Beef cattle that respond differently to fescue toxicosis have distinct gastrointestinal tract microbiota

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Abstract
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Keywords
Fescue toxicosis, tall fescue, microbiota, beef cattle, ITS1, Angus

Disciplines
Agriculture | Animal Sciences | Environmental Microbiology and Microbial Ecology | Large or Food Animal and Equine Medicine

Comments
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Beef cattle that respond differently to fescue toxicosis have distinct gastrointestinal tract microbiota

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Key words: Fescue toxicosis, tall fescue, microbiota, beef cattle, ITS1, Angus
Abstract

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Introduction

Tall fescue (*Lolium arundinaceum*) is a common cool season grass used widely as forage for grazing livestock in the southeastern United States. The grass shares a symbiosis with *Epichloë coenophiala*, a fungus that grows as an endophyte within the plant and provides heat and herbivory resistance from both insects and mammals. The fungus produces ergot alkaloid compounds that have been shown to cause fescue toxicosis (FT) in grazing livestock species such as cattle, sheep and goats [1-3]. Many biologically active alkaloids are produced by the fungus, but ergovaline, an ergopeptide, is commonly thought to be responsible for FT [2].

Induced by the consumption of these toxic ergot alkaloids, FT is a metabolic disease that, in ruminant livestock, manifests as vasoconstriction, higher body temperature, suppressed appetite, and reduced heart rate and prolactin levels [1, 3, 4]. This disease causes an estimated loss of $2 billion US dollars each year due to reduced body weight (BW), milk yields, and rate of calving [5].

Efforts to reduce or eliminate FT included removal of endophyte infected fescue and planting cultivars of endophyte-free [3] fescue. The endophyte free fescue improved cattle performance, but resulted in a general weakening of the plant, reducing tolerance to insects, nematodes, high temperatures and overgrazing [3]. Additionally, researchers focused on the endophyte, identifying strains that produce lower levels of the ergot alkaloids while still providing drought and insect resistance for the grass. Cattle fed fescue infected with low ergot producing strains improved cattle performance similar to endophyte free strains, and are being sold commercially [3, 6, 7]. Finally, management practices such as interseeding higher levels of different plants, rotational grazing, fertilizing with low nitrogen fertilizers and reducing seed heads that contain high-ergot producing endophyte strains in pasture were shown to reduce the toxin levels within the forage [3].
Another strategy to limit the negative effects of the tall fescue endophyte on cattle could be the identification of animals with greater tolerance to FT. Studies on host response of FT are still limited in the literature, with most of them focusing on breed differences [8-11]. Recently, using the same animals as in this study, Khanal et al., 2016 [12] identified pregnant Angus cows showing distinct growth potential under FT. Cows classified as tolerant to FT had higher growth and body condition score, and lower rectal temperature and hair coat score than susceptible animals. Using the same data, Khanal et al., 2018 [13] identified 550 differentially expressed (DE) genes between tolerant and susceptible cows, with in which the most DE genes had functions such as regulation of vasoconstriction and hair coat shedding. With respect to host genomics, the few reports available in the literature have focused on candidate gene approaches [14-16].

The current literature published on the effect of toxic tall fescue effects on gastrointestinal (GI) microbiota is still limited. Most recently, Mote et al., 2019 [17] surveyed fecal bacterial communities of beef cattle during a FT challenge, and described possible connections between the abundance of certain bacterial phylotypes and the host response to FT. Other studies have suggested microbial communities within the cow rumen [18, 19], earthworm’s intestine [20], and soil [21] are able to degrade ergovaline. It is thus conceivable that the GI tract microbiota may be able to alleviate some of the impact of FT symptoms. We hypothesize that both the bacterial and fungal microbial communities within the GI tract are associated with FT tolerance.

In this study, we analyzed fecal microbial communities from cattle with contrasting growth performance during a chronic exposure FT challenge. Our goal was to identify shifts in bacterial, archaeal and fungal microbial populations (using 16S rRNA gene and ITS1 region amplicon sequencing, respectively) between the two tolerance groups across two different locations.
Materials and methods

Ethics statement

All animal procedures were approved by the North Carolina State University (NCSU) Institutional Animal Care and Use Committee (protocol #13-093-A).

Animal trial and selection of animals

An animal trial was conducted to gain insight on the effect of feeding toxic levels of endophyte infected forages. A total of 149 multiparous (parities 2 to 4) pregnant purebred Black Angus cows were used. Approximately half of the animals (78 cows) were located at the Upper Piedmont Research Station (UPRS – Reidsville, NC, NCSU), while the remaining animals (71 cows) were located at the Butner Beef Cattle Field Laboratory (BBCFL – Bahama, NC, NCSU).

Both groups had free access to forage and water during 13 weeks establishing a chronic exposure to toxic fescue (April to July 2016). Cattle at both locations grazed pastures known to be endophyte infected toxic tall fescue for the entirety of the study. Cattle were rotationally grazed every two weeks at each location among select pastures to continue adequate forage management as well as insure sufficient forage was continually available. Forage samples were collected every two weeks to evaluate nutrient quality and percentage of available forage that was fescue.

In addition, fescue tiller samples were collected in November of 2016 to evaluate pasture infection rate for the toxic endophyte. Fescue tiller samples were collected on a particular day, rinsed the following evening, and shipped on ice the following morning to determine pasture infection rate and the average infection rate is reported by experimental period (Agrinostics Ltd.)
Fescue samples from each pasture were sent to the University of Missouri Veterinary Medical Diagnostic Laboratory (Columbia, MO) to analyze the ergot alkaloid amounts present within the grass using HPLC as described by [22]. Fecal material was extracted from all cows following 13 weeks of exposure to endophyte-infected fescue. In brief, a lubricated shoulder length glove was inserted into the rectum and a grab sample of feces were collected from the colon. The fecal samples were labeled and placed immediately on ice and transported back to the lab for further processing. Fecal samples were transferred to labeled 15 ml polystyrene vials (BD Falcon) and stored at -80°C for analysis.

