The gut bacterial diversity of sheep associated with different breeds in Qinghai province

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Abstract
Background: Gut microbiota play important roles in their co-evolution with mammals. However, little is understood about gut bacterial community of Tibetan sheep compared with other sheep breeds. In this study, we investigated the gut bacterial community in 4 different sheep breeds living in the Qinghai-Tibetan Plateau (QTP) of China using high-throughput sequencing (HTS) technique.

Results: The results suggested that bacterial community abundance and breeds diversity of Tibetan sheep (TS) were significantly lower than that of the other three breeds of sheep [Dorset sheep (DrS), Dorper sheep (DrS) and Small Tail Han sheep (STHS)] (p < 0.05). Principal coordinates analysis (PCoA) and nonmetric multidimensional scaling (NMDS) analysis indicated that microbiome composition of TS was significantly different from that of other three sheep breeds (p < 0.01). Firmicutes was the most predominant microbial phylum in the gut, followed by Bacteroidetes. The gut bacterial community of TS showed higher proportions of phylum Spirochaetes, Proteobacteria and Verrucomicrobia, compared to the other three sheep breeds, but the Deferrribacteres was lacked in TS. At the genus level, Treponema, Succinivibrio, 5-7N15 and Prevotella showed significantly higher abundance in TS than in the other three sheep breeds (p < 0.05).

Conclusions: In this study, we first employed HTS to fully understand the gut microbiomes among different sheep breeds in QTP of China.

Background
The significance of gut microbiome is well known, it is an extremely complicated and diverse population and has been explored extensively [1, 2]. Recently, a new term ‘superorganism’ is applied to describe the strong tie between the gut commensal and its reliable host [3, 4]. The intestinal microbiota is linked to a diverse range of conditions, including gathering energy, promoting intestinal epithelial cell proliferation and enhancing the immune system [5, 6], and can be deservedly regarded as an ‘organ’ playing a significant part in the metabolic process [3, 7, 8]. It has been found that more than 3.3 million commensal genes are resided in humans, which is equivalent to 150 times of human genes [9]. In ruminants, the composition of the gastrointestinal commensals, their effect on host immunity and welfare have been explored for several years. A previous study suggested that it was a
great achievement of using gut microbiota by ruminant animals [10]. And it’s true that combining high-throughput ‘omics’ technologies with ruminant’s genomes, the unprecedented well-being of achievement can be harvested [3].

Initially, the researches about gut microorganisms were dependent on the sequence alignment from genomic libraries to screen the functional genes, or through PCR amplification [3]. In *Escherichia coli*, heterologous gene expression was determined by Sanger Chain Termination Method and obtained the autoradiographic map [11]. However, it could only obtain the genes expressed in E. coli, and certainly leads to a large missing of available genes. Through this method, the first gene was acquired from *F. succinogenes* and encoding cellulases [12]. In the following years, seven *F. succinogenes* genes were detected through these traditional genetic methods, which encoded fiber-degrading enzymes [13].

This was certainly an extraordinary progress, but after *F. succinogenes* S85 genomic sequencing was finished, it was found that there were 104 open reading frames to participate in the disruption of plant cell wall [14]. Accordingly, the great advantage of genomic sequencing is intuitively clear, more enzymes were detected than that of previous researches in the *F. succinogenes* genome.

Metagenomics is performed to analyze the genome of microbial communities in an environmental sample, including the genomic sequence-based analysis and functional prediction. It is applied to screening the specific functions and detecting new bioactives in diverse ecosystems [15]. Moreover, it is a vital step to model and connect the microbial structure and function to that of the host [16, 17]. Some researchers found that the gut microbiota of mammals have a large identical part of their functions, suggesting that the understanding of human researches can provide many common views for ruminants [18]. In addition, research has shown that there is a relationship between microbial lineages and their specific environment [19]. However, those gut microbes are not frequently emerged in other environments [20]. Some study also showed that gut microbes were extensively shared among various mammals [21], indicating that some views are commonly applied to both humans and domestic animals.

In the present study, to better analyze and clarify the relationship between microbial lineages and the host breeds from a perspective of gut microbiota, amplification of V3 + V4 region of the 16s rDNA was
performed, which was followed by, the most credible techniques, Illumina MiSeq Reagent Kit PE250 sequencing. Apart from the shared characteristics are presented among all sheep, typical microbial population features corresponding to different sheep breeds are also mentioned in our research.

