

Comparative study on the effects of different feeding habits and diets on intestinal microbiota in *Acipenser baeri* Brandt and *Huso huso*

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Abstract

Background: Siberian sturgeon (*Acipenser baeri* Brandt) and Beluga sturgeon (*Huso huso*) are two important commercial fish in China, and the feeding habits of them are very different. Diets and feeding habits are two significant factors to affect the gastrointestinal microbiota in fish. The intestinal microbiota has been reported to play a key role in nutrition and immunity. However, it is rarely reported about the relationship between the intestinal microbiota and feeding habits/diets on different *Acipenseridae* fish. This study is to comparative analysis of gut microbial community in Siberian sturgeon and Beluga sturgeon fed with the same diet/Beluga sturgeon fed with different diets in order to determine the effects of different feeding habits/diets on the fish intestinal microbiota. Results: According to the experimental objectives, BL and BH groups were Beluga sturgeon (*Huso huso*) fed with low fishmeal diet and high fishmeal diet, respectively. SH group represented Siberian sturgeon (*Acipenser baeri* Brandt) fed with the same diet as BH group. After 16 weeks feeding trial, the intestinal microbiota was examined by 16S rRNA high-throughput sequencing technology. On the phylum level, Proteobacteria and Bacteroidetes were significantly higher in BL group than BH group, and Cyanobacteria showed the opposite trend. Compared with BH group, Proteobacteria and Firmicutes were significantly increased in SH group, whereas Cyanobacteria were clearly decreased. At the genus level, *Pseudomonas* and *Citrobacter* in BL group were significantly higher comparing with BH group, while *Bacillus* , *Luteibacter* , *Staphylococcus* and *Oceanobacillus* was lower in BH group than SH group. Conclusions: Alpha and beta diversities indicated that the intestinal microflora were significant difference between Siberian sturgeon and Beluga sturgeon when they fed with the same diet. Meanwhile, Beluga sturgeon fed with low fishmeal diet can increase the species diversity of intestinal microbiota than it fed with high fishmeal diet. Therefore, feeding habits clearly affected the gastrointestinal microbiota of sturgeons. Moreover, the impact of changes in food on the gut microbiota of sturgeons should be taken into consideration during the process of sturgeon aquaculture.

Background

It is well known that the intestinal tract of vertebrates is a very complex and dynamic ecosystem, which is colonized by a large and diverse microbial community [1]. The intestinal microbiota of vertebrates has been reported to play a key role in nutrition and immunity, such as stimulating the growth and development of the intestinal epithelium, preventing it from pathogen invasion, contributing to the digestion of complex nutrients, and synthesizing beneficial secondary metabolites [2, 3]. As an aquatic vertebrate, the intestinal microbiota of fish has major influences on growth, health and development of fish [3, 4]. The stability of the intestinal microbiota is not only an extremely important factor in the inhibiting colonization of the intestine by pathogens, but also an essential factor for feed digestion [5, 6]. That is, investigation on the intestinal microbiota of fish may reveal intestinal development, homeostasis and the characterization of protection [7].

The composition of intestinal microbiota was impacted by many endogenous and exogenous factors, such as species, lifestyle, feeding habit, diet, nutritional status, living conditions, and so on [1, 3, 8].

Among environmental factors, dietary factors to a large extent affected the structure and abundance of intestinal bacterial community in fish [9]. Moreover, previous study indicated that the intestinal microbiota of omnivorous *Carassius cuvieri* showed the higher diversity than those of carnivorous individuals, which means that feeding habit was also the significant factor to affect intestinal microbiota composition [10]. Therefore, it is crucial to understand how such factors may influence the gut microbiota, with a view to regulate and control the bacterial community.

Sturgeon is the common name used for fish belonging to the Acipenseridae family that includes 27 species, such as *Acipenser*, *Huso*, *Scaphirhynchus* and *Pseudoscaphirhynchus* [11–13]. Siberian sturgeon (*A. baeri* Brandt) and Beluga sturgeon (*H. huso*) are two important species for aquatic farming in China not only for caviar, but also for meat [14, 15]. Siberian sturgeon belongs to the genus of *Acipenser*, and they have a wide range of food, which mainly feed on benthic animals, including chironomus larvae, mollusks, worms, crustaceans and small fish [16]. Besides, the previous study reported that Siberian sturgeon fed total plant-based diets with balance of essential amino acid (EAA) could maintain normal growth performance, which means that although Siberian sturgeon was the omnivorous fish, they also biased towards vegetarian diet [17]. Being carnivorous, Beluga sturgeon was come from the *Huso* genus and has the higher protein requirement to satisfy tissue growth and maintenance [18]. Meanwhile, there was evidence showed that replacing fishmeal with soybean meal without lactic acid significantly reduced growth performances of beluga, which further suggested that animal protein plays a key role in beluga diets due to their feeding habits [19]. Siberian sturgeon and Beluga sturgeon have a close phylogenetic relationship and have a considerable overlap in their spatial and temporal distribution. However, their diets are different, which contributes to their distinct ecological roles. At present, there are not sufficient studies that reference the difference of intestinal microbiota between Siberian sturgeon and Beluga sturgeon. In order to improve the health and productivity of these two important aquaculture species, it is interesting to characterize their gut bacterial communities and the factors that influence the composition and stability of the microbiota. Therefore, the present study was made to determine the intestinal bacterial communities of Beluga sturgeon fed two diets with different fishmeal levels, and to compare the intestinal microbiota of Siberian sturgeon and Beluga sturgeon fed with the same diet in the same environment, respectively.

Methods

Experimental design and diet formulation

Two isonitrogenous (410 g/kg crude protein) and isocaloric (19 MJ/kg) diets were formulated to contain two different fishmeal levels [high fishmeal: 250 g fishmeal/kg diet, FM250] and [low fishmeal: 100 g fishmeal/kg diet, FM100] [14]. The FM100 diet was supplemented with threonine, methionine, lysine and fish oil to keep the same EAA and highly unsaturated fatty acid (HUFA) composition with FM250 diet. The diets were made into sinking extruded pellets by an extruder (MY56X2A, MUYANG Group, Jiangsu province, China) in different pellet diameters (2.0 mm and 3.0 mm) according to fish size. All

experimental diets were air-dried and stored at $-20\text{ }^{\circ}\text{C}$ until use. Diets formulation and proximate compositions are shown in Table 1.

