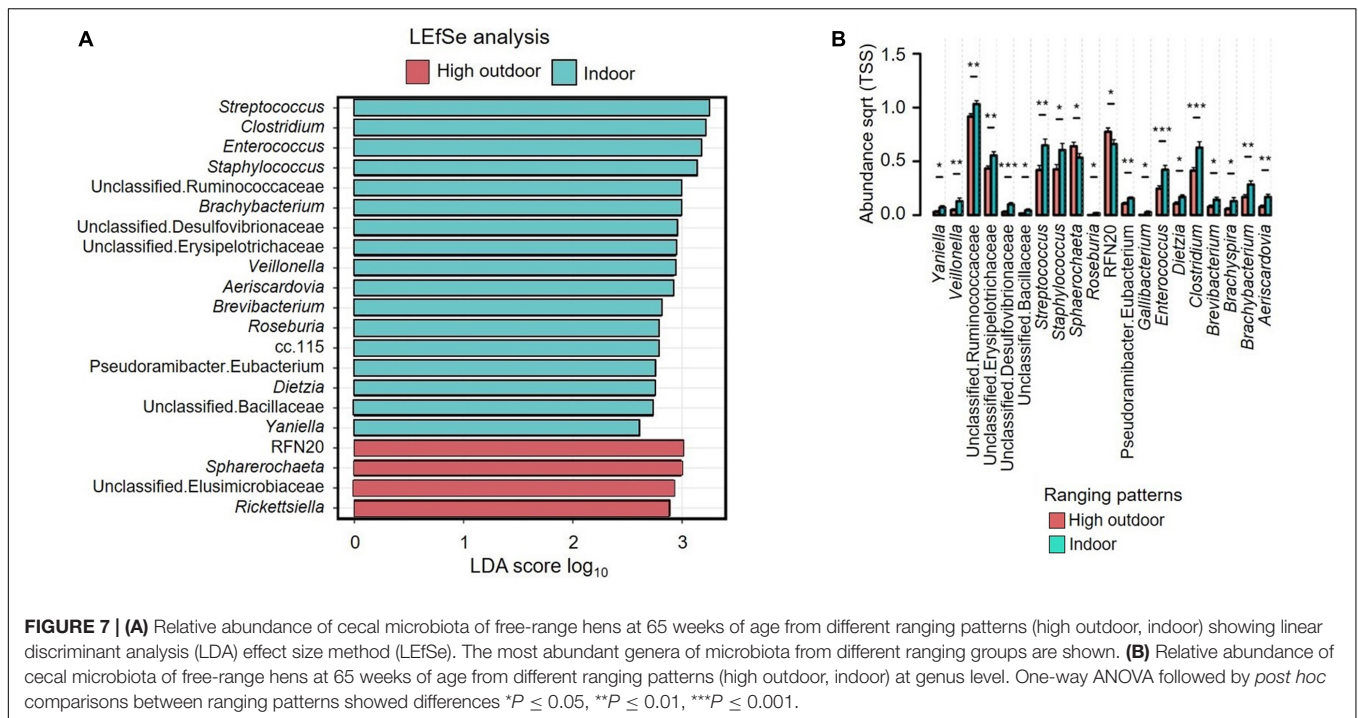




**FIGURE 5 |** Microbial community composition of free-range hens at 65 weeks of age from different ranging patterns (high outdoor, indoor). Clustered bar chart showing 20 most abundant genera.



**FIGURE 6 | (A)** Relative abundance of cecal microbiota of free-range hens at 65 weeks of age from different rearing enrichments (control, novelty, structural) showing linear discriminant analysis (LDA) effect size method (LEfSe). **(B)** Relative abundance of cecal microbiota of free-range hens at 65 weeks of age from different rearing enrichments (control, novelty, structural) at genus level. One-way ANOVA followed by *post hoc* comparisons between rearing enrichment groups showed differences  $^*P \leq 0.05$ ,  $^{**}P \leq 0.01$ ,  $^{***}P \leq 0.001$ .



in the bacteria between the rearing enrichment treatments. Similarly, the differences in beta diversity for ranging patterns (Figures 10C,D) indicated by RDA ( $P = 0.006$ ) and ANOSIM (Bray–Curtis) ( $R = 0.04$ ,  $P = 0.008$ ) showed significant variation in the bacterial diversity between the indoor and high outdoor hens.

