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**Gut microbial community in proboscis monkeys: implications for effects of geographical and social factors**

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Abstract

Recent technological advances have enabled comprehensive analyses of the previously uncharacterized microbial community in the gastrointestinal tracts of numerous animal species; however, the gut microbiota of several species, such as the endangered proboscis monkey (*Nasalis larvatus*) examined in this study, remains poorly understood. Our study sought to establish the first baseline data on the gut microbiota of free-ranging foregut-fermenting proboscis monkeys and to determine how their microbiota are affected locally by environmental (e.g., geographical distance) and social (e.g., group size) factors in proboscis monkeys living in a riparian forest in Sabah, Malaysian Borneo. Using 16S ribosomal RNA gene sequencing of feces collected from more than 300 free-ranging proboscis monkeys, we demonstrated that they were dominated by Firmicutes and Bacteroidetes at the phylum level, with *Ruminococcaceae* and *Lachnospiraceae* being the most abundant families and *Oscillospira* and *Ruminococcus* being the dominant genera; this trend suggests that the microbial community composition of proboscis monkeys is not particularly distinctive compared to other foregut and hindgut fermenting primates. The microbial alpha diversity was higher in larger groups and individuals inhabiting diverse vegetation (i.e., presumed to have a diverse diet). For microbial beta diversity, some measures were significant, showing higher values with larger geographical distances between samples. These results suggest that social factors such as increased interindividual interactions, which can occur with larger groups, as well as physical distances between individuals or differences in dietary patterns, may affect the gut microbial communities of proboscis monkeys.

**Keywords:** Borneo; colobines; foregut-fermenters; gut microbiome; primate
Introduction

The gastrointestinal tracts of animals are colonized by many microorganisms, forming complex microbial ecosystems \(^1\)\(^-\)\(^3\). In general, animal-associated microbial ecosystems have been reported to have a direct effect on host health, contributing not only to daily energy acquisition through the production of vitamins and short-chain fatty acids but also to the host’s immune system and resistance to pathogens \(^4\)\(^-\)\(^6\). Understanding how the microbial community, which serves important functions in animals, is formed in the gastrointestinal tracts would shed light on the survival strategies of diverse animal species. Historically, the study of the human gastrointestinal microbial community has been extensive \(^7\),\(^8\). However, with recent advancements in sequencing technology, the ability to analyze gastrointestinal microbial diversity and community structure based on large amplicon libraries of 16S ribosomal RNA (rRNA) genes, primarily using fecal DNA, has prompted additional research in a variety of nonhuman primates \(^9\)\(^-\)\(^12\). With the recent accumulation of such research findings in nonhuman primates, it is becoming increasingly evident that their living environment influences and shapes the gastrointestinal microbial community \(^13\). For example, within the same primate species, a correlation has been observed between the diversity of dietary items and the microbial community e.g., \(^14\),\(^15\)\(^-\)\(^17\). Alternately, as in primate species that live in groups, it has been hypothesized that social factors may influence the establishment of the microbial community, as the horizontal transmission of the microbiome may occur via direct social interactions between individuals within the group or even indirect interaction via shared environments e.g., \(^18\),\(^19\)\(^-\)\(^21\).

Although the gastrointestinal microbial community, particularly the gut microbiome from fecal samples, has been progressively studied in various primate species over the last decade\(^10\),\(^22\)\(^-\)\(^25\), there are still numerous species in the wild for which even the most fundamental microbial community has not been studied. The foregut-fermenting proboscis monkey (\textit{Nasalis larvatus}), a large, sexually dimorphic, arboreal primate \(^26\), is one of these species for which the gut microbiota
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has not been studied in the wild. They inhabit various riparian and coastal forest environments, including riverine, mangrove, and peat swamp forests, and their dietary patterns have been reported to be flexibly adapted to these environments; diverse diets in riverine forests with higher plant diversity, and low dietary diversity in mangrove forests where plant diversity is extremely low. Due to its adaptable feeding habits, the proboscis monkey is an ideal study species for determining how habitat-specific dietary differences influence gut microbiota. Proboscis monkeys live in social groups, typically consisting of a single adult male and multiple adult females with their offspring, i.e., a harem group (although mixed-sex groups occasionally contain several adult males). There are also all-male groups and solitary males. Notably, the proboscis monkey is a rare species in which harem and all-male groups form complex societies, the so-called multilevel society, regularly. Consequently, it is a fascinating species with which to investigate the relationship between social factors and the intestinal microbial community.

