Disordered intestinal microbes are associated with the activity of Systemic Lupus Erythematosus

Running title: Intestinal microbes in systemic lupus erythematosus

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Abstract

Intestinal dysbiosis is implicated in Systemic Lupus Erythematosus (SLE). However, the evidence of gut microbiome changes in SLE is limited, and the association of changed gut microbiome with the activity of SLE, as well as its functional relevance with SLE still remains unknown. Here, we sequenced 16S rRNA amplicon on fecal samples from 40 SLE patients (19 active patients, 21 remissive patients), 20 disease controls (Rheumatoid Arthritis patients), and 22 healthy controls, and investigated the association of functional categories with taxonomic composition by Phylogenetic Investigation of Communities by Reconstruction of Unobserved States (PICRUSt). We demonstrated SLE patients, particularly those active patients, had significant dysbiosis in gut microbiota with reduced bacterial diversity and biased community constitutions. Among the disordered microbiota, the genera *Streptococcus, Campylobacter, Veillonella*, the species *anginosus* and *dispar*, were positively correlated with lupus activity, while the genus *Bifidobacterium* was negatively associated with disease activity. PICRUSt analysis showed metabolic pathways were different between SLE and healthy controls, and also between active and remissive SLE patients. Moreover, we revealed that a random forest model could distinguish SLE from RA and healthy controls (AUC = 0.792), and another random forest model could well predict the activity of SLE patients (AUC = 0.811). In summary, SLE patients, especially the active patients, show an apparent dysbiosis in gut microbiota and its related metabolic pathways. Among the disordered microflora, 4 genera and 2 species are associated with lupus activity. Furthermore, the random forest models are able to diagnose SLE and predict disease activity.
**Keywords:** Systemic lupus erythematosus; gut microbiota; disease activity; PICRUSt; Random forest

**Introduction**

The gut microbe population, known as “gut microbiota” is heterogeneous and complex, and is composed of more than 1000 different bacterial species (1). Intestinal mucosal immunity have clarified the correlation between the gut microbiota and the host immune system (2). Microbial abnormalities, also known as “dysbiosis”, is thought to be correlated with various diseases, including chronic kidney disease, obesity, type 2 diabetes, atherosclerosis and nonalcoholic fatty liver disease (3-6). Systemic Lupus Erythematosus (SLE) is a heterogenic autoimmune disease promoted by a combination of genetic and environmental factors that bring about an intolerance towards self-antigens (7). Although the etiology of SLE remains unclear, hormonal, environmental and genetic factors are thought to be of importance. Recently, dysbiosis of the gut microbial community in the development of SLE has attracted attention.

Multiple evidences has shown a lower *Firmicutes/Bacteroidetes* ratio and decreased abundance of some families in *Firmicutes* phylum may be involved in remissive SLE (8-10) However, such alterations are also discovered in Intestinal Mucositis and Crohn’s disease (11). It was reported that the presence of *Lactobacillus spp.* in gut could attenuate kidney inflammation in lupus-prone mice in a sex hormone-dependent manner (12), suggesting the gut microbiome may be a possible therapeutic target for SLE. The association of the gut microbiome with different diseases has been shown to be diverse due
to many factors such as host’s age, sex, genotype, diet and geography (13-16). Therefore, alterations of gut microbiome associated with SLE should be variable in SLE patients in Guangdong Province, China compared with other locations.

So far, there are limited studies on SLE and gut microbiota. Only a study observed the gut microbiota in active SLE patients, other studies have just focused on SLE patients in remission (8-10)(17). Whether the gut microbiota is associated with the disease activity still remains unclear. In this study, we recruited both active and remissive SLE patients to investigate the characteristic of intestinal microbes that are associated with disease activity. Since Rheumatoid Arthritis (RA) is another common autoimmune disease, we also included RA patients as the disease control to define the specificity of the SLE-associated gut microbiome. We found that the gut microbiota in SLE patients, especially in active SLE patients, had a distinct dysbiosis in microbiota and its related metabolic pathways. Six disordered genera and two species were revealed to be closely associated with SLE activity. Furthermore, the results suggested the gut microbiota were validated to have strong diagnostic potential for SLE, and even predict the disease activity through random forest analysis.

**Materials and Methods**

**Research participants and sample collection**

40 SLE patients, 20 RA patients and 22 healthy controls were consecutively recruited from Nanfang Hospital, Southern Medical University during 2017. All SLE and RA patients fulfill the American
College of Rheumatology (ACR) classification criteria for SLE or RA disease (18-20). All patients with acute intercurrent illnesses or infections and those who used probiotics or antibiotics within 1 month before admission were excluded (6). The gender- and age-matched healthy controls (HC) who had no known history of autoimmune diseases were also recruited from the Health Examination Centre of Nanfang Hospital. All the participants were female. Average age of SLE, RA and HC group was 37.46 ± 14.17, 44.00 ± 6.53, and 37.18 ± 14.67 respectively (P = 0.142).

Based on the systemic lupus erythematosus disease activity index (SLEDAI) (21), all the SLE patients were divided into the active SLE patients (A) (SLEDAI ≥ 8) (n = 19) and remissive SLE patients (R) (SLEDAI < 8) (n = 21). Exception of the age and gender distribution, patients in group A showed many significant differences from that of group R, having more severe symptoms, including anemia, hypocomplementemia, impaired renal functions and increased autoantibodies, all of which are consistent with the clinical characteristics of SLE (Table 1).