Three experimental groups were designed for this experiment including BL, BH and SH groups. BL and BH groups were Beluga sturgeon fed with low fishmeal diet (FM100) and high fishmeal diet (FM250), respectively. SH represented that Siberian sturgeon fed with high fishmeal diet (FM250) as same as BH group. The purpose of the design was to compare the difference of intestinal microbiota in Beluga sturgeon fed with two difference diets, and to analyze the composition of the intestinal microbial communities between Beluga sturgeon and Siberian sturgeon when they fed with the same diet.

Rearing conditions and sample collection

Siberian sturgeon and Beluga sturgeon were purchased from Zhongketianli Aquatic science and technology Co. Ltd (Beijing, China), where the feeding trial was also performed in a flow-through fish rearing system using recirculation water and under natural photoperiod conditions. Siberian sturgeon and Beluga sturgeon were acclimated to the flow-through fish rearing system and fed with a commercial fish food for 2 weeks before the trials. The system ensured nearly constant and optimal water quality to fishes (temperature $18.0 \pm 1.0\text{ }^{\circ}\text{C}$, dissolved oxygen $> 7.5\text{ mg/L}$, pH 8.1 ± 0.2 , nitrite $< 0.1\text{ mg/L}$). Then, healthy sturgeon ($37.5 \pm 0.0\text{ g}$) were randomly distributed into 9 net cages (capacity: 900 L) at the same cement pool. Triplicate replicate tanks were randomly assigned to three experimental groups (BL, BH and SH), and 60 fishes were batch weighed and stocked in each tank. During the 16-week feeding period, fish were fed with the experimental diets to apparent satiation three times daily at 8:00, 13:00 and 18:00 respectively.

At the end of the feeding trial, all fish were fasted for 24 h before sampling. Two fish in each tank were randomly and quickly captured and weighed individually after being anaesthetized by immersion, until sedate, in a 2-phenoxy ethanol (0.30 ml/L water) solution. After rinsing with sterile distilled water, the surface of the fish is wiped with 70% ethanol to reduce contamination, and then the fish was dissected with sterile scissors. The whole intestinal tract with faecal was removed from abdominal cavity stored in $-80\text{ }^{\circ}\text{C}$ immediately for the following intestinal microbiology analysis. All sampling operations were conducted on the ice [20].

DNA Extraction, purification and PCR amplification

Samples (intestinal content and water sediment) were thawed on ice, and then total bacterial DNA was extracted with Power Fecal DNA extraction kit (Mo Bio, USA) according to the manufacturer's instructions. Using the 0.8% agarose gels checked the purity and quality of the genomic DNA. And the DNA concentration was measured by using a fluorescence spectrophotometer (ES-2, Malcom, Japan). In order to avoid bias, all samples were extracted in duplicates, and the same sample extracts were mixed together [20, 21]. The extracted DNA was stored at $-80\text{ }^{\circ}\text{C}$ until it was used for high-throughput sequencing [22].

In a word, the primer, 336F-806R (F: GTACTCCTACGGGAGGCAGCA; R: GTGGACTACHVGGGTWTCTAAT), was used to conduct PCR amplification of the 16S rRNA V3-V4 region. For each sample, 10-digit barcode sequence was added to the 5' end of the forward and reverse primers (provided by Allwegene Company, Beijing). The PCR was carried out on an APPLIED Biosystems GeneAmp PCR system 9700 using 50 μ l reaction volumes, containing 5 μ l 10 \times Ex Taq Buffer (Mg^{2+} plus), 4 μ l 12.5 mM dNTP Mix (each), 1.25 U Ex Taq DNA polymerase, 2 μ l template DNA, 200 nM bar coded primers 336F and 806R each, and 36.75 μ l ddH₂O. Thermal cycling parameters were 94 °C for 2 min, followed by 30 cycles of 94 °C for 30 s, 57 °C for 30 s and 72 °C for 30 s with a final extension at 72 °C for 10 min. In order to mitigate reaction-level PCR biases, three PCR products per sample were mixed. The PCR products were purified using a QIAquick Gel Extraction Kit (QIAGEN, Germany), quantified using Real Time PCR, and sequenced at Allwegene Company, Beijing.

High throughput sequencing analysis

At last, deep sequencing was performed on Miseq platform at Allwegene Company (Beijing). After the run, image analysis, base calling and error estimation were performed using Illumina Analysis Pipeline Version 2.6. The raw data were first screened and sequences were removed from consideration if they were shorter than 200 bp, had a low quality score (≤ 20), contained ambiguous bases or did not exactly match to primer sequences and barcode tags. Qualified reads were separated using the sample-specific barcode sequences and trimmed with Illumina Analysis Pipeline Version 2.6. And then the dataset were analyzed using QIIME software package (Quantitative Insights Into Microbial Ecology). The sequences were clustered into operational taxonomic units (OTUs) at a similarity level of 97%, to generate rarefaction curves and to calculate the richness and diversity indices [24, 25]. The Ribosomal Database Project (RDP) Classifier tool was used to classify all sequences into different taxonomic groups [26, 27]. Mothur software [28] was used to calculate the alpha diversity indexes of the purified samples to obtain the richness and diversity indices of the bacterial community (i.e., ACE, Chao1, Shannon, and Simpson). To examine the similarity between different samples, heatmap figures, venn diagrams and non-metric multidimensional scaling (NMDS) were used based on the OTU information from each sample using R, while the cladogram was generated using the online LefSe project2.

Statistical analysis

Statistical analysis was carried out using STATISTICA 7.0 for Windows (StatSoft Inc., Tulsa, OK, USA). All data were presented as mean \pm SD and analyzed by one-way analysis of variance (ANOVA). Duncan's multiple range test and critical ranges was used to test differences among individual means. Difference were regarded as significant when $P < 0.05$.

Results

Analysis of 16S rRNA sequencing results

Six samples were taken from each group, but only 5, 5 and 3 samples from BL, BH and SH groups respectively accorded with the requirements of library construction, because the rest samples of PCR products were no purpose bands or low concentrations. After a series of purification and filtration processes on the sequencing results, 13,740 to 28,143 effective sequences were collected from each sample, resulting in a total of 246,700 sequences from the 13 samples. The statistics of the filtered sequencing data of each sample are shown in Table S1 (Additional file 1). The proportion of effective tags among Raw_Tags was in the 60% to 80% range at each sample. The reads sequences in the corresponding length range of each sample after quality control filtration were counted. The effective sequence length distribution is shown in Fig. S1 (Additional file 1). 400 bp to 440 bp were the most effective sequence length distribution region.