Out of the 149 cows enrolled in the trial, 40 cows showing extreme growth performance were selected for further analyses. For each animal, growth during the trial was estimated as the slope of regression analysis of BW on weeks (average weekly gain; AWG). Slopes (i.e. AWG) were estimated based on 3 window periods: weeks 1 through 13 (w1_13), weeks 1 through 7 (w1_7), and weeks 7 through 13 (w7_13) to assess the effect of increase in temperature from April to July, availability of forage and exposure of infected tall fescue. The AWG data for each of these scenarios were analyzed using the following model:

\[ AWG_{ijk} = \mu + L_i + P_j + b_1(iBW_k - \bar{iBW}) + e_{ijk} \]  

where \( AWG_{ijk} \) is the AWG of the cow; \( \mu \) is intercept; \( L_i \) is the fixed effect of the \( i^{th} \) location, \( P_j \) is the fixed effect of the \( j^{th} \) parity; \( b_1 \) is the partial regression coefficient for the covariate of initial BW (\( iBW \)); \( iBW_k \) is the \( iBW \) of the \( k^{th} \) cow; and \( e_{ijk} \) is the residual associated with \( y_{ijk} \), with \( e_{ijk} \sim N(0, I\sigma^2_e) \) where \( I \) is the identity matrix. Statistical analysis was performed in SAS 9.4 (Statistical Analysis System, Cary, NC, USA).
Identification of animals with high (HT) or low (LT) tolerance to FT were based on the residuals from Eq. 1. The top (positive) 20 and bottom (negative) 20 residuals, with equal representation from each location (i.e. 20 from each location), were classified as HT and LT, respectively, for a total of 40 selected animals. Fecal samples from these 40 animals were subjected to amplicon sequencing targeting bacteria and fungi (see below). The aim of this study was to compare the fecal microbiota of those animals that showed most extremes in their performance (based on AWG), to achieve a clearer biological signal. Thus, a non-treatment group (which was not exposed to toxic fescue) was not included in this trial. This analysis of identifying the “best” and “worst” performing animals was done for each of the three window periods, which resulted in different sets of selected animals, depending on the window period. In order to identify which of the three periods better expressed the impact of FT on performance, two additional analyses were performed. First, the residual variance ($\sigma^2_e$) of the data for each of the window periods were estimated with the model below, in order to identify the period in which greater variability of the data was observed, which is an indication of response to diseases [23]:

$$AWG_{ijk} = \mu + L_i + P_j + b_1(iBW_k - \bar{BW}) + e_{ijk}$$

[Eq. 2]

where $AWG_{ijk}$, $\mu$, $L_i$, $P_j$, $b_1$, $iBW_k$, and $e_{ijk}$ are as previously defined in Eq. 1. Analysis was performed in SAS 9.4. The estimated $\sigma^2_e$ of each window period (w1_7, w1_13, and w7_13) was compared between each other and tested using an F-test. In additional, the AWG residuals (AWG_res) of the selected animals based on Eq. 1 were analyzed with the following model:
\[ AWG_{res_{ijk}} = \mu + T_i + L_j + W_k + \text{interactions} + e_{ijk} \]  

where \( \mu \) and \( e_{ijk} \) are as previously defined; \( AWG_{res_{ijk}} \) is the AWG_res of the selected animal; \( T_i \) is the fixed effect of the \( i \)th tolerance group (HT or LT), \( L \) is the fixed effect of the \( j \)th location (BBFCL or UPRS); \( W_k \) is fixed effect of the \( k \)th window period (w1_7, w1_13, or w7_13); and \( \text{interactions} \) represent all possible interactions between these effects. There were no significant effects \((P \geq 0.159)\) for the main effects of \( W \) and \( L \), and for the interactions of \( T*L*W \), \( L*W \), and \( T*L \). There was a significant \((P<0.0001)\) interaction between \( T \) and \( W \), and the main effect of \( T \). Statistical analysis was performed in SAS 9.4.

The estimated \( \sigma^2_e \) for each window period and for each \( T \) by window period are presented in Table S1. The estimated \( \sigma^2_e \) for w1_7 [6.00 (kg/week)^2] was greater \((P<0.01)\) than for w1_13 [0.07 (kg/week)^2] and w7_13 [2.94 (kg/week)^2]. In addition, HT animals for w1_7 had the highest \((P<0.01)\) AWG_res (3.74 kg/week), whereas LT animals for w1_7 (-3.52 kg/week) had lowest \((P<0.01)\) AWG_res, and all of the other AWG_res were not different from each other \((P>0.01)\). Because of the greater estimated \( \sigma^2_e \) and the more extreme AWG_res values, data using w1_7 were used for subsequent analyses.

**DNA extraction**

Fecal material from the selected 40 HT and LT cows was thawed and genomic DNA was extracted from 0.25 grams of sample, using the Qiagen DNeasy Powerlyzer Powersoil kit following the instructions of the manufacturer. Mechanical cell lysis was performed using a Fischer Scientific Beadmill 24. DNA concentrations were determined using a Qubit 3 fluorometer (Invitrogen).
Sequencing and analysis

16S rRNA gene

Briefly, PCR amplicon libraries targeting the 16S rRNA gene present in extracted DNA were produced using a barcoded primer set adapted for Illumina MiSeq [24]. DNA sequence data was generated using Illumina MiSeq paired-end sequencing at the Environmental Sample Preparation and Sequencing Facility (ESPSF) at Argonne National Laboratory (Lemont, IL, USA).

Specifically, the V4 region of the 16S rRNA gene (515F-806R) was PCR amplified with region-specific primers that include sequencer adaptor sequences used in the Illumina MiSeq flowcell [24, 25]. The forward amplification primer also contains a twelve base barcode sequence that supports pooling of up to 2,167 different samples in each lane [24, 25]. Each 25 µL PCR reaction contained 9.5 µL of MO BIO PCR Water (Certified DNA-Free), 12.5 µL of QuantaBio’s AccuStart II PCR ToughMix (2x concentration, 1x final), 1 µL Golay barcode tagged forward primer (5 µM concentration, 200 pM final), 1 µL reverse primer (5 µM concentration, 200 pM final), and 1 µL of template DNA. The conditions for PCR were as follows: 94 °C for 3 minutes to denature the DNA, with 35 cycles at 94 °C for 45 s, 50 °C for 60 s, and 72 °C for 90 s; with a final extension of 10 min at 72 °C to ensure complete amplification. Amplicons were then quantified using PicoGreen (Invitrogen) and a plate reader (Infinite® 200 PRO, Tecan). Once quantified, volumes of each of the products were pooled into a single tube so that each amplicon was represented in equimolar amounts. This pool was then cleaned up using AMPure XP Beads (Beckman Coulter), and then quantified using a fluorometer (Qubit, Invitrogen). After quantification, the molarity of the pool was determined and diluted down to 2 nM, denatured,
and then diluted to a final concentration of 6.75 pM with a 10% PhiX spike for sequencing on
the Illumina MiSeq. Amplicons were sequenced on a 151bp MiSeq run using customized
sequencing primers and procedures [24].