For all participants, the fresh fecal samples were frozen at -80 °C immediately after collection. Ethics approval was granted by the Ethics Committee of Nanfang Hospital, and all of the methods used were in accordance with the approved guidelines. Written informed consent was required from all patients and healthy volunteers in the study.

**Illumina Miseq sequencing of 16S rRNA gene-based amplicons and data processing**

Total DNA was extracted from thawed fecal samples using the LONGSEE STOOL DNA KIT (Longsee...
med Bio Medical., LTD., Guangdong, China) following the manufacturer’s instructions. All the individually processed human fecal DNA extractions were amplified by polymerase chain reaction (PCR). The forward primer (5’-ACT CCT ACG GGA GGC AGC AG-3’) and reverse primer (5’-GGA CTA CHV GGG TWT CTA AT-3’) were used to amplify the 16S rRNA gene V3-V4 variable region from the bacteria by polymerase chain reaction (PCR) as described previously (22). Briefly, amplifications were performed using a step cycling protocol consisting of 98 °C for 30 s, 35 cycles of 98 °C for 10 s, 54 °C for 30 s, and 72 °C for 45 s, ended with the final elongation at 72 °C for 10 min. PCR products were purified using an AxyPrep PCR Cleanup Kit (Axygen, California, USA).

For the sequencing of 16S rRNA gene-based amplicons, the amplicon library was prepared using a TruSeq Nano DNA LT Library Prep Kit (Illumina Inc, CA, USA). The sequencing reaction was conducted using Illumina MiSeq platforms and the data were analyzed by the Quantitative Insights Into Microbial Ecology platform (QIIME, www.qiime.org) using the default parameters (23). The raw sequence data for 16S rRNA gene sequencing data sets was available from the Sequence Read Archive (SRA) database (http://www.ncbi.nlm.nih.gov/sra) at accession number PRJNA493726.

Before assembly, sequence reads were first filtered to remove low-quality or ambiguous reads, including reads lacking exact matching with the primer, sequences with mismatch ratio sequences higher than 0.05 in the overlap region and raw reads shorter than 100 bp with Trimmomatic v.0.32 software (24). Paired-end clean reads were merged using FLASH (25) according to the relationship of the overlap between the paired-end reads when at least 10 of the reads overlapped the read generated from the
opposite end of the same DNA fragment, the maximum allowable error ratio of an overlap region of 0.2, and the spliced sequences were called raw tags.

High-quality Sequences with a distance-based similarity of 97% or greater were grouped into operational taxonomic units (OTUs) using the Vsearch algorithm. Representative sequence was then extracted from each OTU. Next, the chimeric sequences were detected and removed. To assign taxonomy information to each clustered feature, extracted representative sequences were subjected to similarity search against Greengenes sequence and taxonomy database using RDP classifier algorithm (ucluster approach with default settings) and the classify-sklearn plugin within QIime software (version 1.9.1). The phylogenetic relationships were determined based on a representative sequence alignment using Fast-Tree (26). Computation of α-diversity metrics and β-diversity metrics were performed on all samples within the feature table with QIime diversity alpha/beta plugin. Rarefaction curve plots the number of individual’s sample versus the number of species, which was done with QIime diversity alpha-rarefaction plugin. Rank abundance curve portray relative abundance and species diversity within a community by plotting relative abundance of species (y-axis) against their rank in abundance (x-axis), which plotted using QIIME v.1.9.1 software.

Phylogenetic Investigation of Communities by Reconstruction of Unobserved States (PICRUSt) is a bioinformatics software package designed to predict metagenome functional content from marker gene surveys and full genomes. PICRUSt analysis was performed to identify Kyoto Encyclopedia of Genes and Genomes (KEGG) metabolic pathways, and determine functional categories associated with
Comparisons of relative abundance of taxa between groups were performed using Linear discriminant analysis Effect Size (LEfSe), a non-parametric Mann-Whitney U test applied to detect features with significant differential abundance with respect to the groups compared, followed by a Linear Discriminant Analysis (LDA) to estimate the effect size of each differentially abundant feature in Linux platform (28).

**Statistical analyses**

We used the mean (±SD) to express measurement data that obeyed a normal distribution, the median (interquartile range) to express measurement data that obeyed a skewed distribution, and a percentage to express enumeration data. Mann-Whitney U test or Student t test was performed to compare the variables of 2 sample groups. Multiple group comparisons were made by the Kruskal-Wallis test or one-way analysis of variance. False discovery rate (FDR) correction for multiple comparisons was employed, and the statistical power was analyzed via power and sample size calculation in R software (29, 30), then the False Discovery Rate q-value was calculated.

The α-diversity determines the species richness and evenness within bacterial populations. The α-diversity metrics include: Observed species and Chao1 (microbial richness), and Shannon index and Simpson index (microbial diversity) (31). The β-diversity determines the shared diversity between bacterial populations. Different distance metrics reveal distinctive views of community structure.
UniFrac distances measure the shared phylogenetic diversity between communities. A smaller UniFrac distance between two samples indicates a higher similarity among the two microbial communities (32). Principal coordinates analysis (PCoA) was plotted using the package in R software (Version 3.4.4). The Wilcoxon rank sum test was used to determine significance in $\alpha$-diversity and $\beta$-diversity.