The number of sequences in OTU of each sample was obtained within the 97% sequence similarity threshold. Meanwhile, the classification information for each species corresponding to each OTU was also obtained by comparing the OTU representative sequences with a microbial reference database. The bacteria that could be detected were classified into 13 phyla, 21 classes, 41 orders, 77 families, and 120 genera. The composition of each sample community was calculated at phylum, class, order, family, genus and species levels, respectively. Table S2 (Additional file 1) was showed the number of each species at different levels, and Table S3 (Additional file 1) was recorded the total number of OTUs covered by each sample in their subordinate levels. The number of OTUs increased with the depth of sequencing, which was confirmed by the dilution curves of the OTUs measured in this study. The final curve became stable, which signified that the amount of sequencing data is somewhat reasonable (Fig. 1).

Comparison of core intestinal microflora in sturgeons from three experimental groups

Venn diagrams were used to show the core intestinal microflora of the sturgeons in three experimental groups. The core microflora of sturgeons meant the common bacterial populations in three experimental groups. As shown in Fig. 2, the number of OTUs shared by all sturgeon in three experimental groups was 243, while the number of OTUs was 510 sharing by sturgeon in BL and BH groups, and 278 for sturgeon in BH and SH groups. The main bacterial phyla in the intestines of each group of sturgeon are shown in Fig. 3. Three dominant phyla in BL and BH groups were Proteobacteria, Firmicutes and Cyanobacteria, while SH group was Proteobacteria, Firmicutes and Actinobacteria. The core intestinal microflora in fish from three experimental groups were the same, but their relative abundances were different. The top five core intestinal microflora at phyla in BL group showed the order from high to low was Proteobacteria, Firmicutes, Cyanobacteria, Bacteroidetes and Actinobacteria ($P<0.05$), whereas the core intestinal microflora in BH groups showed the difference with the order of Cyanobacteria>Proteobacteria>Firmicutes>Actinobacteria>Bacteroidetes. And the order from high to low of core intestinal microflora in SH group was Proteobacteria, Firmicutes, Actinobacteria, Cyanobacteria, and Bacteroidetes.

A heatmap (Fig. 4) is a graphical representation that uses a system of colored gradients to represent the size of values in a data matrix, and the cluster data are also expressed in heatmap according to species

or the abundance similarity of samples. In order to reflect the similarities and differences between multiple sample communities, high-abundance and low-abundance species are clustered by color gradient and similarity. Based on the species composition and relative abundance of each sample, a heatmap analysis was performed to extract the species at each taxonomic level. Mapping was achieved using R language tools, and a heatmap cluster analysis was performed at the levels of the phylum, class, order, family, genus, and species, respectively. The result was found in Fig. 4 that the vertical clustering between BH and SH groups showed the long branch length, which indicated the richness of intestinal microbiota between two groups was clearly different. Similarly, BL and BH groups have a certain degree of similarity in richness, because the short branch length was found in BL and BH groups at Fig. 4. In addition, the relative abundance of intestinal flora at genus level among BL, BH and SH groups were also showed in Fig. 4. The major intestinal microflora at genus level in BL group were *Acinetobacter*, *Pseudomonas*, *Bacillus*, and so on, as same as the BH group, while the SH group included *Bacillus*, *Staphylococcus* and *Acinetobacter* intestinal microflora at genus level.

Alpha diversity analysis of microbial communities in sturgeons from three experimental groups

Alpha diversity refers to the richness and diversity of a single sample species. There were several indices for measurement of alpha diversity, such as the Chao1, Observed_species, Shannon, PD_whole_tree, and Good's coverage. The Chao1 index measures the richness of species (i.e., the number of species), whereas the Shannon index measure the diversity of species. Observed_species shows the number of OTU was observed with the increase of sequencing depth. PD_whole_tree refers to the number of species observed, reflecting the abundance of the colony. The completeness of the sequencing was tested by good's coverage, which was close to 99% in this study, indicating that the majority of the bacterial species present in the sample had been detected.

Table 2 showed the results of alpha diversity for the gut microbiota in sturgeons from three experimental groups. A total of 826 OTUs were obtained at the 97% similarity level. There was a significant difference in the OTUs, Chao1, Good's coverage, Observed species and Shannon indices between the BH and SH groups ($P < 0.05$), but no significant difference was found in the PD whole tree of the two groups ($P > 0.05$). The result showed that the alpha diversity indices were considered to be significantly different between Siberian sturgeon and Beluga sturgeon when they fed with the same diet ($P < 0.05$).

Meanwhile, although the most alpha diversity indice were no significant difference between BL and BH groups ($P > 0.05$), but the Shannon index was obviously higher in BL group than that in BH group ($P < 0.05$), which pointed that the species diversity of the intestinal microbiota was enhanced when Beluga sturgeon fed with the low fishmeal diet.

Beta diversity analysis of microbial communities in sturgeons from three experimental groups

Non-metric multidimensional scaling (NMDS) (Fig. 5) is a data analysis method that simplifies research objects (samples or variables) in multidimensional space to low-dimensional space for positioning, analysis and classification, while retaining the original relationship between objects. It is applicable to the

case that the exact similarity or heterogeneity data between the research objects cannot be obtained, but only the hierarchical relationship data between them can be obtained. Its basic feature is to treat the data of similarity or dissimilarity between objects as a monotone function of the distance between points, and replace the original data with a new data column of the same order for the metric multidimensional scale analysis on the basis of maintaining the original data order relationship. In other words, when the data is not suitable for the direct multidimensional scaling analysis of variable type, the variable transformation is carried out and then the multidimensional scaling analysis of variable type is adopted. For the original data, it is called non-metric multidimensional scaling analysis. Its characteristics are reflected in multi-dimensional space in the form of points according to the species information contained in the sample, and the degree of difference between different samples is reflected by the distance between points, and finally the spatial locus map of the sample is obtained.

The closer the distance between sample points, the higher the similarity. Generally speaking, the samples within the same circle meant the difference between samples was not obvious, while the sample points within circles with no intersection indicated that there was significant difference between the samples. As shown in Fig. 5, the circles were intersected between BL and BH groups, but no intersection was found in BH and SH groups, which indicated the beta diversity between BH and SH groups was significant difference, whereas BH and BH groups was not.