We sought to establish the first baseline data on the gut microbiota of free-ranging proboscis monkeys and to determine how these gut microbiota were affected at the local scale by environmental factors (e.g., geographical distance) and social factors (e.g., sexes with different life histories and group size). Previous research has demonstrated that the gut microbiota is frequently influenced by dietary habits; therefore, we anticipated that there would be differences between the gut microbiota of proboscis monkeys living near the river mouth, where deforestation by oil palm plantations is more pronounced, and those living in the upper river areas, where relatively large areas of forest remain at our study site. Indeed, since increased dietary diversity has been reported to increase the diversity of the gastrointestinal microbial community, it would be predicted that disturbed habitats would generally experience a reduction in plant diversity and that the monkeys inhabiting these habitats would experience a reduction in dietary diversity. Furthermore, sex differences in social behavior have been associated with sexual biases in the gut microbiota in primates, and female-to-female grooming is the predominant form of
grooming in proboscis monkeys, but rarely between males and females. Thus, female proboscis monkeys are expected to have more contact with more individuals in the group, and accordingly, their gut microbiome may be expected to be more diverse and/or higher similarity than males. Lastly, given that several reports in primates have stated that direct inter-individual contact is associated with the transmission of gut microbiota, it can be predicted that individuals from the same group would have a similar gut microbiome. Additionally, it is likely that individuals in larger groups would have more opportunities for social interaction with more individuals, and as a result, they may possess a more diverse gut microbiome.

Methods

Study site and subjects

The study was carried out in a riverine forest along the Menanggul River, a tributary of the Kinabatangan River, Sabah, Malaysian Borneo (118° 30’ E, 5° 30’ N), inhabited by eight species of diurnal primates, including our study species, the proboscis monkey. For more than a decade, this area has been a popular tourist destination that attracts boat tours; as a result, the proboscis monkeys were well-habituated to human observers. The study site, in a 4 km stretch from the mouth of the Menanggul River upstream was home to over 200 proboscis monkeys, organized into 10 harem groups and various nonharem groups, including all-male, multimale multifemale groups, as well as solitary males. The southern portion of the Menanggul River is dominated by secondary forests, while the northern part has been cleared for oil palm plantations, excluding a protected zone along the river. Daily temperatures in the area were recorded at approximately 24°C (minimum) and 30°C (maximum), with an average annual rainfall of 2,474 mm (Matsuda et al., 2019). The river levels fluctuate by approximately 1 m daily, with seasonal floods causing an average increase of more than 3 m (Matsuda et al., 2010).
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Fecal sampling

Proboscis monkeys in the lower Kinabatangan Floodplain typically prefer to sleep along the river. Therefore, to collect their fecal samples, we conducted our sampling process in the early morning, after the monkeys had left the sleeping trees along the river we had identified the previous night. We only focused on collecting fecal samples presumed to be from adult individuals, and between June 2015 and April 2016, a total of 307 samples were collected. The collection was carried out immediately after defecation, using a sterilized plastic spoon attached to the sampling tube. The spoons were inserted into fresh feces, and only a small amount of the interior was removed and stored in 5-ml lysis buffer (0.5% sodium dodecyl sulfate, 100 mM ethylenediaminetetraacetic acid (pH 8.0), 100 mM Tris-HCl (pH 8.0), and 10 mM NaCl). The feces analyzed for gut microbiota were also used for genetic analysis to determine the sex of the host proboscis monkeys, of which 188 females and 78 males were successfully identified.