We used spearman algorithm to analyze the relationship among microbiota, predicted pathways and SLE activity index. The Random Forest models were trained by “randomForest” package with default parameters in R, then the performance of the model was assessed with a ten-fold cross-validation approach and measured by area under the receiver-operating characteristic (ROC) (33). All tests were performed using GraphPad Prism (v6.0) (GraphPad Software, Inc., CA, USA), SPSS Statistics (V.24.0.0.0) (SPSS Inc., Chicago, USA) or R software (Version 3.4.4).

Results

Characteristics of 16S rRNA sequences.

A total of 82 samples were subjected to 16S rRNA sequencing. These samples were composed of three groups including 22 healthy individuals, 20 RA patients and 40 SLE patients. We obtained 2182143 16S rRNA sequencing reads from stool samples of SLE patients, 976140 reads from RA patients and 1277858 reads from HC, which belong to 714 kinds of operational taxonomic unit (OTUs). The parameters, including Chao1 rarefaction curves, Shannon rarefaction curve, and rank abundance of OTUs, were evaluated to confirm the reliability of the sequencing data (Supplementary Figure S1).
**Difference of the gut microbiota in SLE patients from those of controls**

The α-diversity between two groups was compared using Chao1, Observed species, Shannon index and Simpson diversity indices. Overall, the α-diversity metrics Chao1 and Observed species were significantly higher in healthy controls than in SLE patients (P = 0.038; P = 0.004, respectively), indicating that the gut microbiome in SLE patients exhibited a lower richness than healthy controls (Figure 1A and B and Supplementary Table S1). However, no difference in Shannon and Simpson index (P = 0.089; P = 0.092, respectively) was observed between SLE patients and healthy individuals, suggesting that the evenness of the gut microbiome of the two groups had no significant difference (Supplementary Table S1). There were no associations between α-diversity and drug treatments, such as Hydroxychloroquine, Glucocorticoid, Cyclophosphamide, and Biological agent (Supplementary Figure S2).

To measure the extent of the similarity of fecal microbial communities, β-diversity was calculated using unifrac distances. Principal coordinate analysis (PCoA) based on weighted and unweighted UniFrac distance matrix were used for visualizing sample relationships, and ADONIS analysis was used to test the homogeneity of dispersion among different groups. Our results suggested that there were no associations between β-diversity and medicine treatments, including Hydroxychloroquine, Glucocorticoid, Cyclophosphamide, and Biological agent (Supplementary Table S2 and Figure S3), however, the unweighted UniFrac distance analysis of β-diversity difference demonstrated that the
structure of microbiota of SLE patients differed from healthy controls (ADONIS analysis, $P < 0.001$, $R^2 = 0.054$) (Figure 1C and Supplementary Table S3). Thus, the microbial diversity was significantly different between SLE group and healthy controls.

We then analyzed the phylum-level profiles of feces between SLE patients and healthy controls. The phylum level profiles for gut microbiota of SLE patients and controls were fairly similar, except for reads from the phyla *Fusobacteria* ($P = 0.027$) and *Tenericutes* ($P = 0.002$) (Figure 1D-F). A lower *Firmicutes* / *Bacteroidetes* (F/B) ratio was reported in the feces of remissive SLE patients compared to healthy controls (34). However, we showed the ratio of F/B the feces of SLE had a decreasing trend but no significant difference compared with HC group (Supplementary Table S4).

To further determine the phylogenetic clustering pattern between these two groups, the logarithm linear discriminant analysis (LDA) was performed (Figure 2). The phylum *Tenericutes*, along with *Mollicutes* and *RF39*, were significantly reduced in the intestinal flora of SLE patients compared with healthy controls. In addition, patients with SLE exhibited a significant decrease in the genus *Faecalibacterium* alongside its species *praunstizii*, while the taxonomic clade *Cryptophyta* and genus *Roseburia* were reduced in the gut microbiota of SLE group. On the contrary, the taxonomic clade *Bacilli* from the phylum *Firmicutes* showed clustered differences, while *Streptococcaceae* and *Lactobacillaceae* were expanded in the feces of SLE patients compared with healthy controls. Moreover, the genera *Streptococcus* and *Lactobacillus*, along with their species *Streptococcus. anginosus* and *Lactobacillus. mucosae* were enriched in the intestinal flora of SLE group compared with HC group. In
addition, the feces of SLE patients showed an increase in genus *Megasphaera* (significant taxa \[p<0.005, \text{Kruskal–Wallis test}\] with LDA score >2 were shown). Taken together, sequence profiling of the gut microbiota revealed an apparent dysbiosis of the gut microbiota in SLE patients, which was characterized by reduced bacterial α-diversity and biased community constitutions. These results demonstrated the gut microbiota of patients with SLE differed from those of healthy controls.

To investigate whether the disordered intestinal microbes were specific to SLE patients, we further compared the intestinal microflora distribution between SLE and RA patients. There were no significant difference in α-diversity (Supplementary Table S1) and β-diversity (Supplementary Table S3 and Figure S4C-D) between two groups. LEfSe analysis showed the different microbiota between SLE group and RA group (Supplementary Figure S4). The taxonomic clade *EB1017, Ellin6529*, and *Anaerofilum* were increased in the intestinal flora of RA patients, while the *Lactobacillales* from the *Bacilli*, with its genus *Streptococcus* were enriched in the feces of SLE patients compared with RA patients. In addition, the phylum *Fusobacteria*, along with its taxonomic clade *Fusobacteriia, Fusobacteriales, Fusobacteriaceae, and Fusobacterium* were increased in the feces of SLE patients. The genus *Megasphaera* and *Veillonella*, with its species *Veillonella. dispar*, were also enriched in the feces of SLE group compared with RA group (significant taxa \[p<0.05, \text{Kruskal–Wallis test}\] with LDA score >2.5 were shown).