### Short-Chain Fatty Acids

For the pullet SCFAs, the propionic acid content was higher in the novelty pullets than the control pullets ( $F_{2,27} = 4.04$ ,  $P = 0.03$ ) (Table 1). For the other SCFAs including formic acid, acetic acid, isobutyric acid, butyric acid, isovaleric acid, valeric acid, lactic acid, and succinic acid content, rearing enrichments did not have a significant effect (all  $P \geq 0.10$ ) (Table 1).

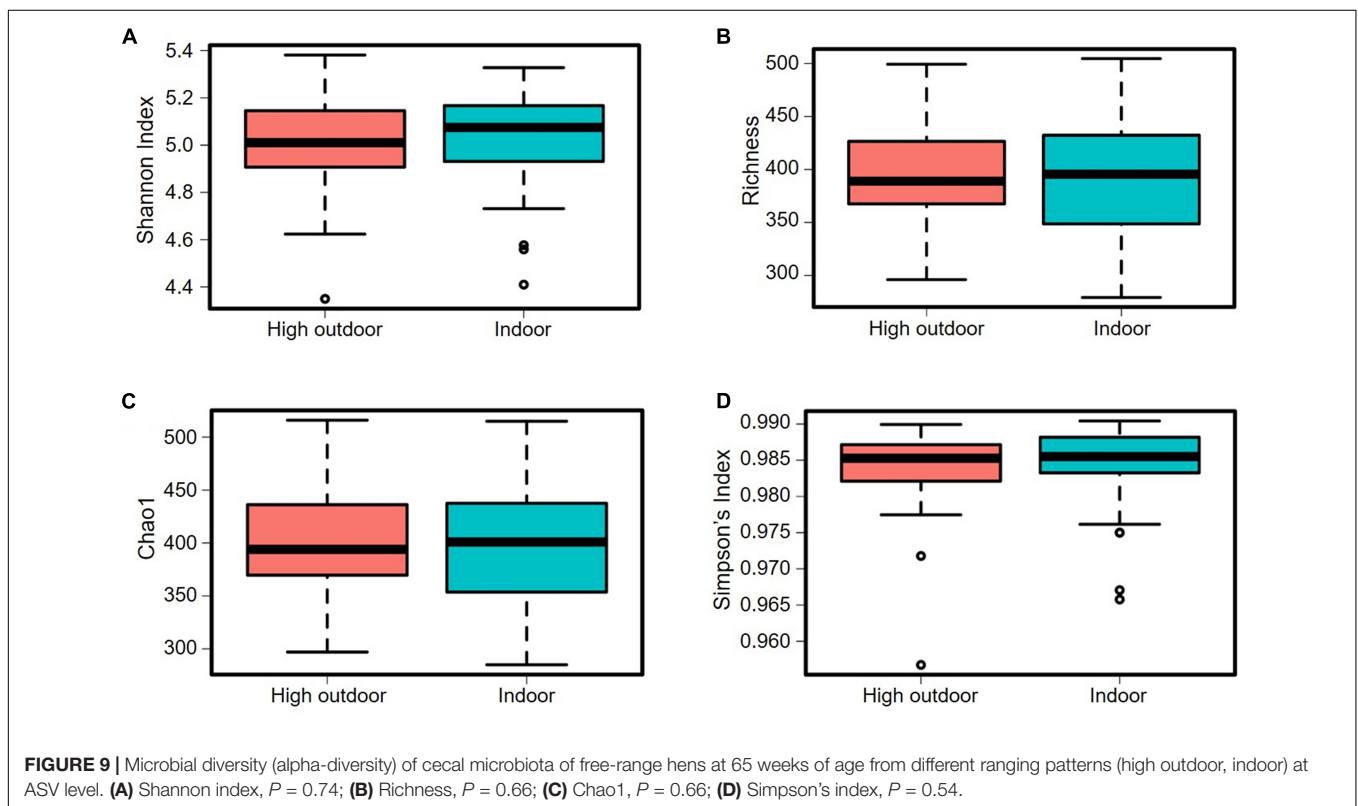
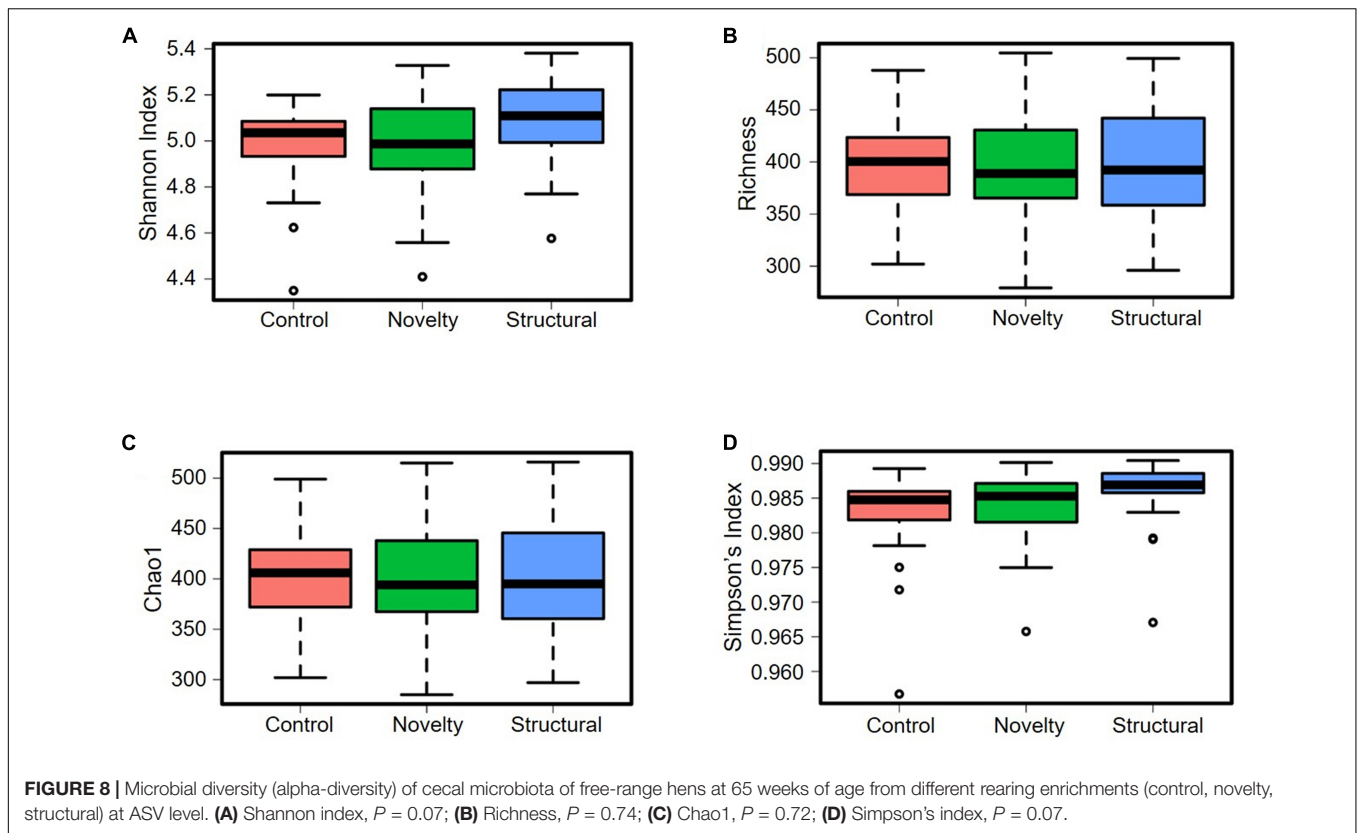
There were no significant effects of both rearing enrichments (all  $P \geq 0.14$ ) and ranging patterns (all  $P \geq 0.17$ ) for the SCFAs including formic acid, acetic acid, propionic acid, isobutyric acid, butyric acid, isovaleric acid, valeric acid, lactic acid, and succinic acid content in the ceca of free-range hens.

