Microbial interaction-driven community differences as revealed by network analysis

Zhe Pan a, Yanhong Chen a, Mi Zhou a, Tim A. McAllister b, Le Luo Guan a,⇑

a Department of Agricultural, Food and Nutritional Science, University of Alberta, Edmonton, AB, Canada
b Agriculture and Agri-Food Canada, Lethbridge Research Centre, Lethbridge, AB, Canada

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Diversity and compositional analysis are the most common approaches in deciphering microbial community differences. However, these approaches neglect microbial structural differences driven by microbial interactions. In this study, the microbiota data were generated from 12 rectal digesta samples collected from steers in which the Shiga toxin 2 gene (stx2) was not expressed (defined as Stx2− group) in the bacteria, and those with stx2 expressed (defined as Stx2+ group) and used to explore whether microbial networks affect gut microbiota and foodborne pathogen virulence in cattle. Although the Shannon and Chao1 indices of rectal digesta microbial communities did not differ between the two groups (P > 0.05), 24 and 13 taxa were identified to be group-specific genera for Stx2− and Stx2+ microbial communities, respectively. The network analysis indicated 12 and 14 generalists (microbes that were densely connected with other taxa) in microbial communities for Stx2− and Stx2+ groups, and 8 out of 12 generalists and 6 out of 14 generalists were designated to Stx2− and Stx2+ group-specific genera, respectively. However, the 66 core genera were not classified as network generalists. Natural connectivity measurements revealed that the higher stability of the Stx2− microbial network in comparison to the Stx2+ network, suggesting that the structure of each microbial community was inherently different even when their diversity and composition were comparable. Group-specific genera intensely interacted with other taxa in the co-occurrence network, indicating that characterizing microbial networks together with group-specific genera could be an alternative approach to identify variation in microbial communities.

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1. Introduction

Assessing microbial profiles using diversity, composition, and abundance measurements are broadly adopted approaches that have been widely applied to determine the role of the microbiome in host health. For example, the lower richness of human fecal microbiota is associated with dyslipidemia and insulin resistance leading to obesity [1], and higher evenness of milk/teat microbiota is associated with dairy cow mastitis [2]. In addition to these quantitative measures, microbial taxa interact within ecological niches, forming “micro-communities” that may function collectively [3]. Such microbial interactions can be influenced by the host, which in turn affect the host’s physiological activities. Hence, microbial interactions, together with microbial-host interactions, are critical for the establishment, maintenance, and function of microbiota [4].

The traditional and commonly used approach to identify microbial interactions is the construction of microbial co-occurrence networks using a correlation-based method (Pearson’s or Spearman’s correlation coefficient) [2,3,5,6]. This method is prone to detecting spurious correlations among low abundance taxa [7] and can lead to an ill-defined understanding of microbial interactions. These conventional methods also adopt subjective thresholds to define significant microbial interactions largely based on known biological information, while appropriate thresholds are hard to select, particularly for less studied and low abundant microorganisms [4,8]. Hence, effective ways to construct microbial co-occurrence networks are needed to enable an in-depth understanding of structural differences in microbial communities and how these could affect host-microbial interactions.

Shiga toxin-producing Escherichia coli (STEC) cause foodborne diseases that can lead to severe human infections (i.e. bloody diarrhea, hemolytic uremic syndrome) [9]. Cattle are the main asymptomatic carriers of STEC with the rectal-anal junction (RAJ) being the main colonization site [10]. As the result, cattle can shed STEC...
to the environment and therefore promote cattle-cattle/human transmission [11,12]. Our recent research has reported that host (i.e. host genes, immunity, microRNAs [13–16])-microbial interactions may play a role in affecting STEC colonization in cattle. In addition, Shiga toxins (stx) are major virulence factors in STEC with prototype toxins being designated as Shiga toxin 1 (stx1) and Shiga toxin 2 (stx2) [17] and our recent study revealed that stx2 expression was associated with host immune gene expression and potential STEC colonization [18]. Although the distinctive fecal microbial communities has been reported in super-shedder (cattle shed >10^6 STEC in feces per gram, SS) compared to non-shedders [19], little knowledge is derived from stx2 in STEC and its relationships with RAJ microbiota and microbial interactions. We speculated that microbial interactions together with the diversity and composition of RAJ microbiota are associated with stx2 expression and STEC colonization in cattle. Hence, this study aimed to assess the rectal microbial communities and interactions in response to stx2 expression in STEC at the RAJ, and the role of microbiota divergent in abundance among microbial interactions using the random matrix theory (RMT)-based method [20] and within-/among-module connectivity [21]. We aimed to identify keystone taxa and low abundant taxa contributing to microbial interaction networks and structural stability for better understanding the role of rectal microbiota in stx2 expression and potential STEC colonization in beef cattle.