DNA purification, 16S ribosomal RNA (rRNA) amplification, and sequencing

After bead-beating using the bead crusher (TAITEC, µT-01, Japan) and centrifuged at 4,200 rpm for 5 minutes, 200 µl of lysis buffer-fecal sample mixture was added with 800 µl of InhibitEX buffer of the QIAamp DNA Stool Mini Kit (Qiagen GmbH, Hilden, Germany). Next, the mixture was centrifuged at room temperature for 1 minute at 13,000 rpm. Then, the lysate was transferred to a new 1.5 ml microcentrifuge tube with 25 µl of proteinase K. This was followed by adding 500 µl of Buffer AL and the manufacturer’s protocols to purify the fecal DNA. Next, the DNA concentration was estimated with a Qubit dsDNA HS Assay Kit and a Qubit fluorometer (Thermo Fisher Scientific). We amplified the V3-V4 region of the 16S rRNA gene using the primer utilized in with slight modification as follows: 16S_V34_F 5′-TCGTCGCGCAGCGTCAGATGTATATTAAGAGACAG-Ns-CCTACGGGNGGCWGG-3′ and 16S_V34_R 5′-GTCTCCTGCGGCTCGAGATGTATATTAAGAGACAG-Ns-GACTACHV
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GG-3’. Subsequently, 3Ns, 4Ns, 5Ns, and 6Ns were inserted in each primer between the specific primer and the adapter. KAPA Pure Beads (KAPA Biosystems, Wilmington, MA) were used to purify the polymerase chain reaction amplicons. The Illumina Nextera XT Index Kit (Illumina, Inc., San Diego, CA) was then used to attach specific dual indices and sequencing adapters to the amplicons for each sample. The resulting products were then combined in equal DNA concentrations to form a pooled sequencing library. Subsequently, the size distribution of the library was then estimated using an Agilent 2100 Bioanalyzer (Agilent Technologies, Inc., La Jolla, CA). The library was then diluted to a concentration of 15 pM and sequenced with a 15% PhiX spike-in on an Illumina MiSeq sequencing platform using the MiSeq Reagent Kit v3 (600 cycles) (Illumina, Inc., San Diego, CA). The resulting read lengths were 301 bp (forward sequences), 8 bp (forward indices), 8 bp (reverse indices), and 301 bp (reverse sequences).

**Data analysis**

*Amplicon sequence variants (ASVs) picking and taxonomic identification*

The demultiplexed sequences were processed using QIIME2 software. In order to ensure high-quality data, sequences with a quality score of less than 30 were discarded. Next, using the remaining sequences, ASVs were generated using the DADA2 pipeline in QIIME2. The ASVs were assigned through the ribosomal database project classifier with GreenGenes v13_8 as the reference database for taxonomic identification. Additionally, we used the built-in function align-to-tree-might-fast tree of QIIME2 to construct a phylogenetic tree of the ASVs.

*Statistical analysis*

Statistical analyses were performed in R Version 4.1.1, with the significance level set at 0.05 using the unrarefied dataset. Results were reported as means with standard deviation. Alpha and beta diversity was calculated using the R package *phyloseq*. The Kruskal–Wallis test from the
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R package dunn.test was used to analyze the differences in alpha diversity among groups. Subsequently, we investigated the effects of social and geographical factors on the microbial alpha diversity of individual samples using a linear model. The alpha diversity of each sample was treated as a normally distributed response variable, while the social (number of individuals within groups) and the geographical factor (distance from the river mouth) were treated as explanatory variables. For all models, we verified that the variance inflation factors were smaller than the cutoff value, i.e., less than $10^{49}$. Therefore, the collinearity between independent factors (explanatory variables) did not affect the results. For the model selection, the possible combinations of the explanatory variables were examined and ranked using the Akaike information criterion (AIC) from the MuMIn package. Following the published guidelines for wildlife research, we selected the best-supported models as those with a $\Delta$AIC score $\leq 2$, where $\Delta$AIC = AIC − minimum AIC within the candidate models.