Collectively, these results demonstrated that the gut microbiota of SLE patients differed from healthy individuals, however, there was no significant difference in gut microflora diversities between SLE and RA patients. The genera *Streptococcus* and *Megasphaera* were specifically increased in the
feces of SLE patients compared with healthy controls and RA patients.

**Difference of microbiota profiling in active SLE patients from remissive SLE patients.**

Given that the gut microbiota was significantly different between SLE patients and healthy controls, we next investigated whether the gut microbiota was associated with disease activity of SLE. Firstly, we compared 16S rRNA sequences of A group (active SLE patients) with R group (remissive SLE patients). The unweighted UniFrac distance analysis of β-diversity difference demonstrated that the structure of the microbiota of A group differed from R group (ADONIS analysis, P =0.047, R² = 0.039) (Figure 3A and Supplementary Table S3), while no obvious difference was observed in α-diversity (Supplementary Table S1), suggesting that the community constitutions in A group were distinctly different from R group, but no difference was found in microbial diversity.

As shown in Figure 3, LEfSe analysis further demonstrated that *Actinomycetales* and *Bifidobacteriales* from phylum *Actinobacteria* showed clustered differences, and the genus *Bifidobacterium* was increased in the feces of remissive SLE patients compared with active SLE patients. In addition, the species *Ruminococcus. gnavus* was reduced in the feces of active SLE patients, whereas *Lactobacillales* from the Bacilli, along with its genus *Streptococcus* and species *Streptococcus. anginosus*, were enriched in the feces of A group compared with R group. Moreover, the genus *Oribacterium* was increased in the intestinal flora of active SLE patients. Furthermore, active SLE patients exhibited a remarkable enrichment of the taxa *Epsilonproteobacteria* from phylum
Proteobacteria, along with its Campylobacterales and Campylobacter (significant taxa [p<0.005, Kruskal–Wallis test] with LDA score >2 were shown). Finally, the ratio of F/B in remissive SLE patient group had a decreasing trend but no significant difference compared with HC group (Supplementary Table S4). Altogether, these results indicated that the gut microbiota profiling of active SLE patients were markedly different from that of remissive SLE patients.

Aberrant microbiome-associated pathway is correlated with the activity of SLE patients.

Another emphasis of our study was to disclose the functional variation in the SLE gut microbiota community. Therefore, we predicted the microbiota-derived pathways using the PICRUSt algorithm with the KEGG database and compared functional abundances among the SLE, RA, and HC groups. In total, we characterized six different pathway categories between SLE group and HC group (Figure 4). The pathways of Apoptosis and Purine metabolism were significantly increased in SLE patient group compared with HC group (Figure 4A), while four pathways, including Pathways in cancer, Bacterial chemotaxis, Bacterial motility proteins, and Flagellar assembly, were decreased in SLE patients (Figure 4B). In addition, nine different functional pathways were identified between A group and R group (Figure 5). Five were related to Synthesis and degradation of ketone bodies, Apoptosis, Lipid metabolism, Secretion system, and Staphylococcus aureus infection, which were significantly higher in active SLE patients than remissive patients (Figure 5A). Conversely, Alanine aspartate and glutamate
metabolism, Carbohydrate metabolism, Primary bile acid biosynthesis, and Secondary bile acid biosynthesis, were obviously increased in remissive SLE patients compared with active patients (Figure 5B). However, there was no different pathway between SLE and RA group (date not shown).

We further examined correlations among SLE/HC-associated taxa and disordered functional pathway to obtain an overview of how specific taxa act during metabolic dysfunction in patient gut. For SLE patients, we characterized a positive correlation between the enrichment of *Streptococcus* and increased Apoptosis pathway ($r = 0.807$, $P < 0.000$, FDR < 0.000) and a negative correlation between *Streptococcus* and Pathways in cancer ($r = -0.550$, $P < 0.000$, FDR < 0.000) (Figure 6A). Further analysis also revealed the active SLE patient-enriched genus *Streptococcus* was negatively associated with pathways of Alanine aspartate and glutamate metabolism, Primary and secondary bile acid biosynthesis ($r = -0.680$; $r = -0.437$; $r = -0.434$, $P < 0.01$, FDR < 0.05, respectively) (Figure 6B), but positively associated with five increased pathways, including Synthesis and degradation of ketone bodies, Apoptosis, Lipid metabolism, Secretion system, and Staphylococcus aureus infection ($r = 0.574$; $r = 0.829$; $r = 0.406$; $r = 0.486$; $r = 0.903$, $P < 0.01$, FDR < 0.05, respectively) (Figure 6C).

Thus, several aberrant microbiome-associated gut metabolic pathways were associated with SLE using PICRUSt analysis. Interestingly, the SLE-enriched genus *Streptococcus* was positively associated with the pathways of Apoptosis, the metabolism of lipid, amino acid and bile acid, Secretion system, and pathogenic bacteria infection.
Association of disordered microbiota and aberrant microbiome-associated pathway with activity of SLE.