Sequence analysis was done with the open source software mothur following the mothur MiSeq
were trimmed using a minimum average quality score of 35, with a sliding window size of 50 bp.
Chimeric sequences were removed with the “Chimera.uchime” command. For alignment, the
SILVA SSU NR reference database v128 [27] and for taxonomic classification the RDP trainset
(trainset16_022016) provided by the mothur website was used. After quality control, 58,400
sequences were randomly subsampled from each sample using mothur. The sequences were
clustered into operational taxonomic units (OTU) with a cutoff of 97% 16S rRNA gene
similarity (=0.03 distance).

**ITS1 region**

Library preparation and amplicon sequencing was performed using Illumina MiSeq sequencing
platform as with the 16S rRNA analysis. The ITS1 region was amplified using ITS1f-ITS2
primers designed to amplify fungal microbial eukaryotic lineages designed by the Earth
Microbiome Project [28]. This generated paired-end reads of 251bp. Sequence analysis was done
with mothur as for 16S rRNA genes. Barcode sequences, primers and low quality sequences
were trimmed using a minimum average quality score of 35, with a sliding window size of 50bp.
Sequences were aligned against themselves using the mothur command “pairwise.seqs”, and the
UNITEV6_sh_99 dataset (provided by mothur) was used to classify the sequences. After quality
control, 10,000 sequences were randomly subsampled from each sample using mothur. The
sequences were clustered into OTUs with a cutoff of 97% ITS1 region similarity (=0.03)
distance) following recent guidelines [29]. Additionally, representative sequences for each OTU
were further classified using NCBI BlastN.

Data availability
The 16S rRNA gene and ITS1 region sequences have been submitted to the NCBI Sequence
Read Archive SRA and are available under the BioProject ID PRJNA498290.

Statistical analysis
The amplicon sequencing data was analyzed with the same model described in Eq. 2. The
OTU count data were analyzed with a negative binomial model [30] including the same effects
in the model described in Eq. 2. In this analysis, the 50 most abundant OTUs for each dataset
(16S rRNA and ITS1) were analyzed. Preliminary analyses indicated statistical problems with
the ITS1 data because of the low counts for some OTUs. Therefore, OTUs with low counts (n =
4) in each location-tolerance group combination were removed. Means (for the diversity data)
and log2 fold-changes (log2FC; for the OTU count data) were separated using Tukey’s test for
the effects of location, tolerance group, and their interaction, when significant (P<0.05). In
addition to these analyses, a canonical discriminant analysis (CDA) was performed with the
objective of identifying OTUs (both 16S rRNA and ITS1 data) that could discriminate groups
with high power. Three CDA analyses were performed: to discriminate between tolerance groups
(2 groups), locations (2 groups), and between the 4 groups from the combination between
location and tolerance group. OTUs were selected using a stepwise approach, with an alpha of
0.15 to enter the model and an alpha of 0.05 to remain in it. In addition, a leave-one-out cross-
validation (LOOCV) was performed in order to assess the classification power of OTU. This analysis was done using the relative abundance data because of its more quantitative characteristics. All were performed in SAS 9.4.

Results

Ergot alkaloid concentrations per farm determined by HPLC

Overall, the percentage of fescue in the pastures was not significantly different between locations (68.1 and 64.3% at UPRS and BBCFL, respectively) throughout the grazing period. Furthermore, the fescue in these pastures were highly infected with toxin-producing endophyte (86.3 and 80.8 % at UPRS and BBCFL, respectively). Ergovaline levels were 1,110 µg/Kg and 1,900 µg/Kg were found at BBCFL and UPRS farms respectively (Table 1). The UPRS farm showed higher ergot alkaloid concentrations than the BBCFL farm, harboring in addition to Ergovaline also Ergosine, Ergotamine, Ergocornine, Ergocryptine and Ergocristine.
Table 1: Ergot alkaloid concentration of tall fescue pastures

<table>
<thead>
<tr>
<th>Ergot alkaloids</th>
<th>Concentration at farm BBCFL(^1) (µg/Kg)</th>
<th>Concentration at farm UPRS(^2) (µg/Kg)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Ergosine</td>
<td>0</td>
<td>1,000</td>
</tr>
<tr>
<td>Ergotamine</td>
<td>0</td>
<td>525</td>
</tr>
<tr>
<td>Ergocornine</td>
<td>0</td>
<td>160</td>
</tr>
<tr>
<td>Ergocryptine</td>
<td>0</td>
<td>450</td>
</tr>
<tr>
<td>Ergocristine</td>
<td>0</td>
<td>145</td>
</tr>
<tr>
<td>Ergovaline</td>
<td>1,110</td>
<td>1,900</td>
</tr>
<tr>
<td>Total</td>
<td>1,110</td>
<td>4,180</td>
</tr>
</tbody>
</table>

\(^1\)BBCFL: Butner Beef Cattle Field Laboratory, Bahama NC
\(^2\)UPRS: Upper Piedmont Research Station, Reidsville NC

Composition of fecal bacterial microbial communities

Overall, 5,320 OTUs were generated after quality control, subsampling and removal of OTUs representing less than ten sequences from the original 4.675 million sequence reads, of which 3.983 million reads (85%) remained after quality control. Most of the reads were bacterial, 0.8% of all reads were classified as *Archaea*. Twenty-three phyla were identified with *Firmicutes* (58.7-67.9%), *Bacteroidetes* (19.7-25.7%), *Proteobacteria* (1.1-12.8%), *Actinobacteria* (1.1-3.7%) and unclassified bacteria (5.2-7.7%) being most abundant (Fig. S1). All other phyla showed relative abundances of less than 1%.