2. Materials and methods

2.1. Animal study and sample collection

The animal trial followed the Canadian Council of Animal Care guidelines and was confirmed by the Animal Care and Use Committee, University of Alberta with Animal Care Committee protocol number AUP00000882. The animal trial and identification of stx2 gene abundance and expression were described in Pan et al. [18]. Briefly, ten cm^2 recto-anal junction (RAJ) tissue and 10 mL rectal contents were collected from a total of 143 feedlot cattle (585.84 ± 64.99 kg) within 30 min after slaughter at a federally approved abattoir.

DNA was extracted from 0.1 g powdered tissue using repeated bead beating and a column (RBBC) method [22], purified using the QiAmp Stool Mini Kit (Qiagen, Germany) and assessed based on absorbance at 260 and 280 nm using ND-1000 spectrophotometer (NanoDrop Technologies, Wilmington, USA). RNA was isolated from 0.1 g powered tissue using trizol reagent (Invitrogen Corporation, Carlsbad, CA, USA) method followed by manufacturer’s protocols and assessed using Agilent 2200 TapeStation (Agilent Technologies, Santa Clara, CA, USA) and Qubit 3.0 Fluorometer (Invitrogen, USA). Tissue extracted DNA was subjected to detection of stx2 gene abundance using quantitative PCR (qPCR) with primers 5’-ACCTGACACCCCTCTC-3’ and 5’-CAGTCTGAAAAGCTC-3’ [23]. Tissue extracted RNA was subjected to the identification of stx2 transcript using reverse transcription real-time quantitative PCR (RT-qPCR) with the aforementioned primers [23]. Both PCR and RT-qPCR were conducted in triplicates for each sample and followed the same thermal program on a StepOnePlus™ Real-Time PCR System (Applied Biosystems, Foster City, CA, USA): one cycle at 95 °C for 10 mins, followed by 40 cycles of 3 s at 95 °C, 30 s at 58 °C. Twelve rectal digesta samples collected from steers whose mucosal samples were confirmed to possess stx2 gene without expression (defined as Stx2– group) in the bacteria, and those with stx2 and were expressed (defined as Stx2+ group, n = 6) in STEC were selected with minimized differences in body weight and age at slaughter between two groups based on one-way ANOVA (Pbody weight = 0.07, Pslaughter age = 0.30, Table S1).

2.2. Amplicon sequencing and microbial community analysis

Total genomic DNA was extracted from frozen rectal digesta samples using the RBB method [21], purified using the QIAmp Stool Mini Kit (Qiagen, Germany) following the manufacturer’s protocols. The concentration and quality of DNA were further determined using the NanoDrop 2000 Spectrophotometer (Thermo Fisher Scientific, USA).

To generate the rectum bacterial compositional profiles, the bacterial V1-V3 region of the 16S rRNA gene was amplified using bacterial primers Ba9F (5’-GAAGTATGATCATGATACG-3’) and Ba515Rmod1 (5’-GCGGCCGCTGCGCAC-3’) [20]. The PCR amplification products were verified using agarose gel (2%) electrophoresis. Two-step PCR was performed for PCR amplicon generation and barcoding. In detail, the PCR was conducted with the following thermal program of initial denaturation at 94 °C for 2 mins, followed by 33 cycles of 94 °C for 30 s, annealing at 58 °C for 30 s, elongation at 72 °C for 30 s, and at last a final elongation at 72 °C for 7 mins. Furthermore, the second PCR was performed using amplicons produced in the first step for barcoding with the following program: initial denaturation at 95 °C for 10 mins, followed by 15 cycles of 95 °C for 30 s, 60 °C for 30 s, 72 °C for 60 s, and a final elongation at 72 °C for 3 mins.