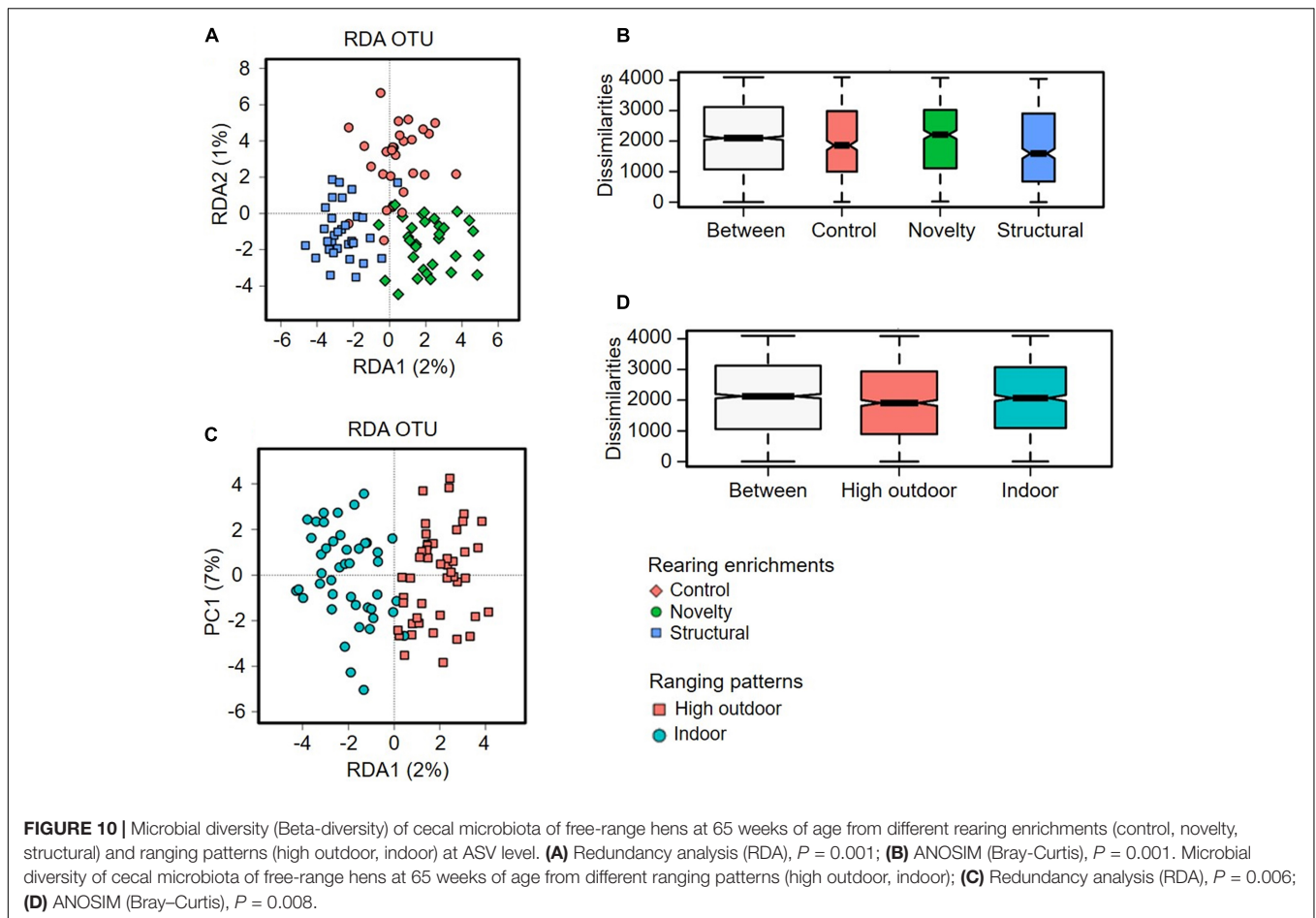
## DISCUSSION

This study was conducted to assess whether different types of environmental enrichments provided during the rearing period of laying hen pullets would affect their cecal microbiota composition and whether these rearing enrichments would continue to have impact on microbiota in adult free-range hens. Additionally, this study assessed the relationship between cecal microbiota composition and individual variation in range use patterns. Overall, the cecal microbiota abundance varied between

the enriched and non-enriched pullets with rearing differences persisting in the adult hens. Cecal microbiota abundance also varied between hens with different ranging patterns. Assessment of short-chain fatty acids detected minimal differences based on rearing treatments in pullets and no difference in adults, or any association with adult ranging patterns. Environmental variation has impact on laying hen microbiota, but the functional impacts of this variation remain to be determined.