The most abundant OTU (OTU 1, comprising 7.6% of all reads) affiliated to the *Ruminococcaceae* UCG-005 group, OTU 2 to *Solibacillus* (99.6% similarity to *Solibacillus silvestris*, 6.1% overall relative abundance), OTU 3 to *Acinetobacter* (100% similarity to...
*Acinetobacter lwoffii*, 4.7% overall relative abundance), OTU 4 to *Bacillus* (99.2% similarity to *Bacillus psychrosaccharolyticus*, 3.2% overall abundance) and OTU 5 to *Monoglobus* (93.7% similarity to *Monoglobus pectinilyticus*, 2.3% overall abundance) (Table S2). Among the most abundant OTUs, a number of OTUs were classified into the same genus such as OTUs 5, 12, 47 (*Monoglobus*); 8, 13, 28, 35 (*Bacteroides*); 11, 15, 40 (*Clostridium*); 9, 14 (*Lysinibacillus*); 2, 46 (*Solibacillus*).

**Composition of fecal fungal microbial communities**

Overall, 1,000 OTUs were generated after quality control, subsampling and removal of OTUs representing less than 10 sequences from the original 7.051 million sequence reads, of which 390,000 reads remained after quality control and subsampling.

OTU 1, OTU 5 and OTU 12 affiliated to *Microsphaeropsis* (Montagnulaceae family), OTU 2 and OTU 3 to *Thelebolus* (Thelebolaceae family), OTU 4 to the Pleosporaceae family, OTUs 14, 22, 34, 35 to *Orpinomyces* and OTUs 25, 26, 31, 45 to *Caecomyces*; both of these genera belong in the family Neocallimastigaceae (Table S3). No sequences were attributed to *Epichloë coenophiala*, the endophyte believed to be responsible for FT, by either the classification against the UNITE database or manual BlastN of representative sequences of each OTU against NCBI nr.

**Alpha diversity of HT and LT cattle fecal microbial communities**

When comparing HT and LT cattle bacterial microbial communities, we observed significant decreases in diversity (Shannon, $P<0.001$) and species richness (Chao, $P=0.0078$; ACE, $P=0.0093$) and an increase in evenness (Simpson, $P=0.005$) in the LT cattle for both sites. For
tolerance group and T*L, we observed a significant difference for species richness estimators
only (Chao, \( P<0.019 \); ACE, \( P<0.025 \)) (Table 2).

Similar to the 16S rRNA gene analysis, fungal microbial community analysis revealed decreased
diversity (Shannon, \( P=0.001 \)) and increased evenness (Simpson, \( P=0.0047 \)) in LT cattle from
both sites considering the tolerance group effect. No change was observed in species richness
between HT and LT cattle when comparing fungal microbial communities (Table 3). The fungal
microbial communities showed higher evenness values than those of the bacterial communities
indicating a less even distribution of the fungal OTUs.
Table 2: Bacterial species richness and diversity estimators in fecal microbial communities across location\(^1\) and tolerance groups\(^2\)

<table>
<thead>
<tr>
<th>Diversity Parameter</th>
<th>BBCFL</th>
<th>UPRS</th>
<th>P-value(^3)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>HT</td>
<td>LT</td>
<td>T</td>
</tr>
<tr>
<td>Ace (richness)</td>
<td>9473.9(^b)</td>
<td>9245.1(^b)</td>
<td>12074(^a)</td>
</tr>
<tr>
<td></td>
<td>(557.9)</td>
<td>(557.9)</td>
<td>(557.9)</td>
</tr>
<tr>
<td>Chao (richness)</td>
<td>6620.9(^b)</td>
<td>6506.2(^b)</td>
<td>8127.3(^a)</td>
</tr>
<tr>
<td></td>
<td>(318.7)</td>
<td>(318.7)</td>
<td>(318.7)</td>
</tr>
<tr>
<td>Npshannon (diversity)</td>
<td>5.80(^a)</td>
<td>5.29(^b)</td>
<td>5.98(^a)</td>
</tr>
<tr>
<td></td>
<td>(0.16)</td>
<td>(0.16)</td>
<td>(0.16)</td>
</tr>
<tr>
<td>Shannon (diversity)</td>
<td>5.71(^a)</td>
<td>5.20(^b)</td>
<td>5.90(^a)</td>
</tr>
<tr>
<td></td>
<td>(0.17)</td>
<td>(0.17)</td>
<td>(0.17)</td>
</tr>
<tr>
<td>Simpson (evenness)</td>
<td>0.018(^bc)</td>
<td>0.057(^ab)</td>
<td>0.017(^c)</td>
</tr>
<tr>
<td></td>
<td>(0.014)</td>
<td>(0.014)</td>
<td>(0.015)</td>
</tr>
</tbody>
</table>

\(^1\)BBFLC, Butner Beef Cattle Field Laboratory (Bahama, NC, USA); UPRS, Upper Piedmont Research Station (UPRS; Piedmont, NC, USA);
\(^2\)HT, High Tolerance; LT, Low Tolerance;
\(^3\)P, Tolerance group (HT or LT); L, Location (BBCFL or UPRS); T*L, interaction between T and L;
\(^a,b,c\) Least-squares means of alpha diversity values lacking common superscripts are statistically different (\(P<0.05\)) based on Tukey’s test;
Numbers within parentheses represent standard error measurements.
Table 3: Fungal species richness and diversity estimators for fecal microbial communities across location\(^1\) and tolerance groups\(^2\)

<table>
<thead>
<tr>
<th>Diversity Parameter</th>
<th>BBCFL</th>
<th>UPRS</th>
<th>(P)-value(^3)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Ace (richness)</td>
<td>4970.96(^b)</td>
<td>7282.02(^ab)</td>
<td>9697.22(^a)</td>
</tr>
<tr>
<td></td>
<td>(1405.16)</td>
<td>(1405.16)</td>
<td>(1405.16)</td>
</tr>
<tr>
<td>Chao (richness)</td>
<td>2732.72(^b)</td>
<td>3270.42(^b)</td>
<td>5256.33(^a)</td>
</tr>
<tr>
<td></td>
<td>(544.6)</td>
<td>(544.6)</td>
<td>(544.6)</td>
</tr>
<tr>
<td>Npshannon (diversity)</td>
<td>3.27(^ab)</td>
<td>2.86(^b)</td>
<td>4.05(^a)</td>
</tr>
<tr>
<td></td>
<td>(0.28)</td>
<td>(0.28)</td>
<td>(0.28)</td>
</tr>
<tr>
<td>Shannon (diversity)</td>
<td>3.1(^ab)</td>
<td>2.69(^bc)</td>
<td>3.82(^a)</td>
</tr>
<tr>
<td></td>
<td>(0.27)</td>
<td>(0.27)</td>
<td>(0.27)</td>
</tr>
<tr>
<td>Simpson (evenness)</td>
<td>0.151(^b)</td>
<td>0.272(^ab)</td>
<td>0.132(^b)</td>
</tr>
<tr>
<td></td>
<td>(0.06)</td>
<td>(0.06)</td>
<td>(0.06)</td>
</tr>
</tbody>
</table>