All amplicon libraries were sequenced using an Illumina MiSeq pair-end 300 bp platform at Centre d’expertise et de services Génome Québec (Quebec, Canada). The raw sequence data were assigned to each sample according to the corresponding barcode and were processed using QIIME2 (Version 2019.10) [24]. Quality control, denoising, removal of chimeric sequences, and generation of amplicon sequencing variants (ASVs) were performed using the QIIME2 plugin DADA2 [25]. Taxonomic classification was performed in QIIME2 using a taxonomic classifier with the SILVA database (version 132) as the reference. The Good’s coverage index was used to evaluate the adequacy of sequencing depth to generate bacterial profiles in each sample. For diversity analyses, alpha diversity was estimated using Shannon (evenness) and Chao1 (richness) indices. Beta diversity was evaluated based on Weighted Unifrac distance using phylogenetic distances across identified taxa in a phylogenetic tree and the abundance of each feature to determine the similarity between Stx2– vs. Stx2+ groups. All diversity metrics were calculated using scripts implemented in QIIME2. The visualization of alpha- and beta- diversity was performed using the ggplot2 package in R. Differentially abundant (DA) genera (Absolute log2 fold change > 1 and false discovery rate <0.05) between Stx2– vs. Stx2+ groups were identified using the DESeq2 package in R. The cut-off of DA genera and group-specific taxa is a relative abundance >0.1% and presence in at least two out of six animals in each group.

2.3. Construction of microbial co-occurrence networks

The random matrix theory (RMT)-based method [8] was employed to construct the microbial co-occurrence network to identify microbial interactions using molecular ecological network analysis (MENA) (http://ieg4.rcc.ou.edu/­mena) [8]. Briefly, the absolute abundance of microbial genera data was uploaded. The absolute abundance microbial genera dataset was appropriately standardized and the pairwise Pearson correlation suggested by the author’s manual was employed to generate correlation coefficients and a similarity matrix firstly [8]. The similarity matrix was subsequently transformed into an adjacency matrix by applying the automatic generated threshold to the correlation values based on the RMT approach [8]. In this study, a connection stands for a strong (Pearson’s r > 0.85) and significant (P < 0.01) correlation. The visualization of the co-occurrence network was performed using gephi (0.9.2).
2.4. Characterization of topological properties of co-occurrence networks

The modularity of each network was estimated using MENA along with the evaluation of other topological properties, including the clustering coefficient [26,27], average path length [28], graph density [29], and average degree [28]. Random networks and power-law distribution assessments were generated to evaluate whether empirical networks were prone to error and to identify microbiota interactions that were due to non-random patterns that represent the empirical structure of microbial communities [30,31]. Random networks were constructed based on the Maslov-Sneppen method [31] using MENA, which kept numbers of nodes (microbial taxa) and edges (connections) unchanged, but rewired positions of all links in the network.

From the network modularity perspective, taxa could be classified into network hubs, module hubs, connectors that represent generalists in the community, and peripherals that represent specialists in the community [32]. Generalists refer to taxa that are highly connected with others both within and among modules (network hubs), within a module (module hubs), and among different modules within a network (connectors). Specialists represent peripheral taxa that interact less with other taxa (including a node that is only connected within a module or at least 60% links within the module) [32]. Within-module connectivity (Zi) and among-module connectivity (Pi) were computed based on the following algorithm [32]:

\[ Z_i = \frac{k_i - \bar{k}_s}{\sigma_{ki}} \]

where \( k_i \) is the number of links of node \( i \) to other nodes in its module \( s \), \( \bar{k}_s \) is the average of \( k \) across all nodes in \( s \), \( \sigma_{ki} \) represents the standard deviation of \( k \) in \( s \). Hence, \( Z_i \) (within-module connectivity) characterizes to what extent node \( i \) is connected to others in its module.

\[ P_i = 1 - \sum_{s=1}^{N_{out}} \left( \frac{k_{si}}{k_i} \right)^2 \]

where \( k_{si} \) is the number of links of nodes \( i \) to nodes in module \( s \), \( k_i \) is the total degree of node \( i \) within a network. Therefore, \( P_i \) (among-module connectivity) of a node stands for evenly distribution of links among all modules if its value is close to 1, and 0 if all its links are within its own module.

Zi and Pi were characterized using MENA with the classification as follows: network hubs (Zi > 2.5; Pi > 0.62), module hubs (Zi > 2.5; Pi < 0.62), connectors (Zi < 2.5; Pi > 0.62); peripherals (Zi < 2.5; Pi < 0.62) [21]. The thresholds for classifying nodes into aforementioned four roles in the network were determined by both heuristic determinations and the concept of 'basin of attraction' [21]. Briefly, Zi and Pi were both computed for each node in the network, and density plots were adopted to visualize the gradient of the value of Zi and Pi that can ‘flow’ to the local minimum (termed as ‘basin of attraction’). In other words, the region of the space that ‘flow’ toward a certain minimum value is therefore regarded as the optimal threshold for Zi and Pi being 2.5 and 0.62, respectively [21].