The microbiota variation in the pullets might have been related to bacterial exposure from different novel objects that were placed in the shed and changed at weekly intervals. The addition of perching structures may have also had impact through accumulation of feces on the solid surfaces. Thus, the effects could have been solely related to external environmental variation in bacterial exposure. Alternatively, or in combination with external bacterial variation, there may have been an interplay between the behavior of the pullets and the colonization process for their microbial communities. Recent evidence from pigs, reported that enriched housing affected microbiota colonization relative to standard control housing (Wen et al., 2021), although this effect was not confirmed across other research (Saladrigas-García et al., 2021). The inclusion of enrichment in the pens resulted in some variation across the rearing period in exhibited behaviors (Campbell et al., 2022), responses to fear tests (Campbell et al., 2021), and body weight (Campbell et al., 2020c). In the week following the sample collection when the pullets were transferred to the laying facility, there were also rearing treatment differences in perching behavior in the new pens (Campbell et al., 2020c). In contrast, there were other behavioral test measures and post-mortem assessment of internal organs that did not show any rearing enrichment effects (Campbell et al., 2020c). Other research with laying hens has shown gut microbiota





composition varies relative to different genetic lines selected for high or low feather pecking behavior (Birkel et al., 2018; van der Eijk et al., 2019). When microbiota is transplanted between the lines, responses in subsequent behavioral testing are affected demonstrating the interaction that is present between gut microbiota and behavioral phenotype (van der Eijk et al., 2020). Similar evidence has been reported in quail selected for high or low emotional reactivity where transplantation between the lines can impact reactivity in subsequent emotional testing, modulating expected responses for the selected line (Kraimi et al., 2019a). Research in quail has also shown probiotics can affect responses in cognitive testing (Parois et al., 2017). Although there were microbiota differences among the rearing treatment groups, and other behavioral differences among the pullets, the causal link between the two is still uncertain with minimal effects of rearing enrichments detected in the short-chain fatty acids. Thus, there is great scope to conduct further studies to identify the causes and consequences of microbiota variation in relation to environmental enrichment and potential behavioral and health effects of the microbiota-gut-brain axis in poultry (Villageliu and Lyte, 2017; Kraimi et al., 2019b).

Rearing enrichments also affected the cecal microbiota abundance in adult hens. Upon transfer from the rearing facility, the adult hens were all housed in the same shed with fresh

litter, although they were separated into adjacent pens. Previous research has shown microbiota composition will change with age (Videnska et al., 2014), particularly following transfer from the rearing to the laying facility (Joat et al., 2021), but will be more similar between hens from the same housing shed compared with hens from different sheds of the same type of housing system (Schreuder et al., 2020). The adult hens retained rearing enrichment differences through to the end of the production cycle, although the specific phylogenetic groups that differed changed across the pullet and adult sampling points. In accordance with these microbiota differences, the rearing treatments had long-term impacts on other behavior, welfare, and egg quality measures in the birds (Bari et al., 2020a,b,c, 2021; Campbell et al., 2020b, 2022), although not on the short-chain fatty acids analyzed in the current study. These enrichment treatment differences support sustained effects of rearing environments regardless of the similar housing conditions experienced as adults. In terms of behavior, there were differences in how the hens used the range with the structural hens ranging for the longest time, the novelty hens showing the fewest range visits, and both the structural and novelty hens spent longer on the range during each individual visit relative to the control hens (Campbell et al., 2020c). These ranging differences likely resulted in corresponding dietary differences

**TABLE 1** | Least squares means (LSM)  $\pm$  SEM of short-chain fatty acids (SCFAs) ( $\mu\text{mol/g}$ ) in cecal contents of pullets and adult free-range hens from different rearing enrichment treatments (control, novelty, structural) and ranging patterns (indoor, high outdoor).