\(^1\)BBCFL, Butner Beef Cattle Field Laboratory (Bahama, NC, USA); UPRS, Upper Piedmont Research Station (UPRS; Piedmont, NC, USA);
\(^2\)HT, High Tolerance; LT, Low Tolerance;
\(^3\)T, Tolerance group (HT or LT); L, Location (BBCFL or UPRS); L*T, interaction between T and L;
\(a,b,c\) Significant differences in alpha diversity values between diversity parameter and effect (T, L, T*L) are designated by lowercase letters (\(P<0.05\))

Numbers within parentheses represent standard error measurements.
Differentially abundant OTUs between HT and LT cattle

Among the 50 most abundant OTUs, statistically significant ($P<0.05$) differences between cattle were seen for 12 bacterial and eight fungal OTUs considering only the effect of location for comparison (Fig. 1).

Considering the effect of tolerance group, only bacterial OTU 3 (Acinetobacter) was significantly ($P<0.0001$) higher in HT cattle and eleven fungal OTUs were significantly ($P<0.05$) different between HT and LT cattle (Fig. 2). Fungal OTUs 2, 3, 13 (all three classified as Thelebolus), and 24 (Coprinopsis) were more abundant in LT cattle, whereas fungal OTUs 1 (Microsphaeropsis), 14 and 22 (Orpinomyces), 20 (Pyrenochaetopsis), 26 and 31 (Caecomyces), and 49 (Pleosporales) were more abundant in HT cattle.

When considering interactions between location and tolerance group for the 16S rRNA, OTUs 6, 11, 15, 18, 19, 20, 26, 27, 38, and 45 showed significantly ($P<0.05$) different interactions, although with opposite trends for each location (Fig. 3). For the fungal OTUs, eleven OTUs were significantly ($P<0.05$) different, out of which six (OTUs 4 (Pleosporales), 5 (Microsphaeropsis), 6 (Psilocybe), 12 (Microsphaeropsis), 27 (Deconica), and 38 (Pleosporales)) were more abundant in the HT cattle and two (OTU 10 and 17; both Thelebolus) were more abundant in the LT cattle at both farms.

**Fig. 1.** Statistically significantly ($P\leq0.05$) different abundant bacterial and fungal OTUs considering location (BBCFL, UPRS) as an effect. The differences in abundance of OTUs are shown as log2 fold changes for each location. Positive values represent higher abundance in farm BBCFL, negative values represent higher abundance in farm UPRS. Error bars represent the 95% confidence interval.
Fig. 2. Statistically significantly ($P \leq 0.05$) different abundant bacterial and fungal OTUs considering tolerance group (HT, LT) as an effect. The differences in abundance of OTUs are shown as log2 fold changes for each tolerance group. Positive values represent higher abundance in HT cattle, negative values represent higher abundance in LT cattle. Error bars represent the 95\% confidence interval.

Fig. 3. Statistically significantly ($P \leq 0.05$) different abundant bacterial and fungal OTUs considering interactions of location (BBCFL, UPRS) and tolerance group (HT, LT) as an effect. The differences in abundance of bacterial (A) and fungal (B) OTUs are shown as log2 fold changes for each interaction. Positive values represent higher abundance in HT cattle, negative values represent higher abundance in LT cattle. Bars in blue and purple represent farms BBCFL and UPRS, respectively. Error bars represent the 95\% confidence interval.

Community-level comparison of microbial communities using canonical discriminant analysis

Results of the CDA analyses are presented in Table 4 and depicted in Fig. 4 and Fig. S2. There were 8, 19, and 14 OTUs selected for the analyses of T, L, and T*L, respectively, with 4 OTUs overlapping between these (bacterial OTUs 19 (Rikenellaceae RC9 gut group) and 21 (unclassified Ruminococcaceae) and fungal OTUs 1 (Microsphaeropsis) and 6 (Psilocybe). The discrimination of groups based on CDA was significant ($P < 0.001$) for all canonical variables (CAN) for both analyses. The squared canonical correlations ($R^2$) were: 94.7\% for T, 99.26\% for L, and 98.4\% (CAN1), 92.5\% (CAN2), and 73.0\% (CAN3) for L*T. For L*T, the proportion of
the total variation explained by each CAN was 80.2% (CAN1), 16.2% (CAN2), and 3.6%
(CAN3). For T, the two OTUs showing the most discriminative power were fungal OTU1
(Microsphaeropsis) and bacterial OTU21 (unclassified Ruminococcaceae), with standardized
canonical coefficients (SCC) of -4.0 and 3.1, respectively. For L, these were (SCC in
parentheses): bacterial OTU1 (Ruminococcaceae UCG-005, -5.3) and OTU2 (Solibacillus, 5.2).
For T*L, these were: fungal OTU6 (Psilocybe, 5.4) and fungal OTU8 (Thelebolus, 2.2) for
CAN1, fungal OTU25 (Caecomyces, -3.0) and OTU1 (Microsphaeropsis, 2.9) for CAN2, and
fungal OTU25 (Caecomyces, 1.5) and bacterial OTU21 (unclassified Ruminococcaceae, 1.4) for
CAN3. The misclassification rates for the CDA of T, L, and T*L were 2.6%, 0%, and 5.3%,
respectively.
Table 4. Canonical discriminant analysis for tolerance group\(^1\) (T), Location\(^2\) (L), and interaction between T and L (T*L)