Results

Description of the sequencing data

The samples we obtained were from the feces of a great variety of sheep, which were comprised 40 subjects, namely 10 Dorset sheep (DrS), 15 Small Tail Han sheep (STHS), 5 Tibetan sheep (TS) and 10 Dorper sheep (DrS). We received 1,694,264 raw bases on a PE250 instrument, as mentioned in the experiment method. Behind quality-filtering (also as described in the methods), 1,359,405 total sequences with an average of 433 bp in length were obtained for the following analysis. In order to normalize the total count for further alpha and beta diversity analysis, the high-quality sequences in all samples were optimized to the minimum number.

Gut microbiota is associated with sheep breeds

After simplifying the original data, 1,359,405 high-quality available sequences were obtained. According to 97% breeds similarity, 7039, 6887, 4112, and 8257 OTUs were separately acquired from samples at groups DrS, DsS, TS, and STHS (Table S2), respectively. A total of 26295 OTUs were detected from all samples, of which 2448 obtained in all groups mentioned as core OTUs (Figure 1(A)). The core OTUs contained nearly 9.31 % of the entire OTUs. Furthermore, 225, 203 OTUs were uniquely detected in DrS and DsS groups, and 168, 654 unique OTUs were found in group TS and STHS, respectively. The number of observed OTUs in the TS samples was fewer than that of the other threes.

To confirm the quality of our sequencing data, we examined alpha and beta diversities of bacterial fraction of the sheep microbiota. Several alpha diversity indices diverged significantly between the four sheep breeds (Fig.1 (B) and (C)). The Shannon-Wiener index may straightly expose the heterogeneity of a community according to the amount of breeds present and their related abundance [22]. The Shannon-Wiener estimator of groups DsS, STHS, TS and DrS were 9.92, 9.33, 8.09 and 9.06, respectively. Within the groups, Chao1 and Shannon index visually reflected that the
abundance and diversity of intestinal microbial population in the TS group were lower than those in groups DsS, STHS and DrS, and the difference between these four groups was significant (P < 0.05; Figure 1 (B) and (C)). In terms of the Chao1 estimator, 3160, 2598, 1585 and 2207 average OTUs were noted in samples at groups DsS, STHS, TS and DrS, respectively, and the ACE estimator indicated 3263, 2646, 1627 and 2267 (Table S3), which was consistent with the results shown in Chao1, suggested that lower OTU richness was got in TS samples than the other threes (Fig.1 (B) and (C)). Collectively, these data pointed towards a more diverse bacterial population in TS compared to the other threes, and indicated differences in intestinal microbial composition in association with sheep breeds.

Comparison of bacterial microbiome diversity among different sheep breeds

The Bray-Curtis distance matrices were measured according to the OTUs abundance of each sample. Based on the distance matrices, the unweighted Unifrac similarity analysis indicated that the similarities among different sheep breeds were significant. The principal coordinates analysis (PCoA) was performed according to the phylogenetic-tree-based Unifrac metric. As shown in Fig. 2 (A), all samples were assembled into three clusters, scattered points in the principal component denoted different breeds and their relationship between each other. There were significant differences between breeds in relation to microbiome composition (PERMANOVA, p < 0.01). TSs were mainly aggregated in cluster B, whereas DsSs were mostly converged in cluster A. Moreover, DrSs and STHSs were more scattered and found between cluster A and cluster B. Both principal components accounted for 28.1% (PC1) and 8.5% (PC2) of the explained variance. Interestingly, two lambs in STHSs were clustered separately as shown in Fig. 2 (A), suggesting that there was a general difference in gut microbiome between adult sheep and lambs.

Nonmetric multidimensional scaling (NMDS) was used to further clarify the difference among all the breeds in bacterial population composition, which was performed using the Bray-Curtis similarity for all the samples at OUT level[1]. Be regarded as a dominant ordination method that could exhibit the non-linear relationship between samples, NMDS has been widely applied in the study of gut microbiome. As shown in Fig. 2 (B), there was distinguishing clustering of TSs samples, meanwhile
samples from DsSs were very close to DrSs. However, the samples from STHS were more dispersed (Fig. 2 (B)).

Additionally, hierarchical clustering analysis of all samples was used to exhibit the similarity between samples, which was performed with Unweighted pair-group method with arithmetic means (UPGMA) and used the Bray-Curtis similarity. Two primary groups were perceived in this analysis. One cluster contains the whole samples of TS and the other cluster contains the all samples from DrS (Fig. 3). It was similar with the results above, that TS samples were distinctive compared with other breeds. And the gut bacterial composition in sheep is largely influenced by the type of samples.