Analysis of the differences in gut microbiota among sturgeons in three experimental groups at the phylum and genus levels

LEfSe (Linear discriminant analysis Effect Size) (Fig. 6) is an algorithm for high-dimensional biomarker discovery and explanation that identifies bacteria of each level of phylum, class, order, family, or genus characterizing the differences among BH, BL and SH three groups (Fig. 6). The cladogram (Fig. 6a) showed differences in 102 taxa among fish in BL, BH and SH groups. BL fish (enrichment in the Gammaproteobacteria and Proteobacteria) have no similar to BH fish, with a lot of changes occurring in the BH group such as enrichment in Actinobacteria, Corynebacteria and Norynebacteria (Fig. 6b). And the SH group was mainly enriched in Bacillales and *Bacillus* genus, which was clearly difference with BH group (Fig. 6b).

In addition, Fig. 7 shows the differences in relative abundances at the phyla level of the top 5 bacterial communities and genera level of the top 8 bacterial communities in the three experimental groups. In the fish from BL group, the relative abundance of Proteobacteria and Bacteroidetes were significantly higher than that in the sturgeons from BH groups ($P < 0.05$). In contrast, the relative abundances of the Cyanobacteria phyla in BL group were significantly lower than in BH group ($P < 0.05$). Both groups did not show any significant differences in the relative abundances of Firmicutes and Actinobacteria ($P > 0.05$) (Fig. 7a).

Compared with the sturgeons of BH group, the relative abundance of Proteobacteria and Firmicutes were significantly increased in sturgeons from SH group, whereas the relative abundance of Cyanobacteria

were clearly decreased ($P < 0.05$). The Bacteroidetes and Actinobacteria relative abundance were no significant differences in sturgeons of BH and SH groups ($P > 0.05$) (Fig. 7b).

At the genus level, the relative abundances of *Pseudomonas* and *Citrobacter* in BL group were significantly higher compared with BH group ($P < 0.05$) (Fig. 7c), while the relative abundance of *Bacillus*, *Luteibacter*, *Staphylococcus*, and *Oceanobacillus* was lower in BH group than in SH group ($P < 0.05$) (Fig. 7d). Three groups did not show any significant differences in the relative abundances of *Alpinimonas* and *Acinetobacter* ($P > 0.05$).

Discussion

With the in-depth study of intestinal microbiota in aquatic animals and the wide application of high-throughput sequencing technology, the progress was visible about the study on the relationship between gastrointestinal microecology and fish health [29]. The research of fish intestinal microbiota can help for understanding how gut microbial communities are assembled and how they impact host fitness. In the same environment, the present study was the first one to compare gut bacterial communities of Beluga sturgeon and Siberian sturgeon fed with the same diet and of Beluga sturgeon fed with two different diets by high-throughput sequencing methodology. Thirteen samples from the intestine contents and mucosa of three groups were analyzed by using high-throughput sequencing of the V3/V4 regions of the 16S rRNA gene.

The analysis results showed that the most effective sequence length distribution region was 400 bp to 440 bp and as the sequencing depth increased, the number of OTUs increased and eventually leveled off, which indicated the amount of sequencing data in this study was reasonable (Figs. 1 and S1). As seen in Figs. 2 and 3, the main composition of gut microbiota of both BH and SH groups are Proteobacteria, Firmicutes, Cyanobacteria, and Actinobacteria when they fed with the same diet. This result was consistent with the majority of studies on the gut microbiota of sturgeons [30–32]. Noteworthy, the relative abundances of intestinal microbiota were completely different in the BH and SH groups, which showed that the intestinal microbiota was difference when the different Acipenseridae fish (Siberian sturgeon and Beluga sturgeon) fed with the same diets. In other words, this result further indicated that feeding habit is an important factor that affects gastrointestinal microbiota. Many previous studies reported that the intestinal microbiota was influenced by the host genotype [33, 34]. Similar results were discovered by Li et al. [35], who point out different species of Carp fed with a commercial fish food in the same earth pond have the difference bacterial communities in the intestines. Meanwhile, three eel species for investigating the autochthonous microbiome using 16S rDNA sequencing indicated that the composition of intestinal microbiome of eel was affected by the characteristics of different eel species [36]. In addition, the alpha and beta diversity indices of the intestinal microbiota between BH and SH groups were also significantly different, which also confirmed the above point of view that the richness and diversity of the intestinal microorganisms are indeed dissimilar in the different species of sturgeons although they fed with the same diet. This result was also consistent with Li et al. [37], who reported that feeding habits and genotype clearly affected the gastrointestinal microbiota of fish.

On the other hand, although the dominant phyla in BL and BH group were Proteobacteria, Firmicutes and Cyanobacteria, while their relative abundance of the dominant phyla was significant difference, which indicated that the same fish (Beluga sturgeon) fed different diets were able to affect the intestinal microbiota. Lv et al. [31] found that the relative abundances of intestinal microbiota were difference between the wild Kaluga (*Huso dauricus*) and cultured Kaluga sturgeon, and their predominant bacteria are Proteobacteria and Fusobacteria, respectively. Besides, there were reported that the microbiota composition of Atlantic Salmon (*Salmo salar*) was influenced by experimental diets when they fed with fishmeal, soybean meal or fermented soybean meal diets [38].

In addition, BL and BH groups had no obviously differences in the alpha and beta diversity indices of the intestinal microbiota except for Shannon index. The result hinted that the different diets had a little effect on the richness of intestinal flora in the same fish (Beluga sturgeon) within a certain period of time. Studies have shown that the normal dominant bacteria in the intestinal tract of aquatic animals are anaerobic bacteria, accounting for more than 99%, and aerobic bacteria and facultative anaerobic bacteria account for about 1% [39]. Due to the symbiosis between anaerobic bacteria and intestinal wall, the content of anaerobic bacteria was stable, while the aerobic bacteria can be free in the middle of the intestinal cavity, so the content of aerobic bacteria was a random fluctuation in the intestinal tract [40]. As a result, the total number of anaerobic bacteria in the intestinal tract of different fishes has little difference, while the total number of aerobic bacteria has great difference [41]. This may explain why the richness of intestinal flora in the same fish with different diets is the same. Interestingly, the Shannon index was significantly higher in BL group compared with BH group, which indicated that Beluga sturgeon fed with low fishmeal diets (fishmeal partially replaced by cottonseed protein) can enhance the diversity of intestinal microbiota. This result was similar with the earlier study on the northern snakehead (*Channa argus* Cantor, 1842), which showed that different dietary soybean meal substitutions significantly affected the intestinal microbiota composition [42]. Similar results were also found in other fish species, such as rainbow trout (*Oncorhynchus mykiss* Walbaum) [43], European sea bass (*Dicentrarchus labrax*) [44] and Atlantic Salmon (*S. salar*) [38].