<table>
<thead>
<tr>
<th>Target</th>
<th>OTU</th>
<th>Classification</th>
<th>Standardized Canonical Coefficients</th>
</tr>
</thead>
<tbody>
<tr>
<td>16S rRNA gene</td>
<td>OTU01</td>
<td><em>Ruminococcaceae</em> UCG-005</td>
<td>CAN1</td>
</tr>
<tr>
<td></td>
<td>OTU02</td>
<td><em>Solibacillus silvestris</em></td>
<td>CAN1</td>
</tr>
<tr>
<td></td>
<td>OTU05</td>
<td><em>Monoglobus pectinilyticus</em></td>
<td>CAN1</td>
</tr>
<tr>
<td></td>
<td>OTU10</td>
<td><em>Rikenellaceae</em> RC9 gut group</td>
<td>CAN1</td>
</tr>
<tr>
<td></td>
<td>OTU12</td>
<td><em>Monoglobus pectinilyticus</em></td>
<td>CAN1</td>
</tr>
<tr>
<td></td>
<td>OTU13</td>
<td><em>Bacteroides plebeius</em></td>
<td>CAN1</td>
</tr>
<tr>
<td></td>
<td>OTU15</td>
<td><em>Clostridium difficile</em></td>
<td>CAN1</td>
</tr>
<tr>
<td></td>
<td>OTU19</td>
<td><em>Rikenellaceae</em> RC9 gut group</td>
<td>CAN1</td>
</tr>
<tr>
<td></td>
<td>OTU21</td>
<td>unclassified</td>
<td>CAN1</td>
</tr>
<tr>
<td></td>
<td>OTU22</td>
<td><em>Psychrobacillus psychrodurans</em></td>
<td>CAN1</td>
</tr>
<tr>
<td></td>
<td>OTU25</td>
<td><em>Corynebacterium kutscheri</em></td>
<td>CAN1</td>
</tr>
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<td>OTU30</td>
<td><em>Christensenellaceae</em> R7 group</td>
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<td>OTU31</td>
<td>unclassified <em>Bacteroidetes</em></td>
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<td>OTU39</td>
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<td><em>Mollicutes RF39</em></td>
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<td><em>Microsphaeropsis arundinis</em></td>
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<td><em>Thelebolus</em></td>
<td>CAN1</td>
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<td>OTU12</td>
<td><em>Microsphaeropsis</em></td>
<td>CAN1</td>
</tr>
<tr>
<td></td>
<td>OTU13</td>
<td><em>Thelebolus</em></td>
<td>CAN1</td>
</tr>
<tr>
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<td>OTU19</td>
<td><em>Phaeosphaeriopsis</em></td>
<td>CAN1</td>
</tr>
<tr>
<td></td>
<td>OTU24</td>
<td><em>Coprinopsis colthurnata</em></td>
<td>CAN1</td>
</tr>
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<td></td>
<td>OTU25</td>
<td><em>Caecomyces</em></td>
<td>CAN1</td>
</tr>
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<td></td>
<td>OTU31</td>
<td><em>Caecomyces</em></td>
<td>CAN1</td>
</tr>
<tr>
<td></td>
<td>OTU34</td>
<td><em>Orpinomyces</em></td>
<td>CAN1</td>
</tr>
</tbody>
</table>

\(P\)-value <0.001 <0.001 <0.001 <0.001 <0.001

\(R^2\) (%) 94.7 99.3 98.4 92.5 73.0

\(^1\) BBCFL, Butner Beef Cattle Field Laboratory (Bahama, NC, USA); UPRS, Upper Piedmont Research Station

\(^2\) HT, High Tolerance; LT, Low Tolerance;

CAN1-CAN3, canonical variables 1-3. The number of CAN in each analysis depends on the number of levels of the discriminated group (i.e. \(n-1\));
Canonical discriminant analysis (CDA) for response to fescue toxicosis. CDA was performed to discriminate animals based on the combination between tolerance group (T; High [HT] and Low [LT] tolerance groups) and location (BBCFL and UPRS) using 14 fungal and bacterial OTUs. Each point represents the canonical score (CS) of each animal based on the respective canonical variable (CAN). The x-axis represents CAN 1 and the y-axis CAN 2. Red and Blue points represent HT and LT animals, respectively, whereas triangles and stars represent animals from UPRS and BBCFL, respectively.

Discussion

In general, our knowledge about FT has significantly advanced during recent years [1, 2, 4, 17, 31-35]. However, the knowledge about a possible involvement of GI tract microbial communities, especially fungal communities, in FT is still highly limited. Recently, Mote et al. found in animals fed toxic fescue, relative abundances of the bacterial families Ruminoccocaceae and Lachnospiraceae were significantly increased [17]. Our study predominantly found consistent changes in the fungal OTUs and only detected a single significantly different bacterial OTU (OTU3, Acinetobacter) within the 50 most abundant OTUs between the tolerance groups, and this OTU was not related to either of the aforementioned families (Fig. 2). It should be noted that the study by Mote et al., used Angus steers, not cows, and was performed at a different location (in Georgia, USA) [17]. Thus, the comparability of these two studies might be limited. Particularly, keeping in mind the strong differences in microbiota between farms found here. Nevertheless, similar to our study, Mote et al. also identified major changes of fecal the microbial communities in response to FT [17]. Another study has shown degradation of fescue
alkaloids by rumen microorganisms without identifying the microbes responsible for the
degradation [19]. Tryptophan-utilizing rumen bacteria can be capable of ergovaline degradation
as shown for a *Clostridium sporogenes*, other *Clostridium* species [36], and a *Prevotella bryantii*
isolate [18]. We did not find *Clostridium sporogenes* or *Prevotella bryantii* OTUs in our dataset;
however, a number of (abundant) *Clostridium* OTUs such as OTUs 11 and 15 (both of them
were more abundant in the high tolerant cattle at the BBCFL farm considering T*L*) were
detected. These OTUs could potentially harbor tryptophan degrading capabilities, possibly
enabling them to degrade ergot alkaloids. Another explanation for the absence of *Prevotella
bryantii* and *Clostridium sporogenes*, which are abundant rumen bacteria, in our samples might
be that we have used fecal (and not rumen) samples and rumen bacteria might not be well
represented in fecal samples. Additionally, a *Rhodococcus erythropolis* strain has recently been
described to degrade ergot alkaloids [21]. We found only very few and extremely low abundant
*Rhodococcus* OTUs in our dataset; based on the reported strain-specific ergot alkaloid
degradation capability of *Rhodococcus* [21], we assume that *Rhodococcus* species are not
involved in ergot alkaloid degradation in the animals analyzed here.