SLEDAI, Complement C3, C reactive protein (CRP), Erythrocyte Sedimentation Rate (ESR) and anti-double stranded DNA (anti-dsDNA) were commonly used to indicate the disease activity of SLE patients (21, 35, 36).

At genus and species levels, *Lactobacillus*, *Streptococcus*, *Megasphaera*, *Fusobacterium*, *Veillonella*, *Lactobacillus. mucosa*, *Streptococcus. anginosus*, and *Veillonella. dispar* were increased in the feces of SLE patients compared with healthy controls or RA patients. Meanwhile, *Streptococcus*, *Oribacterium*, *Campylobacter*, and *Streptococcus. anginosus* were enriched, but *Bifidobacterium* and *Ruminococcus. gnavus* were reduced in the gut microbiota of active SLE patients compared with that of remissive SLE patients. Except *Bifidobacterium*, five changed genera were positively associated with disease activity (Figure 7A-F and Supplementary Table S5). For example, the abundance of *Streptococcus* was positively correlated to SLEDAI ($r = 0.492$, FDR $q = 0.008$), while negatively associated with Complement C3 ($r = -0.502$, FDR $q = 0.008$) (Figure 7A). *Campylobacter* and *Streptococcus. anginosus* also showed a positive correlation with SLEDAI ($r = 0.470$, FDR $q = 0.009$; $r = 0.388$, FDR $q = 0.040$, respectively) (Figure 7). Moreover, the abundance of *Veillonella* and its species *Veillonella. dispar* showed negative correlations with Complement C3 ($r = -0.475$, FDR $q = 0.008$) (Figure 7).

The genus *Streptococcus*, which was specifically associated with the activity of SLE, was related to
eight aberrant microbiome-associated pathways (Figure 6). We further explored whether these eight disordered pathways were also related to the activity of SLE (Supplementary Table S5). Alanine aspartate and glutamate metabolism, Secondary bile acid biosynthesis, and Lipid metabolism were closely associated with SLEDAI (r = -0.376; r = -0.382; r = 0.318, FDR q <0.001, respectively) (Figure 7G-I). As such we hypothesized that the genus *Streptococcus* might play an important role in the disease progression of SLE through these three pathways.

**Potentials of gut microbiota for SLE diagnosis or disease activity monitoring**

Given that the gut microbiota in SLE patients, especially in active SLE patients, had a distinct dysbiosis in microbiota, we next addressed the potential diagnostic value of the gut microbiota as potential biomarkers for SLE by ROC curve analyses. Due to its non-parametric assumptions, random forest was used to detect linear and nonlinear effects and potential taxon–taxon interactions, to identify taxa that could differentiate SLE subjects from control subjects (healthy controls and RA patients), and to discriminate active SLE patients from remissive patients. We used 10-fold cross-validation approach to evaluate the performance of model, and predictive power was scored in ROC analysis. We first made the mode to differentiate the SLE patients from healthy controls and RA patients based on the genus and species levels. We showed that the area under the curve (AUC) was 0.792 (95% CI: 0.750–0.835) (Supplementary Table S6 and Figure 8A), suggesting that the gut microbiota had the potential to diagnose SLE from healthy and disease controls (RA patients). We observed that in the model, out of the
top 10 genera and species, 8 belonged to the phylum *Firmicutes*, 1 belonged to *Fusobacteria*, and 1 belonged to *Actinobacteria*. Of the 8 genera in the *Firmicutes* phylum, 5 were part of *Clostridia* class, and 3 were *Bacilli* (Supplementary Table S6). Furthermore, among the 10 genera and species, the *mucosa*, *Lactobacillus*, *Megasphaera*, and *Streptococcus* were significantly enriched, while *Faecalibacterium* was decreased in the feces of SLE patients compared with healthy controls. In addition, both *Veillonella* and *Fusobacterium* were increased in the gut microbiota of SLE patients than RA patients (Figure 2; Supplementary Figure S4 and Table S5). Accordingly, most of the genera and species in the model were the disordered genera in the feces of SLE group compared with healthy controls and RA patients.

We further built another model to distinguish active SLE patients from remissive patients based on the genus and species levels. In this model, the AUC was 0.811 (95% CI: 0.754-0.869) (Supplementary Table S7 and Figure 8B), suggesting that the gut microbiota had the potential to monitor the activity of SLE. Anti-dsDNA was reported to be reasonably sensitive and specific in the diagnosis of SLE, and raised titers of anti-dsDNA along with hypocomplementemia were associated with the activity of SLE(37). We showed that the AUC value for combination of Complement C3 and anti-dsDNA was only 0.773 (95% CI: 0.597–0.949) (Supplementary Figure S6). These results indicated that the combination of the gut microbiota might have a better surveillance value for SLE activity than the combination of Complement C3 and anti-dsDNA. Moreover, as shown in the model, out of the top 10 genera and species, 5 belonged to the phylum *Firmicutes*, 4 belonged to *Actinobacteria*, and 1 belonged to *Proteobacteria*. 