In our multivariate analysis of microbiome composition, we calculated Bray–Curtis dissimilarity, along with weighted and unweighted UniFrac indices with the R package vegan. We constructed nonmetric multidimensional scaling (NMDS) for visualization using Bray–Curtis dissimilarity and principal coordinate analysis (PCoA) plots by weighted and unweighted UniFrac indices, respectively. In order to further assess the effect of social and geographical factors and to test their correlation with microbial beta diversity, Mantel tests were conducted. The social factor, known as ‘demographic distance,’ was calculated based on the composition of the differences in the group and the number of individuals within the groups, while the geographical factor, known as ‘geographical distances,’ was calculated based on the distance from the river mouth.

Results

Phylogenetic profile of the fecal microbiome
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In 307 fecal samples, we detected 18,948 ASV, classified into 27 phyla, 65 classes, 94 orders, 120 families, 137 genera, and 51 species. Figure 1 depicts the general distribution of the top five taxa at the phylum, family, and genus levels. At the phylum level, the top five taxa were consistent across sexes; Firmicutes dominated the gut microbial community (male: 81.8%; female: 82.3%), followed by Bacteroidetes (male: 8.4%; female: 8.0%), Cyanobacteria (male: 1.5%; female: 1.8%), Proteobacteria (male: 1.5%; female: 1.3%), and Actinobacteria (male: 1.1%; female: 1.1%) (see Supplementary Table 1 for details). At the family and genus level, the top five patterns of the gut microbial community in both sexes were also consistent (see Supplementary Table 2 and Table 3 for details), i.e., family level: Ruminococcaceae (male: 46.2%; female: 44.6%), Lachnospiraceae (male: 14.6%; female: 15.3%), S24-7 (male: 6.9%; female: 15.3%), Christensenellaceae (male: 2.0%; female: 2.2%), and [Mogibacteriaceae] (male: 1.9%; female: 1.8%); genus level: Oscillospira (male: 10.5%; female: 10.2%), Ruminococcus (male: 4.7%; female: 4.5%), Dorea (male: 2.3%; female: 2.1%), Blautia (male: 1.2%; female: 1.4%), Clostridium (male: 1.2%; female: 1.2%). In addition, we found that the pattern of the top five taxa (at the phylum, family, and genus levels) was consistent across different group types (harem and non-harem groups), with only minor differences found in solitary males (likely due to small sample sizes) (see Supplementary Table 4-6).

Gut microbial diversity

Alpha diversity

The mean observed richness and Shannon’s $H'$ of the gut microbiome in all fecal samples were $410.3 \pm 59.9$ and $5.0 \pm 0.20$, respectively. There were no significant differences in observed richness between sexes (Kruskal–Wallis $\chi^2 = 2.50$, df = 1, $p = 0.11$; Figure 2a), nor between the group types (harem and non-harem groups and solitary males) (Kruskal–Wallis $\chi^2 = 4.96$, df = 2, $p = 0.08$; Figure 2b). Shannon’s $H'$ did not differ significantly by sex (Kruskal–Wallis $\chi^2 = 1.72$, $p = 0.19$).
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df = 1, p = 0.19; Figure 2c) or group type (Kruskal–Wallis $\chi^2 = 0.751, df = 2, p = 0.69$). Richness values ranged between 73 and 734. In contrast, the Shannon diversity index values ranged between 3.76 and 5.48. The best-fit model to explain the observed richness, as determined by AIC, included both the number of adult females in the harem groups and the location of the collected samples represented as the geographical distance from the mouth of the river (Table 1 and Figures 3a, b), although the AIC value of the following model, which included only the number of adult females, was also <2.0. The observed richness increased as the number of adult females in the harem groups, and the distance from the river mouth increased (upper river). The linear models for Shannon’s $H'$ revealed a similar pattern to the observed richness, with a positive effect on the number of adult females and geographical distance (Table 2 and Figures 3c, d). In contrast, model selection of the model for the investigation of whether the observed richness and Shannon diversity $H'$ were affected by social and ecological factors, i.e., the number of individuals, including adult males and females and subadult males within non-harem groups, and the location of the collected samples, represented as the geographical distance from the river mouth, indicated that the null model was the best. The second- and third-best models, with observed richness and Shannon diversity $H'$, incorporated social and ecological factors with AIC values below the cutoff of 2.0. This result indicates that a similar pattern was observed in nonharem groups, albeit with weaker effects than in harem groups.