We chose to study the fecal microbiota as fecal samples allow for periodic observations of
microbiota with large sample sizes in a non-invasive way. One major limitation of using fecal
samples to study GI tract microbiota is the fact that the fecal microbiota primarily reflects
luminal microbiota, which can be significantly different from GI tract mucosal microbiota.
Moreover, fecal samples are more representative of the digesta in the lower GI tract and do not
necessarily adequately represent rumen microbial communities. It should thus be noted that the
findings found here using fecal samples might not reflect changes of the microbiota of e.g. the
rumin. In the future, an investigation of microbiota changes in response to toxic fescue feeding should be performed using rumen samples as FT is considered a rumen metabolic disease. 16S rRNA gene amplicon sequencing revealed highly different microbial communities in the HT and LT cattle at both farms analyzed. The shifts in microbial community composition were more pronounced at farm UPRS, where the animals were fed a diet containing higher ergovaline levels and more diverse ergot alkaloids, suggesting a tolerance-by-location interaction (T*E), which is consistent with [37] (Table 1). This suggests that the amount of alkaloid toxins in the diet determines not only the severity of FT but also changes in microbial community composition. In the LT cattle, species richness and diversity were lower than in the HT cattle. These results indicate a shift towards less diverse microbial communities in the LT cattle GI tract microbiota in response to a toxic fescue diet. Whether this reduced diversity is a sign of a functional dysbiosis – an imbalanced microbiota – remains to be tested in future experiments. Again, the changes in species richness and diversity were more prominent in farm UPRS with the higher ergot alkaloid levels. Similarly, on a community level, LT and HT cattle fecal microbial communities showed a significantly different composition as highlighted by the CDA analysis which revealed a clear clustering of fecal communities with respect to location and tolerance group. Also, on OTU-level, significant differences between fecal microbiota were revealed in our study for location as well as for tolerance group.

Twelve bacterial OTUs and eight fungal OTUs were significantly different when considering only location. Bacterial OTUs 3 (Acinetobacter), 9 (Lysinibacillus), 10, 33, and 48 (Rikenellaceae RC9 gut group), 29 (Sporobacter), 43 (Christensenellaceae), 47 (Monoglobus), and 50 (Molllicutes) were more abundant at farm UPRS and 21 (unclassified Ruminococcaceae), 25 (Corynebacterium), and 36 (Arthrobacter) were more abundant at farm BBCFL. Fungal
OTUs 3 and 7 (*Thelebolus*) were more abundant in UPRS and 1 (*Microsphaeropsis*), 14 and 34
(*Orpinomyces*), 25 and 26 (*Caecomyces*), and 49 (*Pleosporales*) were more abundant at BBCFL.
These differences in abundance may be caused by the different levels of toxins or different
management and feeding strategies and feed composition between the two farms.

For tolerance group, only one bacterial OTU (OTU 3, *Acinetobacter*) was significantly different
and higher in HT cattle. *Acinetobacter* species have been found in ruminant GI tracts [38]. It is
currently unclear whether these *Acinetobacter* phylotypes may be opportunistic pathogens or are
part of the physiological GI tract microbiota. In contrast, eleven fungal OTUs were significantly
different between HT and LT cattle when considering tolerance group. Fungal OTUs 1
(*Microsphaeropsis*), 14 (*Orpinomyces*), 20 (*Pyrenochaetopsis*), 22 (*Orpinomyces*), 26 and 31
(both classified as *Caecomyces*), and 49 (*Pleosporales*) were found to be significantly more
abundant in HT cattle, whereas OTUs 2, 3, 13 (all three classified as *Thelebolus*), and 24
(*Coprinopsis*) were significantly more abundant in LT cattle.

When considering T*L interactions, for the 16S rRNA data, differing results in abundance for
specific bacterial OTUs were observed for the two farms, suggesting that different bacteria may
be associated with the different response to FT at each of the farms. These bacteria may derive
from the different environments at the two farms, possibly resulting from different feeding and
management strategies at both farms. This could suggest that bacteria may not be of key
relevance for different response to FT. For the fungal data, more coherent results were found for
the interactions between T*L: A number of fungal OTUs were consistently more abundant in HT
cattle at both farms: OTUs 4 (*Pleosporales*), 5 (*Microsphaeropsis*), 6 (*Psilocybe*), 12
(*Microsphaeropsis*), 27 (*Deconica*), and 38 (*Pleosporales*), and other fungal OTUs were more
abundant in the LT cattle at both farms (OTU 10 and 17, both classified as *Thelebolus*).
Some of the abundant fungal OTUs could not be classified to genus level and it is thus hard to speculate about a possible function of these fungal phylotypes. Nevertheless, we assume that OTUs which are more abundant in the HT cattle, could potentially contribute to the better performance indicated by their weight gain observed in the HT cattle. Conversely, the OTUs found in LT cattle may be associated with more severe negative effects of FT. The fact that we observed consistent changes in the abundant fungal OTUs which were higher in HT cattle at both farms, and that their abundance was consistently higher at the UPRS farm (which is characterized by higher ergot alkaloid toxin levels in the diet) suggests that these fungi may be involved in mitigating the effects of FT in those animals. In spite of the differences caused by performing our animal trial at two different locations, the observation of similar and consistent changes in abundance of certain fungal phylotypes in response to FT, provides more support to the hypothesis that these phylotypes could potentially be positively associated with the higher AWG in HT cattle in our study. It should be noted that this potential beneficial role of fungi in FT tolerance needs to be investigated in more detail in future studies.