Gut microbial diversities and community composition among different sheep breeds

In order to clarify the diversity of gut bacterial composition in different sheep breeds, we estimated the gut microbiota in different taxonomical levels. The overall bacterial composition of different groups at the phylum level is identified in Fig. 4 (A) which shows that Firmicutes was the most predominant phylum in the 40 samples, followed by Bacteroidetes. The higher abundance of phylum Spirochaetes, Proteobacteria and Verrucomicrobia were found in TSs than those in the other three breeds, but the Deferribacteres was lacked in TSs (Fig. 4 (A)).

When analyzed on the family level from all samples, as shown in Fig. 4 (B), no significant differences were detected between these four groups. Ruminococcaceae and WCHB1-25 were the most abundant families in DsS, DrS and STHS groups, whereas Alcaligenaceae, Desulfovibrionaceae and Barnesiellaceae were almost absent. As for TS groups, the most abundant families were Spirochaetaceae, S24-7, Prevotellaceae, Barnesiellaceae and Succinivibrionaceae, while BS11 and WCHB1-25 were almost absent in the TS samples (Fig. 4 (B)).

In contrast to the family level, there are significant differences between TS group and the other three groups on the genus level. The main genera in TS group included Treponema, Succinivibrio, 5-7N15 and Prevotella (Table 1), while Bifidobacterium, Sharpea and YRC22 were absent (Fig. 4 (C)). Moreover, in the groups of DsS, STHS and DrS, Treponema remained the predominant population, and Coprococcus and Roseburia were relatively less abundant. However, it's worth noting that a large number of microbes in TS samples were relatively abundant, when compared to other three groups.
Discussion
This study was aimed at acquiring insight into the gut bacterial composition of four sheep breeds living in the QTP of China using next-generation sequencing technique. In our study, the gut bacterial community of four different sheep breeds was estimated by PCR-retrieved microbial 16S rDNA gene libraries. In our research, the intestinal microflora of fecal inclusion has been examined by bacterial diversity and abundance in different sheep breeds. These results indicate that there are significant differences of the gut microbiota between TSs and the other three sheep breeds (DrSs, DsSs and STHSs), besides, the bacterial diversity and composition in TSs are lower. However, the bacterial abundance in TSs are higher than those in the other three sheep breeds. The microbial diversity of TSs altered significantly compared with the other three breeds, which is similar with the earlier reports in high-altitude mammals [23, 24]. PCoA clustering analysis reveals that the microbial structure is distinct between the TSs and the other three breeds (Figure 2A). Besides, hierarchical clustering analysis shows that TS samples clearly cluster together, indicating that the intestinal microbial population of TSs are highly conserved for the comparison between interbreeds. From different taxonomical levels, the abundance of gut bacterial composition is also distinct among different sheep breeds. This phenomenon is probably due to the fact that Tibetan sheep have adapted to the high-altitude environment, while the other three breeds, as introduced later, are convergent to the commensal composition of Tibetan sheep. In addition, our study shows that the gut bacterial composition of lambs is quite different from that in adult sheep, suggesting that the gut microbial composition in lambs will develop towards adult sheep under their survival environment.

This study suggested that Firmicutes and Bacteroidetes were the most abundant phyla in the gut microbiomes of all samples, which accounted for 55.83% and 24.39% of the total microbial abundance, respectively. It was consistent with the prior studies in herbivores [25]. The functions of Firmicutes and Bacteroidetes were closely associated with carbohydrate, protein, and fiber metabolism [26]. The crop straws mainly contained cellulose, hemicellulose and lignin, and the Firmicutes contained a lot of fiber-decomposing bacteria, including Butyvibrio, Ruminococcus, Oscillibacter and Eubacterium, which may explain why the Firmicutes were dominant in the rumen
bacterial community of ruminants [25]. *Bacteroidetes* was the major degraders for decomposing non-fibrous plant components in sheep gut, and the *Prevotella* had the highest composition in this bacteria group. It has been suggested that *Prevotella* can account for 60%-70% of the overall microbial communities in rumen, and it contained highly active hemicellulose decomposing microbes, which was important for the decomposition of non-fibrous polysaccharides or proteins in crops [27, 28].