*Beta diversity*

According to PCoA, using weighted and unweighted UniFrac and NMDS plots with Bray-Curtis dissimilarity, individuals of different sexes or group types did not exhibit visually distinguishable patterns (Figure 4). Conversely, PERMANOVA analysis revealed weak but significant differences between both sexes (PERMANOVA, Bray–Curtis, $R^2 = 0.00481, p = 0.012$; weighted UniFrac, $R^2 = 0.0023, p = 0.77$; unweighted UniFrac, $R^2 = 0.0042, p = 0.037$) and among groups.
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(PERMANOVA, Bray–Curtis, $R^2 = 0.00956, p = 0.022$; weighted UniFrac, $R^2 = 0.0101, p = 0.047$; unweighted UniFrac, $R^2 = 0.0084, p = 0.005$) on the gut microbiome.

In order to examine the correlation between beta diversity measures and the two factors, i.e., geographical distances and social factors, a Mantel test was conducted separately on a dataset containing only samples from harem groups and one containing only samples from nonharem groups. Mantel tests performed with different beta diversity measures revealed significant correlations between weighted UniFrac and geographical distances from different individuals ($r = 0.09, p = 0.01$) and also between unweighted UniFrac and geographical distances from different individuals ($r = 0.06, p = 0.02$). This indicated that individuals living further apart had a gut microbial community that was less similar, although the explanatory power of geographic distance was relatively weak. In contrast, Bray–Curtis dissimilarity measures of diversity, which do not account for the phylogeny of microbial composition, did not exhibit a significant correlation with geographic distance ($r = 0.01, p = 0.38$). In addition, there was no correlation between the different beta diversity measures and the distance of group composition between the various harem groups (weighted UniFrac: $r = 0.09, p$-value = 0.96; unweighted UniFrac: $r = 0.03, p = 0.18$; Bray–Curtis: $r = 0.01, p = 0.38$). For samples from non-harem groups, unweighted UniFrac ($r = 0.08, p = 0.03$) and Bray–Curtis dissimilarity ($r = 0.20, p = 0.001$) were significantly correlated with geographical distance, whereas there was no significant correlation between weighted UniFrac and geographical distance ($r = 0.03, p = 0.29$). Only Bray–Curtis dissimilarity exhibited a significant correlation with the similarity distance of the group composition between different groups ($r = 0.15, p = 0.02$), while the other measures did not (weighted UniFrac: $r = 0.09, p = 0.18$; unweighted UniFrac: $r = 0.02, p = 0.20$).

**Discussion**
Overall gut microbial composition

The present study was the first comprehensive analysis of the intestinal microbial community based on feces from more than 300 free-ranging proboscis monkeys. At the phylum level, the composition of their gut microbial community was nearly identical to that of other foregut-fermenting primates. Firmicutes and Bacteroidetes were predominant in *Colobus polykomos*, *Procolobus badius*, *P. verus*, *Rhinopithecus brelichi*, and *R. roxellana*[^12^][^52^][^54^], with the exception of *Pygathrix nemaeus*, which includes Bacteroidetes in the top five, but Firmicutes and Tenericutes dominate the top two phylum[^35^]. This trend is not exclusive to foregut-fermenters but is also observed in hindgut-fermenters, such as *Cercocebus atys*, *Cercopithecus campbelli*, *C. diana*, *C. petaurista*, *Chlorocebus sabaeus*, *Lemur catta*, *Macaca fuscata*, *M. mulatta*, *M. thibetana*, *Pan troglodytes*, *Propithecus verreauxi* and *Theropithecus gelada*[^12^][^52^][^55^][^59^]. At the family level, the predominance of Ruminococcaceae, or Lachnospiraceae is comparable to that of the foregut-fermenting *R. brelichi* and *R. roxellanae*. *Macaca fuscata*, a hindgut-fermenting primate, exhibited a similar trend[^55^], and Ruminococcaceae were frequently the most dominant in other hindgut-fermenting primates[^12^][^56^][^57^]. Although the proportion taxonomically assigned at the genus level was limited, the top two genera, *Oscillospira* and *Ruminococcus*, are listed among the top five in *P. nemaeus*, a foregut-fermenter as well as in the proboscis monkey. *Ruminococcus*, in particular, is a common genus, as it frequently ranks first not only in foregut-fermenters but also in hindgut-fermenters[^55^][^57^]. However, *Blautia* is not among the top genera in *P. nemaeus*, but it is among the top genera[^55^][^57^] in the hindgut fermenters. In light of these findings, the composition of the gut microbial community of proboscis monkeys would not be significantly different from that of other foregut-/hindgut-fermenting primates.