Network stability is a critical component that tests if a network is resilient to perturbations sourced from external factors other than interactions [33,34]. Here, natural connectivity [35] was introduced to describe network stability differences in response to the stx2 expression in STEC. The estimation of natural connectivity was based on the following algorithm:

\[ \text{ave}(\lambda) = \ln \left( \frac{1}{N} \sum_{i=1}^{N} e^{\lambda_i} \right) \]

where \( \text{ave}(\lambda) \) is the natural connectivity, \( N \) is the number of nodes in the network, \( \lambda_i \) is the eigenvalue of the adjacency matrix. A total of 137 nodes representing 80% of total nodes were randomly removed from the adjacency matrix and \( \lambda_i \) and \( \text{ave}(\lambda) \) were recalculated after each removal. The visualization of the natural connectivity was performed using the ggplot2 package in R.

The identification of potential hub microbial taxa in the network was validated using Lasso regression in R package ‘glmnet’. Twelve samples were regarded as the training data with 9 samples (6 Stx2+; 3 Stx2−) as the external test data. The accuracy rate (the number of samples recognized correctly / total number of samples) was estimated to determine the model classification performance.

2.5. Data availability

All the sequencing data used in the current study has been submitted to NCBI Sequence Read Archive (SRA) under the accession numbers from SRR14769039 to SRR14769050 (BioProject ID PRJNA736180).

3. Results

3.1. Taxonomic assessment of the RAJ content-associated microbiota

A total of 179,917 filtered pair-end reads were generated with 14,993 ± 437 (mean ± SE) sequences per sample, and a total of 2,798 amplicon sequence variants (ASVs) were identified ranging from 174 to 275 (Table S2). The Good’s coverage was >99.9% for all samples, indicating adequate sequencing depth.

From all samples, >99% of reads were classified into 13 phyla, with Firmicutes (72.7 ± 2.0%) and Bacteroidetes (24.6 ± 1.9%) being the most predominant phyla (relative abundance >10%, Additional file 1). At the family level, 59 families were identified with 7 unclassified and 52 classified (Additional file 1, of which Ruminococcaceae (47.2 ± 1.5%), Lachnospiraceae (9.2 ± 1.2%), and Prevotellaceae (8.8 ± 1.1%) were the most predominant families. At the genus level, 154 taxa were identified (Additional file 1, 20 unclassified; 134 classified), of which Ruminococcaceae UCG-005 (23.1 ± 2.1%) and coprostanoligenes group (9.1 ± 0.7%) from Ruminococcaceae family and Christensenellaceae R-7 group (7.3 ± 0.7%) from Christensenellaceae family were the most abundant genera. The most frequently detected genera (average relative abundance >0.5%; Additional file 1) belonged to Firmicutes (19 out of 27, 4 unclassified, 15 classified) and Bacteroidetes (8 out of 27, 1 unclassified, 8 classified), respectively.

There were 52 to 75 genera identified from each sample with 15 core genera shared by all 12 samples (Fig. 1A, Table S4). Specifically, 30.7% (4 out of 13 phyla, Firmicutes, Bacteroidetes, Actinobacteria, Proteobacteria), 18.6% (11 out of 59 families, 6 out of 11 belonged to Firmicutes; 5 out of 11 belonged to Bacteroidetes; Additional file 1) and 9.7% (15 out of 154 genera, 9 out of 15 belonged to Firmicutes; 6 out of 15 genera belonged to Bacteroidetes; Fig. 1A, Table S4,) were present in all samples. In addition, twenty-four genera were Stx2− specific and 13 genera were Stx2+ specific with 66 genera shared by both groups (Fig. 1B). Genera belonging to Firmicutes were the most predominant taxa in Stx2− (10 out of 24) and Stx2+ (4 out of 13) groups (Table S5).

3.2. Comparable diversity and composition of RAJ content-associated microbiota between Stx2− and Stx2+ groups

Neither Shannon nor Chao1 indices differed (Kruskal Wallis test, P > 0.05, Fig. 2A) between the bacterial communities from Stx2− and Stx2+ groups. Further comparison of the similarity of microbiotal communities between the two groups using ANOSIM (Analysis of similarities) revealed no clustering patterns (P = 0.52, Fig. 2B) at the phylum (P > 0.5, Fig. 3), family, or genus.
No differentially abundant taxa at the phylum, family, or genus level were identified between Stx2+/C0 and Stx2+ groups.