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Among the 5 genera in the *Firmicutes* phylum, 3 were from the *Clostridia* class, 1 was *Erysipelotrichi* and 1 was *Bacilli* (Supplementary Table S7). In this case, the *Campylobacter*, *Streptococcus*, and *Oribacterium* were enriched, while the *gnavus* and *Bifidobacterium* were reduced in active SLE patients compared with remissive SLE patients (Figure 3 and Supplementary Table S7). Altogether, a great part of the genera and species in the model were disordered genera in the feces of active SLE patients, suggesting that the disordered intestinal flora might have potential to diagnose SLE, even monitor disease activity.

**Discussion**

SLE is an autoimmune disease that affects multiple tissues, and causes joint pain, renal disease, muscle pain, fever, poor circulation, inflammation, fatigue, loss of appetite and other symptoms (38). Though the cause of SLE still remains unclear, it is thought to be involved with hormonal, genetic and environmental factors (39). The gut microbiome was believed to be a key factor in influencing predisposition to autoimmunity diseases (40). Recent studies further supported that gut microbiome dysbiosis could act as an important factor in promoting chronic inflammation into autoimmune diseases (2, 41, 42). However, there were only limited works in exploring the potential relationship of gut microbiome with SLE (8-10, 17, 39). In this study, we have provided new evidence about the gut microbiome dysbiosis in female SLE patients by fecal bacteria sequencing. Importantly, we for the first time explored whether the gut disordered microbes were associated with the activity of SLE.
We investigated the profiling of the gut microbiota and showed a distinct dysbiosis of the gut microbiota in SLE patients, which was characterized by reduced bacterial α-diversity and biased community constitutions. Most of the patients in our study were currently on various immunosuppressants and glucocorticoids treatments. Veena Taneja et al. have demonstrated that RA patients using methotrexate (MTX) and hydroxychloroquine exhibited an increase in species richness and diversity (43). However, our results showed no significant relationship between drug treatments and the abundance diversity of gut microbiota in the SLE patients, which might because the most of enrolled patients were treated with steroids or immunosuppressants, while only 4 patients did not use any drugs.

Phyla *Firmicutes* together with *Bacteroidetes* usually account for more than 90% of all phylogenetic species, were involved in host metabolism and immunity (44). In our study, the Phyla *Firmicutes* and *Bacteroidetes* occupied the most abundant microorganism, consistent with the typical human intestinal microbiome structures. It was reported that the *Firmicutes*/*Bacteroidetes* ratio was significantly lower in the feces of SLE patients in remission (8). However, no significant different for our cohort of remissive SLE patients and healthy controls (P > 0.05). Also, there was no significant difference in the *Firmicutes*/*Bacteroidetes* ratio between active SLE and healthy controls (P > 0.05), consistent with the available data (17). The changes of the genera in SLE patients of our study were only partly consistent with previous studies (8, 10, 17, 45), which might partially due to the sample size and geographical locations of patients. It is well known that cohorts with different patient characteristics, including disease stage, geographical locations, diet and status, might exhibit different gut microbiota
profiling (15, 16, 46-48). Therefore, the alterations of gut microbiome associated with SLE should display differences among different geographical locations and disease status.

In this study, we found that the abundance of pathogenic genus *Streptococcus*, with its species *anginosus*, and genus *Megasphaera* were significantly enriched in the feces of SLE patients compared with healthy controls; genus *Streptococcus* and its species *anginosus* were positively correlated to the activity of SLE. In addition, the genus *Veillonella* and its species *dispar* were significantly increased in the gut microbiota of SLE patients compared with RA patients and had a positive association with the activity of SLE. The association of these disordered genera with the activity of SLE was most striking, and to our knowledge, this is the first study to describe such a significant relationship with SLE. The genera *Streptococcus* and *Megasphaera* were reported to be closely related to the intestinal disturbance of autoimmune disorders. For example, *Streptococcus* and *Megasphaera* were enriched in primary biliary cirrhosis (49) and Pediatric Autoimmune Neuropsychiatric Disorders (50). Also, *Streptococcus* was relatively increased in RA patients (43). It was demonstrated that *S. anginosus* rarely caused infections in healthy individuals, but caused infections in the immunodeficient individuals (51). As reported, genera *Streptococcus* and *Veillonella* had pro-inflammatory effects. For example, the combination of *Streptococci* with *Veillonella* appeared to negate IL-12p70 production, while augment IL-8, IL-6, IL-10, and tumor necrosis factor alpha (TNF-α) response (52).

The SLE patients, especially the active patients, had an increased population of oral bacteria, which is an interesting phenomenon that occurred in the intestinal flora of SLE. However, the gut
microbiome of liver cirrhosis, colorectal cancer, RA, and ACVD patients also showed an increase in the abundance of oral bacteria in gut microbiota (53), and only RA and ACVD have been epidemiologically associated with periodontitis. Interestingly, our results suggested that the abundance of genus *Streptococcus* were enriched in active SLE patients, suggesting that the oral microbiota might be overrepresented in the lower gastrointestinal populations of patients with active SLE. Besides, more severe forms of periodontitis were found in SLE subjects that had higher bacterial loads (54), resulting in an increase in oral bacteria entering the intestine.

Furthermore, our data showed that many beneficial commensal microbes, such as *Roseburia*, *Faecalibacterium* and its species *prausnitzii* were depleted in SLE patients. Meanwhile, the genus *Bifidobacterium* was adversely correlated with activity of SLE. These microbes belongs to the phylofunctional core of the intestinal microbiota (55, 56), which can produce short chain fatty acids (SCFAs), especially butyrate-acid, to play multiple critical roles in the maintenance of human health, including producing energy components and intestinal epithelial nutrition (57), reducing the severity of inflammation (58), maintaining intestinal barrier functions (59) and enhancing colon motility functions (60).