Microbial patterns in relation to social factors
There were no differences in the alpha diversity index (number of ASV and $H'$) of the proboscis monkey gut microbiota between sexes or group types, indicating that alpha diversity is not affected by differences in social factors such as sex differences in life history and/or social composition of individuals. Conversely, significant differences in beta diversity were observed between the sexes, suggesting that differences in life history and the frequency of social interactions may have influenced the composition of the gut microbiota. At maturity, females transfer from their natal harem groups to other harem groups, whereas males disperse from their natal harem groups in the early stages to join all-male groups. Additionally, grooming within harem groups occurs primarily between females, with males rarely participating. Consequently, it is likely that these differences in the life histories of the sexes and in the frequency of social interactions between the sexes influence the composition of the gut microbiota in proboscis monkeys. Several other nonhuman primates have been reported to exhibit sexual biases in the gut microbiota, which could be attributed to differences in such interactions and life histories between sexes as a result of group living, e.g., *Callithrix jacchus*, *Rhinopithecus bieti*, and *Alouatta pigra*, though such sexual differences are little in some species, e.g., *Propithecus verreauxi*. In contrast, it is difficult to determine its ecological significance without knowing how differences in microbiota composition affect factors such as food digestion and immunity. As expected, the positive correlation between the number of females in harem groups and the alpha diversity index suggests that increased individual interactions result in an increase in alpha diversity. This result is consistent with previous research indicating that direct physical contact between social partners is a major factor in the transmission of gut microbiota.

**Microbial patterns in relation to geographical factors**

In addition to the number of females within the harem groups, individuals who resided in areas further upstream of the river mouth at our study site had higher alpha diversity indices. This may
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be due to differences in the food diversity consumed by proboscis monkeys in upstream and downstream regions. In general, individuals with a more varied diet have a greater variety of symbiotic bacteria in their gastrointestinal tracks. Indeed, given that the downstream area of the study site is more heavily affected by deforestation and has lower plant diversity (IM uppub data), individuals in the upstream area may have had access to a greater variety of food sources, resulting in a tendency for a higher alpha diversity index in their gut microbiota. In the nonharem groups, a similar relationship was observed between alpha diversity and the number of individuals in the group and the geographical conditions they inhabited, although this trend was weaker with insignificance. This difference may be attributable to the fact that nonharem groups, particularly all-male groups, exploit a broader range of riparian habitats than harem groups, i.e., the bias in consumed food items may be less pronounced than in the harem group. However, the social relationships between individuals within nonharem groups remain unclear, making further discussion impossible at this time. The key to advancing this discussion in the future would be the collection of additional ecological and social observation data on non-harem groups.