Anaerobic fungi of the phylum Neocallimastigomycota are effective fiber degrading organisms in the herbivore gut and have been reported to improve feed intake, feed digestibility, feed efficiency, and daily weight gain and milk production [39, 40]. Of the 50 most abundant OTUs, OTUs 14, 22, 25, 26, 31, 34, 35, and 45 were classified as members of the Neocallimastigomycota. Of these, OTUs 14, 22, 26 and 31 were found to be more prevalent in HT cattle. Although we provide no functional data here, this result adds to the accepted consensus these fungi positively contribute to the ruminant system and underlines the importance of gaining functional data in future research to increase efficiency and overall health in livestock species.
The genus *Thelebolus* was attributed to 11 of the 50 most abundant fungal OTUs. Of these OTUs, OTUs 2, 3, 10, 13, and 17 were found to be significantly more abundant in LT cattle, suggesting a negative effect of *Thelebolus* on ruminant health and performance. Members of the genus *Thelebolus* have been found in ruminant samples before [41]. Although there is little published data on this genus, some recent publications have suggested species within this genus can produce a cytotoxic exopolysaccharide designated as Thelebolan [42] and has been recently tested for its apoptotic effect on cancer cells [43]. It is unknown whether this compound contributes to the negative effects in LT cattle during chronic exposure to toxic fescue, and what conditions allow for increased abundance of this genus. It may be possible that the exopolysaccharide has an apoptotic effect on healthy cells in the GI tract, therefore limiting the absorption of nutrients and reducing integumentary strength within the gut.

OTUs attributed to unclassified Pleosporales were identified within our samples consistent with a recent study that found Pleosporales in the ruminant GI tract [41]. Members of the order Pleosporales including *Microsphaeropsis* have been identified as saprobic, endophytic and pathogenic fungi, and they are often are present within animal dung [44]. It is noteworthy that of the 16 OTUs assigned to the order *Pleosporales* within the 50 most abundant fungal OTUs, six OTUs (1, 5, 12, 20, 38, and 49) were found to be significantly more abundant in HT cattle whereas none were found to be more abundant in LT cattle suggesting a potential – although yet to be verified - beneficial role of these Pleosporales phylotypes during a FT challenge. Similar to the anaerobic fungi of the phylum Neocallimastigomycota, it is important to identify possible functional traits in these microorganisms that could be associated with the positive health of these animals with additional research.
Interestingly, we did not find any fungal OTUs related to *Epichloë coenophiala*. This may either be explained by the absence of *Epichloë coenophiala* in the samples sequenced here, or that *Epichloë coenophiala* might not be targeted by the primers used for ITS1 amplicon sequencing, or a different, yet unidentified, fungus might be responsible for the alkaloid production resulting in FT in our study. It is also conceivable that the aerobic nature of *Epichloë* as an endophyte prevents its growth under anaerobic conditions of the mammalian GI tract. In addition, fecal samples were collected in the end of the trial, when endophyte infection levels were lower in the forage, which could contribute to this lack of identification of this fungus.

It may be the case that certain fungal species degrade the alkaloid causing FT, thus decreasing the effect of FT directly. Another explanation may be the cellulolytic and fiber-degrading capabilities of fungi. Fungal phylotypes which are more abundant in HT cattle may be producing higher amounts of absorbable nutrients, thereby compensating for the negative impact of FT on GI tract systems. Many fungi are known to produce bioactive antibacterial compounds and could influence microbial community composition and, indirectly, an animals’ response to FT. Members of the genus *Microsphaeropsis*, attributed to three of the 50 most abundant fungal OTUs, including OTU1, are known to produce antimicrobial compounds [45], which could potentially alter microbial community composition.

**Conclusion**

This study compared the fecal fungal and bacterial communities of Angus cattle that exhibited contrasting tolerance to fescue toxicosis. Both microbial communities were significantly distinct between the HT and LT cattle after chronic exposure to toxic fescue, and may contribute to the animals’ physiological response to FT. HT cattle had more even and diverse fecal microbial
610 communities, suggesting LT cattle may be experiencing dysbiosis. Cattle with higher tolerance
611 to FT were associated with higher abundances of anaerobic fungi of the phylum
612 Neocallimastigomycota known to breakdown cellulose, and uncharacterized members of the
613 Pleosporales order. Cattle with lower tolerance to FT were found to have higher abundance of
614 phylotypes within the Thelebolus genus. This shift in the GI microbiota was more evident at the
615 UPRS location characterized by higher levels of infected fescue, suggesting a tolerance-by-
616 location interaction. In addition, this was the first study to analyze fungal communities
617 associated with contrasting growing performance under FT in cattle. To better understand the
618 contribution of the microbiota, particularly of fungi, to mitigate FT, functional data using rumen
619 samples will be needed in the future. The availability of such data might allow identifying
620 additional ways to mitigate the negative impact of FT on grazing livestock.
621
622 Acknowledgements
623 This work was supported by North Carolina Cattlemen’s association and North Carolina
624 Agricultural foundation.
625
626
References


14. Looper ML, Reiter ST, Williamson BC, Sales MA, Hallford DM, Rosenkrans JCF. Effects of body condition on measures of intramuscular and rump fat, endocrine factors, and


Supporting information captions

Table S1: Residual variance ($\sigma_e^2$) for each window period (WP), and AWG_res2 means for each WP by genetic group (GG).

Table S2: The 50 most abundant 16S rRNA gene OTUs

Table S3: The 50 most abundant ITS1 OTUs

Figure S1: Mean relative abundance of the five most abundant bacterial phyla across all sample sites and groups. The error bars represent SEM.

Figure S2: Canonical discriminant analysis (CDA) for response to fescue toxicosis (FT). For A, CDA was performed to discriminate animals based on location group (L): BBCFL and UPRS can be found in blue and gold, respectively. For A, CDA was performed to discriminate animals based on tolerance group (T): High (HT) and Low (LT) tolerance to FT can be found in green and red, respectively. For A, the x-axis represents the CS for CAN 1 and y-axis represent the density of the CS data.
Figure 1

The figure shows a bar chart with Log2 Fold Changes on the y-axis and various bacterial OTUs on the x-axis. Two groups are compared: 16S rRNA in red and ITS1 in green. The groups are labeled as BBCFL and UPRS. The chart highlights significant changes in bacterial abundance for different OTUs across the two groups.
Figure 2

The figure shows a bar graph with log 2 fold changes on the y-axis and various OTUs on the x-axis. The graph compares 16S rRNA and ITS1 sequences. The bars are color-coded, with red representing 16S rRNA and green representing ITS1. The graph includes OTUs labeled as Acinetobacter, Thelebolus, and others, with log 2 fold changes ranging from -10 to 10.