Parameters	Rearing enrichments			F ratio, P value	Ranging patterns		F ratio, P value
	Control	Novelty	Structural		Indoor	High Outdoor	
<b>Pullets</b>							
Formic acid	0.68 $\pm$ 0.10	0.80 $\pm$ 0.10	0.72 $\pm$ 0.10	$F_{2,27} = 0.09, 0.92$	–	–	–
Acetic acid	61.60 $\pm$ 5.95	78.75 $\pm$ 5.95	63.70 $\pm$ 5.95	$F_{2,27} = 2.48, 0.10$	–	–	–
Propionic acid	9.89 $\pm$ 0.86 <sup>b</sup>	13.19 $\pm$ 0.86 <sup>a</sup>	10.72 $\pm$ 0.8 <sup>ab</sup>	$F_{2,27} = 4.04, 0.03$	–	–	–
Isobutyric acid	0.55 $\pm$ 0.06	0.47 $\pm$ 0.06	0.53 $\pm$ 0.06	$F_{2,27} = 0.54, 0.59$	–	–	–
Butyric acid	14.12 $\pm$ 1.82	16.94 $\pm$ 1.82	4.24 $\pm$ 1.82	$F_{2,27} = 0.77, 0.47$	–	–	–
Isovaleric acid	0.17 $\pm$ 0.02	0.14 $\pm$ 0.02	0.18 $\pm$ 0.02	$F_{2,27} = 0.78, 0.47$	–	–	–
Valeric acid	1.34 $\pm$ 0.13	1.49 $\pm$ 0.13	1.33 $\pm$ 0.13	$F_{2,27} = 0.44, 0.65$	–	–	–
Lactic acid	1.08 $\pm$ 0.97	2.87 $\pm$ 0.97	1.15 $\pm$ 0.97	$F_{2,27} = 0.35, 0.71$	–	–	–
Succinic acid	1.42 $\pm$ 0.16	0.97 $\pm$ 0.16	1.35 $\pm$ 0.16	$F_{2,27} = 2.24, 0.13$	–	–	–
<b>Adult hens</b>							
Formic acid	1.01 $\pm$ 0.08	0.79 $\pm$ 0.08	0.80 $\pm$ 0.08	$F_{2,18} = 1.76, 0.20$	0.88 $\pm$ 0.07	0.84 $\pm$ 0.07	$F_{1,19} = 0.02, 0.89$
Acetic acid	66.75 $\pm$ 5.42	82.22 $\pm$ 5.47	77.06 $\pm$ 5.49	$F_{2,13} = 2.10, 0.16$	73.50 $\pm$ 4.47	77.19 $\pm$ 4.45	$F_{1,13} = 0.34, 0.57$
Propionic acid	27.37 $\pm$ 2.32	32.53 $\pm$ 2.34	31.92 $\pm$ 2.35	$F_{2,13} = 1.47, 0.27$	29.28 $\pm$ 1.91	31.93 $\pm$ 1.91	$F_{1,14} = 0.95, 0.35$
Isobutyric acid	1.28 $\pm$ 0.18	0.98 $\pm$ 0.18	1.03 $\pm$ 0.18	$F_{2,14} = 0.80, 0.47$	1.09 $\pm$ 0.15	1.10 $\pm$ 0.15	$F_{1,14} = 0.003, 0.96$
Butyric acid	11.23 $\pm$ 1.12	14.51 $\pm$ 1.13	13.55 $\pm$ 1.13	$F_{2,15} = 2.26, 0.14$	12.16 $\pm$ 0.93	14.03 $\pm$ 0.91	$F_{1,16} = 2.05, 0.17$
Isovaleric acid	0.45 $\pm$ 0.09	0.30 $\pm$ 0.09	0.32 $\pm$ 0.10	$F_{2,14} = 0.68, 0.52$	0.36 $\pm$ 0.08	0.36 $\pm$ 0.08	$F_{1,14} = 0.0002, 0.99$
Valeric acid	2.15 $\pm$ 0.24	2.47 $\pm$ 0.24	2.10 $\pm$ 0.24	$F_{2,17} = 0.72, 0.50$	2.19 $\pm$ 0.20	2.28 $\pm$ 0.19	$F_{1,17} = 0.11, 0.74$
Lactic acid	1.62 $\pm$ 1.05	1.47 $\pm$ 1.06	2.15 $\pm$ 1.05	$F_{2,17} = 0.34, 0.72$	1.47 $\pm$ 0.87	2.02 $\pm$ 0.85	$F_{1,18} = 0.44, 0.52$
Succinic acid	1.04 $\pm$ 0.15	0.99 $\pm$ 0.15	1.40 $\pm$ 0.15	$F_{2,17} = 2.03, 0.16$	1.17 $\pm$ 0.13	1.12 $\pm$ 0.13	$F_{1,17} = 0.10, 0.76$