3.3. General co-occurrence patterns in each co-occurrence network

The non-random co-occurrence patterns were observed based on the significant power-law distribution in each group ($R^2_{\text{Stx2+}} = 0.98, P < 0.05; R^2_{\text{Stx2-}} = 0.97, P < 0.01$) as well as the greater value of structural properties in the empirical as compared to the random networks (Table 1). Hence, non-random empirical co-occurrence networks were established to uncover real-world microbial interactions. The empirical network consisted of 86 nodes (genera) with 322 edges (mean of 7.49 edges per node) for Stx2− (Table 1, Fig. 4A), and 77 nodes with 243 edges for Stx2+ (Table 1, Fig. 4B). The average network distance between all paired nodes (average path length, APL) was 2.84 edges with a diameter (longest distance) of 8 edges in the Stx2− network and 2.86 edges with a diameter of 7 edges in the Stx2+ network, respectively (Table 1). The clustering coefficient (the degree to which nodes tend to cluster together) was 0.32 for the Stx2− network and 0.33 for the Stx2+ network, respectively (Table 1). The modularity index was 0.44 for the Stx2− network and 0.46 for the Stx2+ network (values >0.4 suggest that the network has a modular structure [36]) (Table 1).

3.4. Identification of keystone taxa and their associations with different co-occurrence patterns

High modularity was observed for both networks with 7 and 6 observed modules in the Stx2− and Stx2+ co-occurrence networks, respectively (Fig. 4A, B) with no network hubs being identified. More than 80% of nodes (Stx2−: 74 out of 86, 86.0%; Stx2+: 62 out of 77, 80.6%) with an abundance >0.2% were classified as
peripherals (Fig. 5A). Only 14% (connectors: 11 out of 86; module hubs: 1 out of 86; Stx2 group) and 19.4% (connectors: 14 out of 77; module hubs: 1 out of 77; Stx2+ group) of taxa from Stx2 and Stx2+, respectively, were designated as specialists (Fig. 5A). Specifically, >85% (23 out of 27) of connectors had a lower abundance (<0.2%) with a high Pi value (0.62–0.77) (Fig. 5B).

Fig. 3. Comparison of average relative abundance at the phylum level between Stx2− and Stx2+ groups. Circos plots were used for visualization of average relative abundance at the phylum level between Stx2− and Stx2+ groups. Phyla in each group with a total relative abundance of >0.1% in at least three samples were included. The Kruskal-Wallis test was used to determine the average relative abundance of phyla was comparable (P > 0.05) between Stx2− and Stx2+ groups. P-values were false discovery rate (FDR, q = 0.05) adjusted.

Table 1
Topological properties of empirical and random network between Stx2− and Stx2+ groups.

|                  | Empirical network | Random network |
|------------------|-------------------|----------------|
| Group            | Stx2−             | Stx2+          | Stx2−            | Stx2+           |
| Nodes            | 86                | 77             | 86               | 77              |
| Edges            | 322               | 243            | 322              | 243             |
| Modularity (MD)  | 0.44              | 0.46           | 0.29 (±0.011)    | 0.31 (±0.012)   |
| Clustering coeff. (CC) | 0.32          | 0.30           | 0.12 (±0.014)    | 0.12 (±0.016)   |
| Average path length (APL) | 2.84         | 2.86           | 2.47 (±0.030)    | 2.58 (±0.033)   |
| Graph density (GD) | 0.088            | 0.083          | 0.09             | 0.08            |
| Average degree (AD) | 7.49           | 6.31           | 7.49             | 6.31            |

Fig. 4. Co-occurrence networks of bacterial genera in (A) Stx2− and (B) Stx2+ groups. The size of each node is proportional to the number of connections (that is, degree); the color of connections between two nodes represents a positive (red) or a negative correlation (blue). Each module is presented as a specific color. (For interpretation of the references to color in this figure legend, the reader is referred to the web version of this article.)
Furthermore, generalists formed differential clustering patterns between Stx2− and Stx2+ networks. Generalists were evenly distributed across identified ecological clusters in the Stx2− network. Four out of fourteen generalists belonged to module 3, whereas other generalists were in module 1 (3 out of 11), module 2 (2 out of 11), module 4 (1 out of 11), module 5 (1 out of 11) and module 6 (1 out of 11). However, in the Stx2+ network generalists were not evenly distributed with 7, 4, 3, and 1 out of 14 generalists belonging to modules 1, 3, 2, and 4, respectively.