There was a weaker but significant correlation between the physical distance between individuals and their similarity in the composition of their intestinal microbial communities, regardless of a group type. It should be noted, however, that the significance of the correlation varied slightly between the various group types based on the various indices, namely weighted UniFrac, unweighted UniFrac, and Bray–Curtis. As previously mentioned, differences in vegetation between upstream and downstream areas of this study site may influence the alpha diversity of individual proboscis monkeys; it cannot be ruled out that the physical distance between individuals may lead to differences in diets consumed, which may, in turn, lead to differences in the composition of the intestinal microbial community between individuals. However, no correlation was found between similarities in group composition and similarities in gut microbial community composition. Compared to seasonal and clumped food sources such as fruits and
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flowers, the higher ubiquitous and abundant availability of leaves as the main food source for proboscis monkeys in this study site and the lack of a clear hierarchy between individuals, which occurs particularly within harem groups. This may have contributed to the lower likelihood of dietary bias across group composition in proboscis monkeys, and hence there might be no differences in the similarity of the composition of gut microbial community between individuals. Conversely, there have been no studies on the hierarchy among males in nonharem groups, particularly in all-male groups, and the finding that significant differences were only observed in nonharem groups by Bray–Curtis dissimilarity may suggest that there may be a severe hierarchy among males, which resulted in differences in their dietary composition. To verify this, however, more comprehensive behavioral observations of all-male groups are required.

**Outlook**

We were interested not only in determining the general trends in the intestinal microbiota of proboscis monkeys but also in determining how social and geographical factors affected this microbial community. In contrast, the biological significance of such differences in the intestinal microbial community between individuals and between groups could not be determined. Future research must examine how differences in the gut microbial community between individuals influence their feeding strategy through more detailed functional analysis, such as by isolating and cultivating characteristic strains. Lastly, the present study suggested that the degree of deforestation may influence the gut microbiota, although quantitative comparisons of these effects were not possible. With advancements in technology for the analysis of gut microbiota, it may be possible in the future to transition from non-invasively collected free-ranging primate feces to studies that contribute to animal conservation, such as assessing the effects of forest disturbance. Multiple studies have demonstrated their potential use in animal conservation.
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Author contribution

B.G., I.M. and V.S.K. conceptualized the initial idea; I.M. collected fecal samples; A.T. and J.T. arranged the sampling in the field; L.J., G.H., and W.L. conducted a genetic experiment. I.L, W.L., and I.M. performed and interpreted the statistical analyses; L.J., I.M., and V.S.K. drafted the manuscript. All authors contributed to the final version of the manuscript.

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Declarations

Conflict of interest: the authors declare no competing interests.

Ethics approval: This study was reviewed and approved by the Sabah Biodiversity Council (Access License Reference: JKM/MBS.1000-2/2 JLD.5) and was conducted in compliance with animal care regulations applicable to Malaysian laws.
Data Availability statement

The original contributions presented in the study are included in the article/Supplementary Material, further inquiries can be directed to the corresponding author/s. The sequencing data have been deposited to NCBI under PRJNA928786.
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https://doi.org:10.1007/s10329-011-0259-1


https://doi.org:10.3106/041.041.0201
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Table and figure legends

Table 1 Summary of model selection using linear models. This method was used to investigate whether observed richness (A) and Shannon diversity $H$ (B) were affected by social and ecological factors. An analysis of the number of adult females within harem groups and the location of samples collected was conducted. These represent the geographical distance from the mouth of the river, respectively (only models with $\leq$AIC 2 are shown).

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Figure 1 The general pattern of the top five taxa. These were analyzed at the phylum (a), family (b), and genus (c) levels.

Figure 2 Comparison of the mean observed richness (a) and the Shannon's $H'$ (b) for different sexes and different groups. The central line of the box represents the median, and the lower and upper bounds of the box represent the first and third quartiles.

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**Figure 4** Unweighted UniFrac principal coordinate analysis (PCoA) plot. This was analyzed by sex by group type (a); unweighted UniFrac PCoA plot, i.e., by sex by group type (b); and Bray–Curtis nonmetric multidimensional scaling plot (c).
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<th>Intercept</th>
<th>Adult female</th>
<th>Distance from river mouth</th>
<th>df</th>
<th>Log-likelihood</th>
<th>AIC</th>
<th>$\Delta$AIC</th>
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(B)

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