<sup>a,b</sup>Dissimilar superscript letters indicate significant differences between rearing enrichment treatments ( $P < 0.05$ ).

across treatment groups in how often they consumed the formulated feed located inside the shed. Hens on the range would have had the opportunity to consume vegetation, insects, and grit with diet demonstrated to directly affect microbial communities (Borrelli et al., 2017; Zhou et al., 2021). The control hens also showed the poorest plumage condition across time indicating higher levels of feather pecking in these birds (Bari et al., 2020b). Similar to the pullets, the casual relationship between rearing enrichments, microbiota and other behavioral and welfare measures is unable to be confirmed from this study but warrants further investigation to understand these associations.

There was distinct variation in abundance of bacterial genera between the indoor and high outdoor hens showing these two groups could be differentiated by their microbiota profiles. While several previous studies have compared housing system effects on microbiota, including birds housed in indoor versus outdoor systems (Cui et al., 2017; Chen et al., 2019; Hubert et al., 2019; Schreuder et al., 2020, 2021; Seidlerova et al., 2020), to the authors' knowledge, this is the first study that has showed differences in these indoor/outdoor subpopulations of birds from the same communal housing system. These microbiota results align with other behavioral evidence distinguishing hens that range frequently versus remain inside the shed within the same free-range flock. Studies across multiple experimental and commercial systems have shown greater fear and anxiety in the indoor-preferring birds (reviewed in Campbell et al., 2020a). In an associated study conducted on a subsample of birds which included the individuals sampled in the current study, the indoor hens

showed poorer plumage condition, more comb wounds, but higher body weight than the high outdoor hens (Bari et al., 2020c). The differences in ranging patterns also likely led to differences in diet as hens outside had the opportunity to forage for insects and consume grit. As per the results with rearing enrichments, the casual relationship between the microbiota profiles and ranging patterns was unable to be established from the current study and is an area for further research to better understand and refine management practices for free-range hens. Similarly, individual resource use patterns by hens within an indoor system, such as the distinct variation demonstrated in aviary-housed hens and the areas of the system they frequent (Campbell et al., 2016; Rufener et al., 2018), may also correspond with variation in gut microbiota, but this would need to be confirmed.

## CONCLUSION

The different enrichments during rearing influenced the cecal microbiota composition of the pullets and this effect remained in the adult hens when they were moved to the same free-range shed and housed in identical pens. The subsequent individual differences in the ranging patterns also corresponded with differences in microbiota profiles with a greater abundance of different types of cecal microbiota genera in the indoor hens compared with the high outdoor ranging hens. Short-chain fatty acids predominantly did not differ across enrichment and ranging groups and further research is needed to understand

the causal relationships among the microbiota differences that were found.

## DATA AVAILABILITY STATEMENT

The data presented in the study are deposited in the NCBI repository, accession number PRJNA786377 (<https://www.ncbi.nlm.nih.gov/bioproject/786377>).

## ETHICS STATEMENT

The animal study was reviewed and approved by the University of New England Animal Ethics Committee (AEC17-092).

## AUTHOR CONTRIBUTIONS

MB and DC conceived and designed the experiment. CK initiated and assisted in the cecal sample collection. MB collected and analyzed the data, prepared the figures and tables, and drafted the manuscript. SK and YB contributed to the data processing

and analysis. S-BW provided expert guidance. DC wrote sections of the manuscript. All authors reviewed the manuscript critically and approved the final version.

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