In this study, the relative abundance of *Firmicutes* and *Bacteroidetes* were not consistent among the different sheep breeds, *Bacteroidetes* had higher abundance in TS than that in other three breeds. In order to investigate the reasons of this result, we estimated the genera level of taxonomy and found that within the phylum *Bacteroidetes*, three genera had significantly higher abundance in TS than that in other three sheep breeds, including 5-7N15, *Prevotella* and *Bacteroides*, all of them belonged to the class *Bacteroidia* (Table 1). Previous study has shown that the *Prevotella* played an important influence on the fermentation process of feed in the rumen of sika deer [29]. Other studies also suggested that *Prevotella* was predominant in ruminants [28, 30, 31]. *Prevotella* was used to degrade lignocellulosic feedstock with excreate xylanase and CMCase [32]. Besides, some genera of the class Bacteroidia had high active hemicellulose decomposition and provided the hosts with the abilities to digest and extract available nutrition from fibrous plants [33]. In our research, we hypothesized that this phenomenon is probably due to the fact that TS have adapted to the high-altitude environment, while the other three breeds are convergent to the commensal composition of TS.

A special bacterial phylum was identified in the four sheep breeds was *Proteobacteria*, which was observed in various ruminants, such as sheep and cattle, but a relative low proportion (0.8%) was found in yaks [34-36]. In this study, *Proteobacteria* had significantly higher abundance in TS than that in the other three sheep breeds (p < 0.01). It has been reported that phylum *Proteobacteria* had the highest richness in the bovine rumen[37], and played a crucial role in the biofilm formation, fermentation and the soluble carbohydrates digestion [38]. We inferred that the reason for the higher abundance of *Proteobacteria* in TS was related to the different breeds of sheep. TSs have adapted to surviving in this harsh plateau environment, where Tibetans raise these animals for food and
sustenance [23, 39]. Several studies revealed that the gut microbes of ruminants help them survival at high altitude, which involve in the energy metabolism pathway [23, 24, 40]. Further study will be necessary for investigating their nature and effect in the gut.

Conclusion

On the whole, our 16s rDNA analyses provide essential views into the gut microbiome of different sheep breeds and highlight the difference between TSs and the other three sheep breeds (DsSs, STHSs and DrSs). This study also showed a lot of high abundance species, which may play some specific important roles in the hosts. The fluctuation of gut bacteria composition indicated that gut microbes could be changed along with the differences of sheep breeds. Different breeds caused a shift on the microbial community structure and decreased the bacterial species diversity in the gut of TS. And we have to expand study quantity in the future. Furthermore, the investigation of their distinct microbial structures may provide theoretical basis for the light on ecology of these comparatively abundant, but enigmatic microbial symbionts of ruminants.

Methods

Description of samples

Sheep were obtained from specialized farms (Qinghai Province, China) and were analyzed faecal 16s rDNA sequences from 40 individuals including 10 Dorset sheep (DrS), 15 Small Tail Han sheep (STHS), 5 Tibetan sheep (TS) and 10 Dorper sheep (DrS). Each sheep breeds we selected were self-propagated and fed by the profit-making sheep farm and had similar genetic context. All screened animals were healthy and no other diseases appeared prior to the sample selection. Table S1 provides the detailed information of each sheep we sampled.

DNA Extraction

Following the manufacturer’s instructions of Omega Bio-tek, microbial genomic DNA was extracted from 500 mg of each fecal sample using the fecal DNA kit. Meanwhile, we measured the DNA quality with 1% agarose gel electrophoresis, and examined the concentration through the NanoDrop Spectrophotometer, storing DNA samples at \(-20\cdot^\circ\text{C}\).

PCR Amplification of 16s rDNA
The V3-V4 region, which was 468bp within the 16s rDNA gene, was used to build the illumine sequencing library and amplified with the broadly conserved primers 341F (5′-CCTACGGGNGGCWGCAG-3′) and 805R (5′-GACTACHVGGGTATCTAATCC-3′). Different identifier codes were added at each primer for the further illumina sequencing. Polymerase chain reaction (PCR) was applied in a 50 µl reaction system including 2x Phanta Max Master Mix (Vazyme, China), 10 mM each primer, 16 µl each ddH2O and 5 µl DNA template. The PCR program was initial denaturation at 95 °C, with 8 cycles of denaturation at 95°C for 30 s, and followed annealing at 55°C for 30 s, extension at 72 °C for 45 s, with a final elongation phase at 72 °C for 5 min. The PCR products were performed by Quant-It Pico Green kit (Invitrogen, United States) and put them together for further library preparation. Barcoded samples were combined equal concentrations according to volume of sequencing. By Agilent 2100 Bioanalyzer (Agilent Technologies, United States), we performed the library concentration and eluted with Tris_HCl (pH 8.5). After denaturation, barcoded samples were combined following the volume of sequencing and sequenced on a PE250 v3 instrument using 600 cycles MiSeq Reagent Kit on a MiSeq Platform (Illumina; United States).