Moreover, we observed an increased abundance of beneficial commensal genus *Lactobacillus* and its species *Lactobacillus mucosae* in the feces of SLE cohort compared to healthy controls. Supportive of a role for *Lactobacilli* in the pathogenesis of lupus, taxa in this genus were found to be enriched in female NZB/W F1 mice, the model of systemic lupus. In this study, *Lactobacillus spp.* were
associated with more severe disease, whereas they were reduced as disease is controlled with
dexamethasone (17). As reported (61) Lactobacillus reuteri increased over time in the feces of mice
from both lupus models as their disease progress, in addition, Lactobacillus spp. were increased in a
longitudinal cohort of SLE patients compared with healthy controls. In this study, the
pDC/IFN-promoting properties of L. reuteri in the context of a lupus-prone host suggest a paradigm in
which a bacterium that is normally considered a probiotic may become harmful under certain genetic or
environmental conditions. We also observed that Lactobacillus were enriched in feces of SLE patients,
suggesting a potential role for these taxa in SLE pathogenesis, which need further research in the future.

Our study has demonstrated that some pro-inflammatory bacteria in genera Streptococcus, and
Campylobacter expanded, while some anti-inflammatory bacteria in genera Roseburia, Faecalibacterium, and Bifidobacterium reduced in the feces of SLE patients, especially the active
patients, resulting in the release of inflammatory factors, then aggravating the systemic inflammation
level. Some pro-inflammatory pathogens increased accompanied with the intestinal mucosal barrier
compromised, which lead to more bacterial LPS transferring into lymph nodes and blood to stimulate
the TOLL-like pathway of the host cells, and produce inflammatory cytokine (62). SLE patients
generally used massive immunosuppressive agents and glucocorticoids during the active period, which
could inhibit the immune system and might cause a large increase in opportunistic pathogens (63, 64).
Notwithstanding, it is questionable whether such changes in gut bacterial profile are a cause or
consequence of SLE. However, to posit further on this, is beyond the scope of this study, and we will focus on this in the future research.

In addition, several aberrant microbiome–associated gut metabolic pathways were revealed to be associated with SLE using PICRUSt analysis. We found that SLE patients were enriched in multiple metabolic pathways containing gene functions of Apoptosis, Purine metabolism, and the Apoptosis were positively associated with the genus *Streptococcus* that was highly enriched in SLE patients and especially in active patients. As reported, Apoptosis pathway played an important role in the pathogenesis of SLE (65). Besides, among these altered pathways, the alanine, aspartate and glutamate metabolism, Secondary bile acid biosynthesis, and Lipid metabolism were not only related to the disease activity, but also significantly associated with *Streptococcus*. The alanine, aspartate and glutamate metabolism, which was identified to be increased in remissive SLE patients in our study, had been previously reported to play a pivotal role in resting or activated T cells (66). Lipid metabolism participates in the regulation of many cellular processes such as cell growth, proliferation, differentiation, survival, apoptosis, inflammation, motility, etc (67). In active SLE patients, the dyslipidemia was more prevalent, suggesting that inflammation may be related to lipid metabolism (68). Thus, *Streptococcus* might play an important role in the pathogenesis of SLE through these pathways.

Due to the heterogeneous presentation of SLE patients and their unpredictable disease course, there is a great need for accurate assessment of disease activity. Several immunologic markers including anti-double stranded DNA (dsDNA) antibody and complement are common used in laboratory
monitoring of disease activity, however, these traditional biomarkers are better related to certain clinical manifestations of the disease, especially nephritis, rather than to the activity of the disease itself (69). Currently, disease activity in SLE can be assessed using composite disease activity indices, such as SLEDAI score and British Isles Lupus Assessment Group (BILAG) score (70). However, the composite disease activity indices depend on differential organ involvement and physical assessments (71, 72). Besides, they could be complex for use in routine clinical practice. Thus, there is a great urgent for the identification of new biomarkers that can quantify disease activity (73, 74).

Moreover, due to the existence of a remarkable difference in microbiota between SLE status, the random forest models were built in this study to examine whether microbiota composition could identify their disease status. Of note, a random forests model was identified for diagnosing SLE from healthy controls and RA patients with a AUC value of 0.792. To be mentioned, another random forest predictive model showed to be a suitable model for the prediction of disease activity of SLE with the AUC of 0.811, which was higher than the combination of Complement C3 and anti-dsDNA (AUC=0.773). Accordingly, our results suggested that the gut microbiota might be potential biomarkers for diagnosis of SLE and even monitoring SLE activity in a non-invasive method. However, the sample size enrolled in our study was relatively small, therefore, more samples are needed to evaluate the performance of the disordered genera in the future.

In summary, these disordered bacteria and related metabolic pathways might provide clues in studying of the SLE pathogenesis, and in searching for suitable biomarkers for the diagnosis SLE or
monitoring SLE activity in a non-invasive method. Specific microbial clades might be viable targets for the therapeutic manipulation by dietary interventions, prebiotics, probiotics and specifically tailored antibiotics. Determining the functions of the microbial clades that expand or contract in SLE will contribute to developing effective strategies to target them. However, the key role of microbiota in SLE pathogenesis and prospective mechanistic studies still need to be further investigated.