At the phylum level, the richness of Proteobacteria and Bacteroidetes in the intestines of BL group was clearly higher than that of BH group, whereas for Cyanobacteria was exactly the opposite (Fig. 7a). Proteobacteria, as the most abundant in the fish ponds, participated in various biogeochemical processes (for example, carbon, nitrogen, and sulfur cycling) in aquatic ecosystems [45, 46]. Previous studies have indicated that Proteobacteria were one of the most abundant phyla in the intestinal samples of fish as well as in most mammals gut samples [47]. Some researchers have already verified that some bacteria of Proteobacteria in the gut of healthy fish may significantly contribute to the digestive function [3]. Bacteroidetes is responsible for the metabolism of steroids, polysaccharides, and bile acids, helping the host in the absorption of polysaccharides and the synthesis of protein [48, 49]. Moreover, Bacteroidetes was also typically enriched in other herbivores [50–52]. In other words, herbivores were more likely to accumulate more Bacteroidetes compared with carnivores. Hence, we may infer that the intestines of the BL group have the higher amount of Proteobacteria and Bacteroidetes might be because the fishmeal was partially replaced by cottonseed meal in the diet of the BL group.

The ratio of Firmicutes to Bacteroidetes ratio (the F/B ratio) is significantly correlated with obesity [53–56]. There were also reported that the F/B ratio of the gastrointestinal microbiota of carnivorous fishes were obviously higher than other fishes [37]. Coincidentally, the result of this study showed that the F/B ratio of SH group was 13.8, while the F/B ratio in BH group was 14.6 (Fig. 7b), which indicated that Beluga sturgeon is actually bias carnivorous than the Siberian sturgeon. This result accorded with previous study, which reported that Siberian sturgeon was the omnivorous fish, they also biased towards vegetarian diet when they compared with beluga [17].

At the family and genus levels, the relative abundance of *Pseudomonas* and *Citrobacter* in BL group was significantly higher than those in BH group ($P<0.05$) (Figs. 4, 6 and 7c). Previous studies have reported that *Citrobacter* was one of the important cellulose-degrading bacteria [57–59]. This point of view can explain our results that the abundance of *Citrobacter* was increased when Beluga sturgeon fed with low fishmeal in order to improve the digestion of cellulose. Similar results were reported by Liu et al. [60], who found that *Citrobacter* was one of the most abundant bacteria in the herbivorous fish. Wu et al. [20] also discovered that *Citrobacter* was one of the most abundant in grass carp samples. In addition, the previous work reported that the *Pseudomonas* was more frequently found in grass carp (*C. idella*) fed a soybean meal diet compared to a casein meal diet. The current study also showed that the abundance of *Pseudomonas* was enhanced in Beluga sturgeon when they fed with high vegetable protein diets. Therefore, the increase of these pathogenic bacteria may indicate that Beluga sturgeon might not be adapted to the low fishmeal diet very well, increasing the risk of intestinal infections.

Another result (Figs. 4, 6 and 7d) indicated that the relative abundance of *Bacillus*, *Luteibacter*, and *Staphylococcus* of the sturgeons in SH group was significantly higher than those in BH group on the family and genus levels ($P<0.05$). The previous studies reported that the genus of *Bacillus*, as a typical probiotic, was used for enhancing host immunity and extracting nutrients consumed in fish diets [61, 62]. The function of *Bacillus* was able to improve the activity of digestive enzymes, enhance the immune system response, improve survival rate and confers disease resistance against pathogenic *Vibrio* species [63]. Moreover, *Bacillus* was also one of the important cellulose-degrading bacteria [57–59]. Hence, there were two points to contribute to the present result. On the one hand, Siberian sturgeons had the better colonization of *Bacillus* than Beluga sturgeon when both of them fed with the same diet. On the other hand, the increasing of *Bacillus* can improve the cellulose digestion in the Siberian sturgeon, which means that Siberian sturgeon had higher adaptability for vegetarian diet when they compared with Beluga sturgeon.

Conclusions

In conclusion, Siberian sturgeon and Beluga sturgeon, as the two different species from the same family, were significantly different in feeding habits. When they fed with the same diets in the same environment at the present study did not result in similar intestinal bacteria, suggesting that the specific endogenous factors outweighed by far the environmental factors to mould the composition of microbiota [35]. Meanwhile, our results showed that Beluga sturgeon fed with the low fishmeal diets can increase the

species diversity of intestinal microbiota compared with it fed with high fishmeal diets. Therefore, the impact of changes in food on the gut microbiota of the sturgeons should be taken into consideration during the process of sturgeon aquaculture.

Abbreviations

EAA: Essential amino acid; FM: Fishmeal; EPA: Eicosapentaenoic Acid; DHA: Docosahexaenoic Acid; one-way ANOVA: One-way analysis of variance; SD: Standard deviation; PCR: The polymerase chain reaction; RDP: The ribosomal database project classifier; NMDS: Non-metric multidimensional scaling; LEfSe: Linear discriminant analysis Effect Size

Declarations

Ethics approval and consent to participate

This paper does not contain any studies with human participants by any of the authors. No specific permits were required for the described field studies. No specific permissions were required for access to the artificial pond in Zhongketianli Aquatic science and technology Co. Ltd, Beijing, China. The field studies did not involve endangered or protected species.

Consent for publication

Not applicable.

Availability of data and materials

All data generated and analyzed during this study are included in this published article. The raw data are available from corresponding author under on reasonable request.

Competing interests

The authors declare that they have no competing interests.

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Authors' contributions

GX drafted the manuscript; GX, WX and TL participated in DNA extraction, 16S rRNA sequencing and sample collection; ZM, NJ and LL participated in study design and performed data analysis; MX

participated in study design. All authors read and approved the final manuscript.

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Tables

Table 1 Ingredient composition and proximate composition of experimental diets *

Ingredient (g/kg diet)	FM 100	FM 250
Fish meal ¹	100	250
Soybean meal	230	250
Wheat flour	163.4	221
Krill meal	60	60
Wheat gluten	50	50
Brewer's yeast	50	50
Soybean lecithin	20	20
Fish oil ¹	36.3	25
Soybean oil	17	24
Premix ²	20	20
Cottonseed protein	190	0
65% Lysine	9	0
Methionine	1.5	0
Threonine	1.8	0
Ca(H ₂ PO ₄) ₂	51	30
Proximate composition (g/kg)		
Crude protein	410	407
Crude fat	91	92
Moisture	80	94
Ash	95	79
Gross energy (MJ/kg)	19.1	19.1
Total phosphorus	16.1	16.3
Total calciums	14.7	15.3
Threonine	15.3	14.8
Methionine	7.7	7.4
Lysine	23.3	22.6
EPA ³	5.5	5.3
DHA ³	5.1	5.2

¹ Fish meal and fish oil were produced in Peru and supplied by the International Fish Meal and Fish Oil Organization (IFFO, Hertfordshire, UK); Soybean meal, soybean oil and lecithin were supplied by YiHai Kerry Investment Company Limited, Shandong, China; Wheat flour were supplied by Guchuan Group, Beijing, China.