Moreover, decreased stability of the network was evidenced by reduced natural connectivity in the Stx2+ network in comparison to the Stx2− network (Fig. 6). The natural connectivity that supported network stability in each group gradually decreased with the increasing number of removed nodes, while natural connectivity in the Stx2+ network always being lower than that in the Stx2− network regardless of the numbers of nodes removed from each network (Fig. 6).

3.5. Group-specific taxa as keystone taxa in microbial interactions

More than 50% of the group-specific genera (8 out of 12 in Stx2−; 6 out of 12 in Stx2+, Table 2) belonged to generalists with a lower abundance (relative abundance <0.2%). Group-specific generalists in the Stx2− network included: Parvibacter, Candidatus saccharimonas, Acetitomaculum, as well as unknown genera within the Bacteroidetes, Proteobacteria, Tenericutes, Cyanobacteria, and the Veillonellaceae. Besides, among generalists in the Stx2+ network, Prevotellaceae Ga6A1 group, Flexilinea, Ruminiclostridium 9, Ruminococcus 1, Acetobacter, and Streptomyces were group-specific genera in microbial communities in the Stx2+ group. All the core microbes shared by two groups (Fig. 3B) were regarded as specialists that were poorly connected with other taxa within networks.

4. Discussion

This study assessed microbial interactions in response to stx2 expression in STEC at RAJ in beef steers, shedding light on microbial mechanisms regulating STEC colonization as well as providing novel approaches to understanding differences in microbial communities. The compositional profiles of microbial communities identified in our study were comparable to those identified from the rectum content of dairy cattle [37], and fecal microbiota of beef cattle [19,38] with the most dominant phyla being Firmicutes and Bacteroidetes with an accumulative relative abundance of up to 94.1%, 89.5%, and 80.6%, respectively. The proportion of three main families (Ruminococcaceae (47.2 ± 1.5%), Lachnospiraceae (9.2 ± 1.2%), and Prevotellaceae (8.8 ± 1.1%)) were also similar to

![Figure 5](image-url) Distribution and average relative abundance of genera based on their network roles. A. Distribution of bacterial genera based on their network roles. Nodes in the network were classified as peripherals, modular hubs, or connectors. No network hubs were identified in networks from both groups. B. Average relative abundance of connectors.

![Figure 6](image-url) The natural connectivity representing the network stability of co-occurrence networks in both Stx2− and Stx2+ groups.

Table 2

| Phylum               | Genus                        | Stx2− specific | Stx2+ specific |
|----------------------|------------------------------|----------------|----------------|
| Actinobacteria       | Parvibacter                  |                |                |
| Bacteroidetes        | f_Bacteroidales RF16 group   | Stx2− specific |                |
| Cyanobacteria        | o_Gastranaerophilales        | Stx2− specific |                |
| Firmicutes           | f_Veillonellaceae            | Stx2− specific |                |
|                      | Acetitomaculum               | Stx2− specific |                |
| Patesicibacteria     | Candidatus saccharimonas     | Stx2− specific |                |
| Planctomycetes       | p-1088-a5 gut group          | Stx2− specific |                |
| Proteobacteria       | Parasutterella               | Stx2− specific |                |
| Tenericutes          | o_Izimaplastales             | Stx2− specific |                |
| Stx2+ group          | Saccharopolyspora rectivirgula | Stx2+ specific |                |
| Actinobacteria       | Prevotellaceae Ga6A1 group   | Stx2+ specific |                |
| Bacteroidetes        | Prevotellaceae UCG-001       |                |                |
|                      | Parabacteroides              |                |                |
| Chloroflexi          | Flexilinea                   | Stx2+ specific |                |
| Firmicutes           | Oscillibacter                | Stx2+ specific |                |
|                      | Ruminiclostridium 9          | Stx2+ specific |                |
|                      | Ruminococcus 1               | Stx2+ specific |                |
|                      | f_Erysipelotrichaceae        |                |                |
|                      | Cellulosilyticum             |                |                |
| Planctomycetes       | p-1088-a5 gut group          |                |                |
| Proteobacteria       | Acetobacter                  | Stx2+ specific |                |
|                      | Parasutterella               |                |                |