Bioinformatics and Statistical Analysis

In our research, the generating sequences have been uploaded to the National Center for Biotechnology Information (NCBI) under accession number AR180907. The QIIME (Quantitative Insights Into Microbial Ecology, v1.8.0) was performed to process the raw reads, and then the paired reads were assembled by FLASH v1.2.7 [41, 42]. Subsequently, QIIME was used to filter and analyze the joined sequences. By UPARSE 7.0, operational taxonomic units (OTUs) were obtained with based on a 97% identity threshold. Eventually, the whole OTUs were categorized to distinct taxonomic levels by Ribosomal Database Project (RDP) classifier 2.2 [43]. Based on the OTUs information, R package VennDiagram was performed to complete the venn diagram. In addition, the phylogenetic tree was obtained by MAGA 5.2 after sequences alignment. Alpha diversity was measured by MOTHUR, which was referred to the microbial community diversity. Bray-Curtis distance and unweighted Unifrac was evaluated the similarities of different samples with R package vegan. The Bray-Curtis distance is founded on common OTUs among samples to provide equal weight to differences in each taxa [1, 44].
The Unifrac is used to create the phylogenetic tree for samples and the taxa which are phylogenetically related will give less divergent Unifrac values, while the unrelated taxa will generate larger differences [45]. QIIME was performed to generate phylogenetic beta diversity, and further to do principal coordinate analysis (PCoA) and hierarchical clustering analysis by R program based on Bray-Curtis distance and unweighted Unifrac. PERMANOVA and student’s t-test were performed to exam significant differences between various groups.

Declarations

**Abbreviations**

DrS: Dorset sheep; STHS: Small Tail Han sheep; TS: Tibetan sheep; DrS: Dorper sheep; NCBI: National Center for Biotechnology Information; QIIME: Quantitative Insights Into Microbial Ecology; OTU: operational taxonomic units; RDP: Ribosomal Database Project; PCoA: principal coordinate analysis; PICRUSt: Phylogenetic Investigation of Communities by Reconstruction of Unobserved States; NMDS: Nonmetric multidimensional scaling; UPGMA: Unweighted pair-group method with arithmetic means; QTP: Qinghai-Tibetan Plateau.

**Ethics approval and consent to participate**

All experiments were approved by the Ministry of Health in China for the care and use of laboratory animals and supervised by the Research Ethics Committee of Northwest A&F University. Informed consent was obtained from the animal owners in advance.

**Consent for publication**

Not applicable

**Availability of data and materials**

All data generated or analyzed during this study are included in this published article, and also available from the corresponding author on reasonable request.

**Competing interests**

The authors declare that they have no conflict of interest.

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Authors’ contributions

WTM and DKC conceived and designed experiments; XTY and JJC performed all experiments. XTY, CXZ and YXQ collected and analyzed the data. XTY and JJC drafted the manuscript. All authors read and approved the final manuscript.

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Table

[Please see the supplementary files section to view the table.]

Figures
Figure 1

The community composition and microbial diversity index analysis. (A) Venn diagram showing overlap in OTUs of differential abundance in DrS, DsS, TS and STHS. (B) Shannon index. (C) Chao1 index. Different asterisks represent statistical significance (*p < 0.05, ** p < 0.01, *** p < 0.001).
Figure 2

Compositional analyses of the gut microbiome of different sheep breeds. (A) PCoA plot of similarities between the different groups. Principal component (PC) 1 and 2 accounted for 28.1% and 8.5% of the variance, respectively. (B) NMDS showing the alteration of bacterial population based on Bray-Curtis distance.
Hierarchical clustering of bacterial communities according to Bray-Curtis distance
Figure 4

Microbial composition of different samples. Each bar represents the average relative abundance of each bacterial taxon within a group. (A) Taxa assignments at Phylum level. (B) Taxa assignments at Family level. (C) Taxa assignments at Genus level.

Supplementary Files
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Supplementarytable.pdf
revisedtable.pdf