² Including vitamin premix (mg/kg diet): vitamin A 20; vitamin B₁ 12; vitamin B₂ 10; vitamin B₆ 15; vitamin B₁₂ 8; niacinamide 100; ascorbic acid 1,000; calcium pantothenate 40; biotin 5; folic acid 10; vitamin E 400; vitamin K₃ 20; vitamin D₃ 10; inositol 200; corn protein powder 150. Mineral premix (mg/kg diet): CuSO₄×5H₂O 10; FeSO₄×H₂O 300; ZnSO₄×H₂O 200; MnSO₄×H₂O 100; KIO₃ (10%) 80; Na₂SeO₃ (10% Se) 67; CoCl₂×6H₂O (10% Co) 5; NaCl 100; Zeolite 16138.

³ EPA: Eicosapentaenoic Acid; DHA: Docosahexaenoic Acid.

* All diets were produced at National aquafeed safety evaluation station, Beijing, China, as extruded pellets.

Table 2 Number of reads, reads assigned to OTUs, good's coverage and alpha diversity indices of intestinal microbiota composition in sturgeons from three experimental groups

	BL	BH	SH
OTUs	357.2±32.27 ^b	314.4±30.14 ^b	191.0±9.64 ^a
Chao1	390.05±28.66 ^b	328.61±39.76 ^b	214.21±19.3 ^a
Good's coverage	0.994±0.0005 ^a	0.995±0.0010 ^a	0.998±0.0007 ^b
Observed species	335.32±26.36 ^b	271.92±27.92 ^b	186.47±7.84 ^a
PD whole tree	108.49±18.45 ^b	58.67±23.33 ^{ab}	18.67±1.01 ^a
Shannon	5.62±0.08 ^b	4.37±0.27 ^a	5.41±0.02 ^b

Note: means in the same row with different superscripts are significantly different ($P<0.05$)

Additional File

Additional file 1:

Table S1. Statistics of sequencing data of each sample after filtration.

Table S2. Statistics of OTU species of samples on various levels.

Table S3. Statistics of OTU clustering results of samples on various levels.

Figure S1. Effective sequence length distribution.

Figures

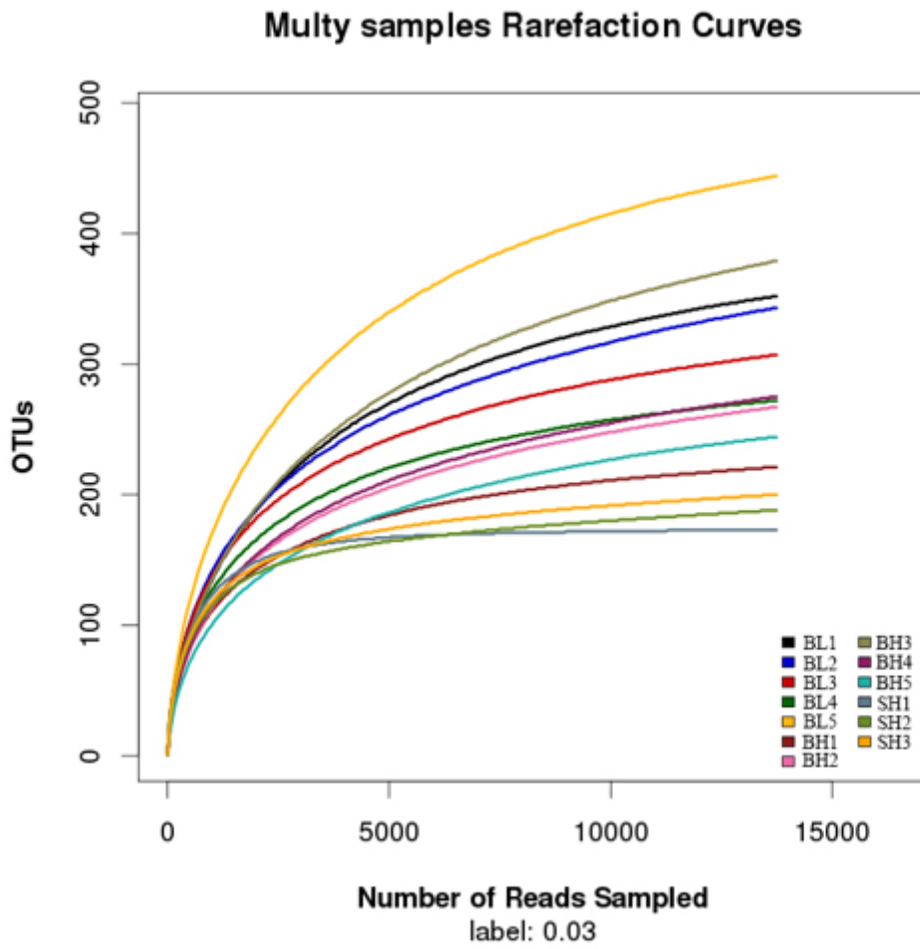


Figure 1

Rarefaction curve.

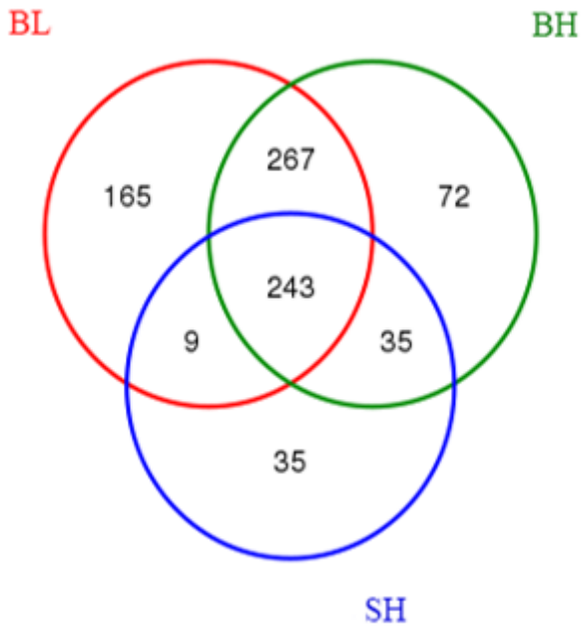


Figure 2

Venn diagram.