the core microbes shared by two groups (Fig. 3B) were regarded as specialists that were poorly connected with other taxa within networks.
the bacterial taxa identified in a study where cattle were shedding >10^4 cfu/g of *E. coli* O157 in that 31.8%, 10.5%, 9.0% of total reads from fecal microbiota were assigned to *Ruminococcaceae*, *Prevotellaceae*, and *Lachnospiraceae*, respectively [19]. Although all cattle were raised under the same high-grain diet and management conditions and were similar in age and body weight, individual variations of microbial communities were still observed including the relative abundance of each microbial community and the proportion of predominant taxa shared by each microbial community. Variation in the microbial fecal profile of individual cattle has been previously reported [19], a finding that has been attributed to differences in age, weight, and diet. As these variables were relatively consistent in our study, other host factors along with microbial crosstalk within the microbiota might be the main drivers for the individualized microbial composition.

Despite highly similar microbial profiles including comparable alpha and beta diversity between Stx2− and Stx2+ groups, microbial interactions within the microbiota varied. Microbes can interact with each other for the purpose of co-evolution [39] leading to adaptation and specialization [39] of certain microbial taxa, which promote future alterations in the microbial community. To date, the widely used approaches to study the microbial interactions is based on network construction mainly using correlation-based (Pearson’s and Spearman correlation coefficient) and maximal information coefficient (MIC) methods that have proven to be less useful in inferring microbial ecological networks assessed by area under the precision-recall curves (AUPR) [3]. These metrics (i.e., correlation-based approaches, MIC) are less applicable to microbial compositional data as the assumption of independent variables can be satisfied, leading to the generation of spurious correlations [3,40], and therefore are less powerful for inferring real microbial interactions. To overcome such limitations in this study, similarity matrix was first established using Pearson’s correlations and then the RMT-based approach was employed to determine the reference point which enables automatic threshold selection and minimizes noise. The RMT-based approach was established based on two universal roles of random matrix theory: the distribution of two nearest eigenvalues follows Gaussian orthogonal ensemble (GOE) statistics correlations exist (a true correlation will follow GOE distribution in RMT theory), while it follows Poisson distribution if there is no correlation [41]. Particularly the transition between GOE and Poisson distribution serves as the reference point to distinguish non-random relationships (that is true correlations) in the data matrix from background noise [41]. In other words, a correlation refers to the GOE distribution while non-correlation represents the Poisson distribution and the transition point from GOE to Poisson distribution is the reference point for the automatic generated threshold used for the construction of the microbial network and the identification of meaningful correlations from the noise (i.e. certain correlations might not follow GOE and therefore could be noise or spurious correlations). Hence, compared to studies that only adopt correlations (Pearson’s and Spearman correlation coefficient) for constructing microbial networks, the RMT-based approach is more effective at identifying true interactions within the response networks to stx2 expression.

Comparison of random and empirical networks based on topological properties, confirmed constructed networks were effective for investigating interactions between stx2 expression and microbial communities. Particularly, modules (clusters) were powerful topological features to reflect network differences, referring to the fundamental units whose constituent elements (nodes) are functionally similar in terms of specific chemical, and biological processes [42]. Previous explorations in soil-microbial interactions revealed that clusters in the network have specific and different functions that enable microbes to respond to different soil conditions [43,44]. For instance, three major modules with diverse functions (electron-transfer, biogeochemical C- and N-cycles, organic contaminant degradation) were characterized for microbial communities from soil contaminated with oil [44]. Clusters were shown to be crucial components of microbial communities in the network and necessary to develop an understanding of modularity in networks and microbial interactions. The Stx2− and Stx2+ groups formed comparable numbers of clusters while clusters in each group included different taxa, suggesting functions of clusters in each group differed in their interactions in response to stx2 expression in STEC at RAJ. Regardless, the highly connected genera among densely connected clusters of nodes (that is, modules) were observed in both Stx2− and Stx2+ groups resulting in the formation of ‘small world’ topologies, indicative of empirical networks that are more clustered than random networks.