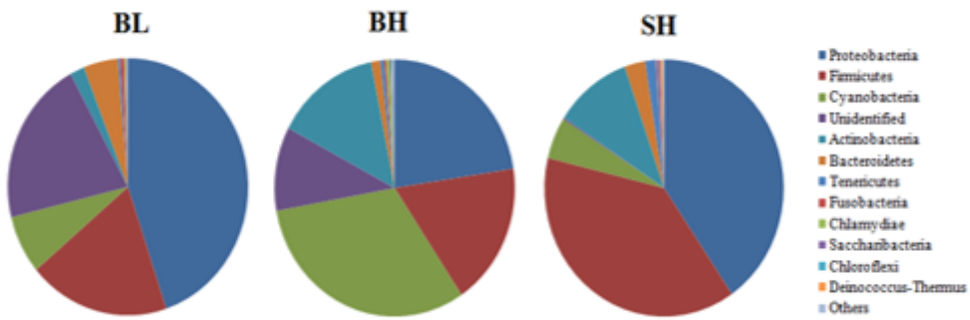


Figure 3

Pie charts.

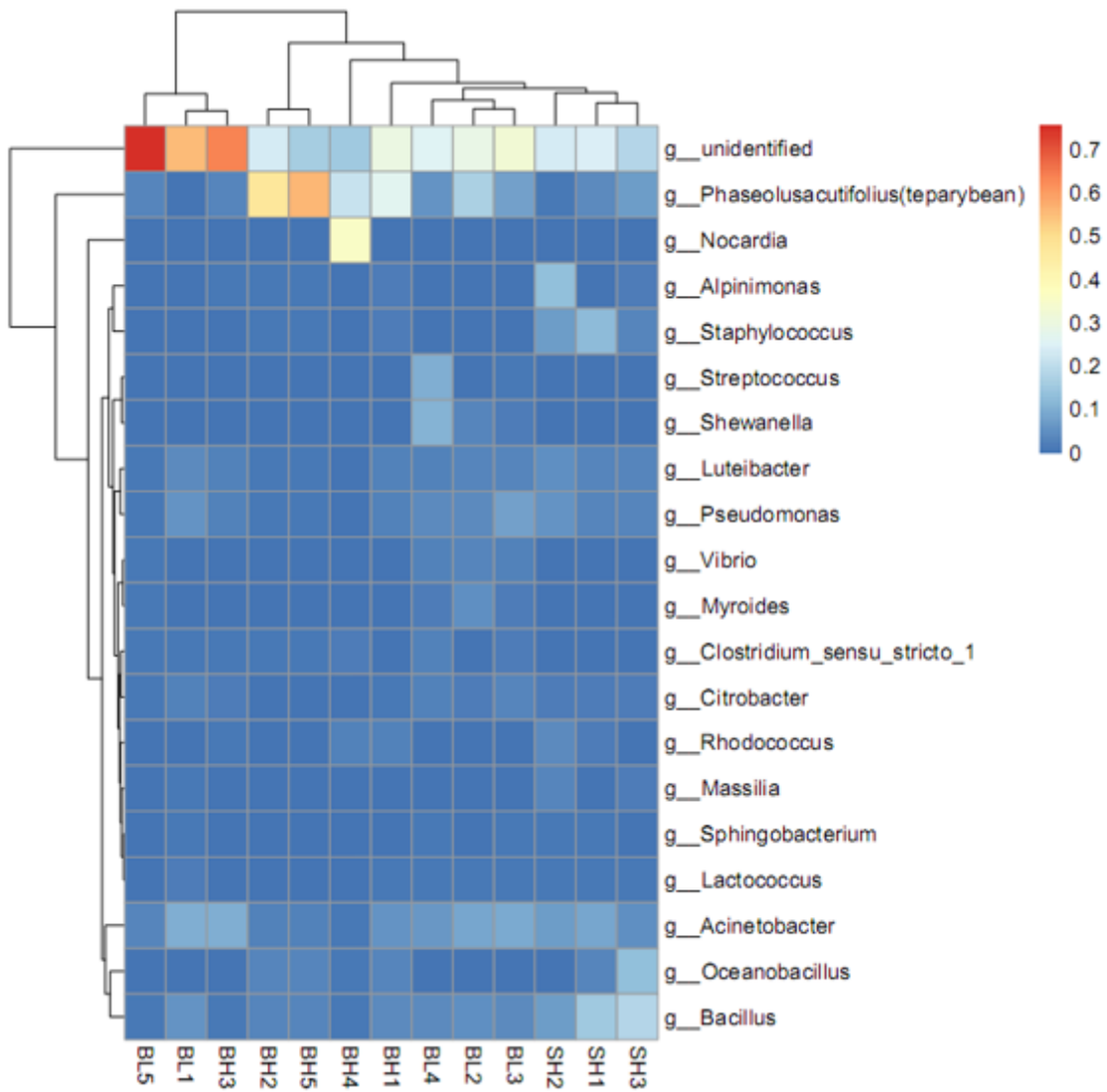


Figure 4

Heatmap showing richness of species at each level.