Identifications of generalists also furthered the understanding of microbial community structure and differential microbial interactions, which play an empirical rather than theoretical role among microbial interactions. Particularly, identified generalists in our study are group-specific taxa, which are with low relative abundance but might play unique roles such as regulating microbial interactions within each network through nutritional supplementation or competition. *E. coli* O157:H7 can utilize ethanolamine as the free nitrogen source in the bovine small intestine, thus the presence or absence of ethanolamine utilizing bacteria could be a contributing factor to diverse microbial communities [19,45]. In our study, *Streptomyces* identified as Stx2+ specific genera were capable of metabolizing ethanolamine [46], which might generate a niche for the survival of stx2 expressed bacteria. However, not all group-specific generalists in microbial communities play a role in nutrient supplementation for STEC. For instance, *Acetobacter* (a group-specific generalist from the Stx2+ group) is capable of converting ethanol to acetic acid which can inhibit STEC [47]. A similar case is also observed in Stx2− group that *Acetitomaculum* (a group-specific generalist from Stx2− group), an aceticogenic bacteria [48], may also suppress STEC. It is noticeable that >85% (23 out of 27) generalists were low abundant taxa with an average abundance <0.2%, highlighting the irreplaceable role of rare abundant taxa among microbial interactions. For instance, rare *Methanotrophs* (*i.e.* *Methylocaldum*) acting as ‘primary producers’ in methane-driven food webs [49], rare symbionts (*i.e.* *Symbiodinium*) increase coral-algal assembly stability with environmental alterations [50]. Hence, these results raise the possibility that less abundant group-specific generalists contribute to the differential degradation of organic matter and play a practical role in microbial interactions by mediating nutrient availability in a manner that may have positive or negative effects on STEC colonization. The lack of well-defined approaches for determining keystone taxa from networks embedded our understanding of microbial interactions. Previous studies adopted general topological properties (*i.e.* betweenness centrality) for inferring keystone taxa [51], which neglects the fact that networks tend to be modular. Identification of generalists (or keystone taxa) following the concept of among/within-module connectivity provides the added advantage for further understanding of microbial interactions. Thus, the approach in our study of using the concept of modularity to compute the role of each node is more representative of network modularity and the role of each node with/among the overall structure.

As a supportive approach, lasso regression revealed group-specific microbes contributed to the mucosa-associated stx2 expression. Among the 7 selected genera based on lasso regression, 2 out of 7 were core and 3 out of 7 were group-specific genera (Table S6). However, all of the core microbes were identified as peripherals that were poorly connected with other taxa and were not expected to affect the robustness of the network [52]. Compared to studies that emphasize the value of core microbiota in the maintenance of microbial communities, group-specific genera
exhibiting diverse functions could potentially interact with the host and Stx2-carrying/Stx2-expressed bacteria, leading to the establishment of a different co-occurrence network. Hence, our study highlights the valuable role of group-specific taxa in the microbial co-occurrence network which advances the understanding of microbial interactions in terms of stx2 expression and potential STEC colonization. It is noticeable that the identified microbial networks should be validated in the future. In addition to the mathematical algorithm that reduces spurious correlations, adding stable isotope probes in the DNA-derived microbial community and analyzing heavy-isotope enriched DNA could be alternative approach, which could differentiate inactive microbial interactions from ecological meaningful microbial crossstalk left [53,54]. For instance, a DNA-based stable isotope probing approach using [13C]CH₄ has been employed to identify real microbial interactions by co-occurrence analysis in ombrotrophic peatlands [53]. However, using heavy isotope probes to examine true microbial interactions within the dense microbial communities of the gut needs further evaluation.

5. Conclusions

Overall, our results revealed comparable diversity and composition of RAJ microbiota were observed between Stx2- and Stx2+ animals with >60% genera recognized as members of the core microbiome. However, RMT-based network analysis revealed varied microbial interactions, keystone taxa, and stability of microbial communities in response to stx2 expression. Group-specific taxa play an unusual role in the network which might drive microbiota-stx2 interactions. This study also constitutes an in-depth understanding of host-STE ucz networks and highlights the possibility of altering the gut environment to mitigate stx2 expression through modifying the gut microbiota. However, future validations using a larger sample size are needed to verify the proposed methods for deciphering microbial community differences as well as the principles of microbe coexistence that determine microbial interactions. Regardless, our findings highlight the critical role of group-specific genera among microbial interactions and shed light on using an integrated approach that integrates group-specific genera with network analysis to identify and characterize differences in microbial communities with comparable microbial profiles.

Author statement

ZP involved in concept development, performed data analysis and manuscript writing. YC contributed to sample collection, lab work and data generation. MJZ involved in microbiota data analysis and manuscript revision. TAM involved in data interpretation and manuscript revision. LLG involved in the concept of the study, data analysis and manuscript writing.

Declaration of Competing Interest

The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

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Appendix A. Supplementary data

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