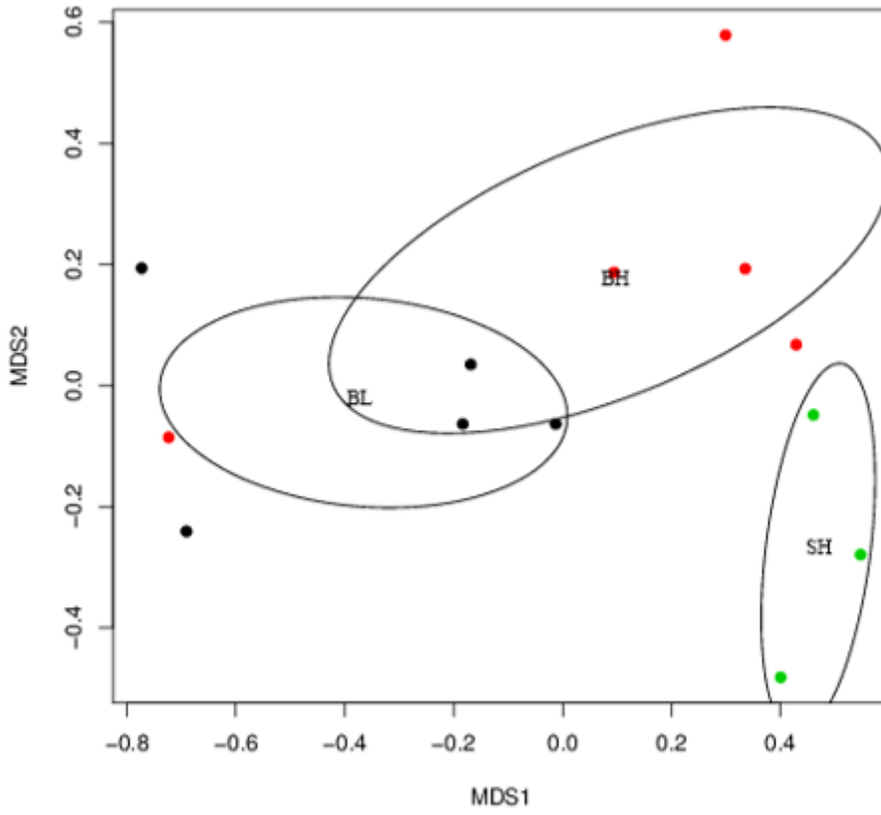


Figure 5

NMDS analysis.

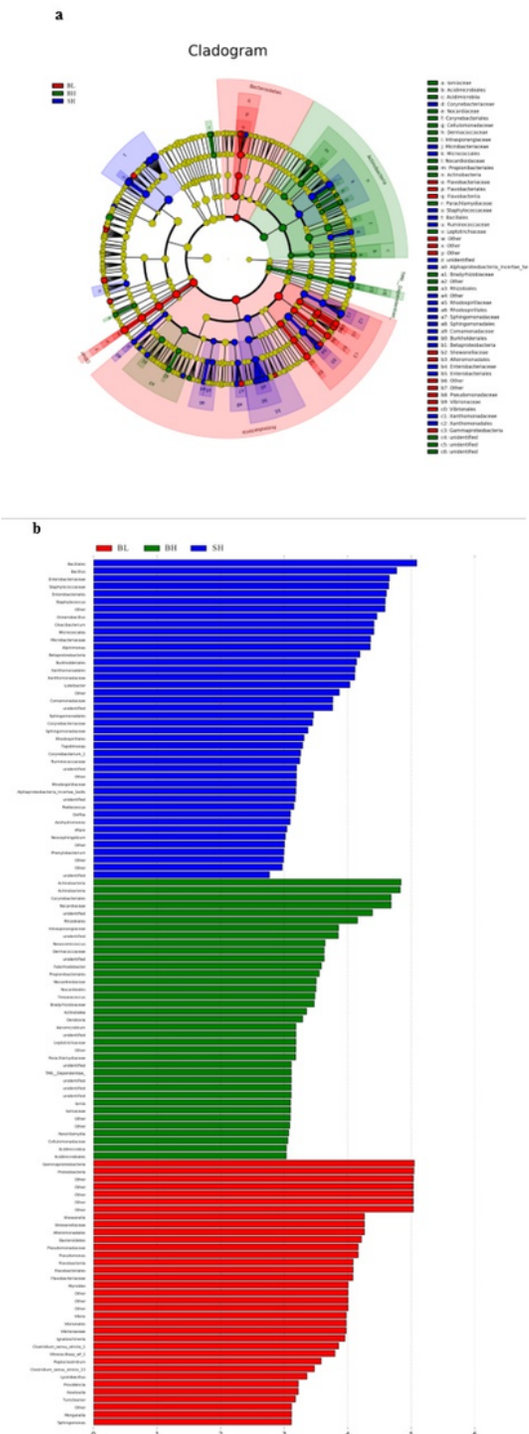


Figure 6

Taxonomic cladogram obtained from LefSe analysis of 16S rRNA sequencing (a). Only taxa with LDA score >2 are shown (b).

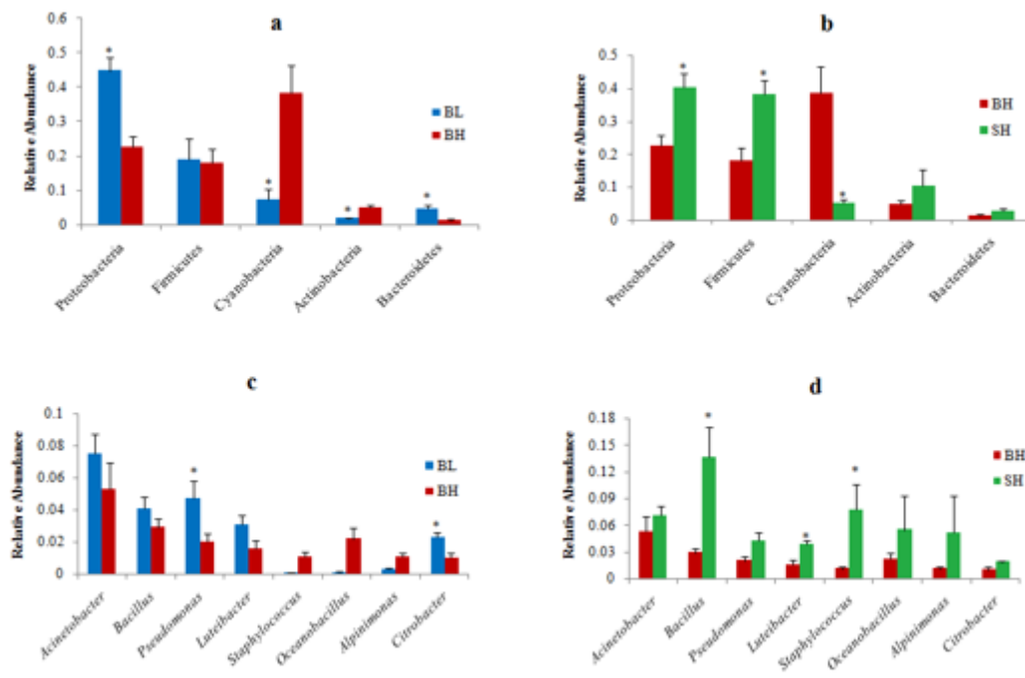


Figure 7

(a and c) Relative abundances (mean % SD) of five major bacterial phyla and eight major bacterial genera between BL and BH groups. (b and d) Relative abundance (mean % SD) of five major bacterial phyla and eight major bacterial genera between BH and SH groups.

Supplementary Files

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