**Conclusions**

In this study, we reveal that intestinal dysbiosis and aberrant metabolism pathways are existed in SLE patients, especially in active SLE patients. Notably, there are 4 disordered genera and 2 species that are associated with the clinical disease activity in our patient cohort. Furthermore, there are two kinds of genera-panels can be the indicators for diagnosing or monitoring disease activity of SLE by random forest algorithm. However, we also recognize the limitations of our study. Since the results are deduced by a single-center study with a relatively small sample size, larger and prospective cohort studies will be required to verify and validate this predictive model.

**Declarations of interest**

None.
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Authors contribution statement

Yao Li, Haifang Wang, Xin Li, Ji-Liang Li, and Yurong Qiu were responsible for the overall design and interpretation of the study; Yurong Qiu and Ji-Liang Li conceived and designed the study; Yao Li, Haifang Wang, Xin Li, Qiong Zhang and Chen Fu performed the experiments; Yao Li, Haifang Wang, Xin Li and Xiaohe Zhang contributed to the experimental data collection and analysis; Hongwei Zhou, Yan He and Pan Li contributed to the bioinformatics analysis; Yao Li, Haifang Wang, Xin Li wrote the manuscript. All authors read and approved the final manuscript.

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**Table**
Table 1: Characteristics of the active SLE patients and remissive SLE patients.

|                          | A (n = 19)          | R (n = 21)          | P-value   |
|--------------------------|---------------------|---------------------|-----------|
| Age, years               | 34.05 (13.92)       | 40.57 (13.63)       | 0.143     |
| WBC, 10^9/L #            | 7.40 (5.32-8.53)    | 6.72 (5.72-8.22)    | 0.955     |
| RBC, 10^12/L #           | 4.17 (3.49-4.35)    | 4.33 (4.00-4.58)    | 0.0389*   |
| HGB, g/L #               | 106 (95-117.50)     | 126 (115-133.25)    | < 0.001 ***|
| PLT, 10^9/L #            | 205 (152.50-308.00) | 235.50 (202.25-272.50) | 0.558     |
| C3, g/L                  | 0.68 (0.32)         | 0.93 (0.23)         | 0.0013**  |
| CRP, mg/L #              | 1.68 (0.65-4.66)    | 1.56 (0.57-4.07)    | 0.765     |
| ESR, mm/h #              | 21.50 (11-53.50)    | 19.00 (5.00-60.00)  | 0.457     |
| Pyuria, n (%)            |                     |                     |           |
| - (negative)             | 6 (31.58)           | 17 (80.95)          | 0.002**   |
| + (positive)             | 13 (68.42)          | 4 (19.05)           |           |
| Albuminuria, n (%)       |                     |                     |           |
| - (negative)             | 3 (15.79)           | 15 (71.43)          | 0.000***  |
| + (positive)             | 16 (84.21)          | 6 (28.57)           |           |
| Hematuria, n (%)         |                     |                     |           |
| - (negative)             | 4 (21.05)           | 19 (90.48)          | 0.000***  |
| + (positive)             | 15 (78.95)          | 2 (9.52)            |           |
| 24-UTP, mg/24h #         | 0.68 (0.22-2.29)    | 0.17 (0.11-0.46)    | 0.048*    |
| Anti-dsDNA, UI/ml #      | 65.64 (16.24–156.66)| 20.07 (2.88–62.20)  | 0.0186*   |
| Lupus nephritis, n (%)   | 15 (78.95)          | 9 (42.86)           | 0.20      |
| SLEDAI #                 | 12 (9.5-14.0)       | 4 (1.5-6.0)         | < 0.001 ***|
| Medication use           |                     |                     |           |
| Hydroxychloroquine       | 12                  | 10                  | 0.119     |
| Glucocorticoid           | 17                  | 16                  | 0.527     |
| Cyclophosphamide         | 3                   | 3                   | 0.574     |
| Biological agent         | 5                   | 2                   | 0.894     |
A, the active SLE patients; R, the remissive SLE patients; WBC, white blood cell; RBC, red blood cell; HGB, haemoglobin; PLT, platelet; C3, Complement component 3; CRP, C-reactive protein; ESR, erythrocyte sedimentation rate; 24-UTP, 24-hour urine protein; Anti-dsDNA, anti-double stranded DNA; SLEDAI, Systemic Lupus Erythematosus Disease Activity Index. Data represent the mean (standard deviation), and the data # represents the median (interquartile range). The P values were calculated by Mann-Whitney U test or Chi-square test. *P < 0.05; **P < 0.01; ***P < 0.001.

**Figure legend**

**Figure 1.** The different microbial diversity between SLE patient group and healthy controls. A-B. Significantly different richness of α-diversity between the gut microbiota of SLE and HC. C. Principal coordinate analysis illustrating the grouping patterns of SLE and HC group based on the unweighted UniFrac distances. Each closed circle represented a sample. Distances between any pair of samples represented their dissimilarities. D. The average relative abundances of the predominant bacterial taxa at the phylum level in the SLE patients and HC group. E-F. The significantly different phyla in SLE patients compared to healthy controls. HC, healthy controls; SLE, Systemic lupus erythematosus patients.
Figure 2. The differentially abundant taxa between the feces of SLE patients and healthy controls. LEfSe analysis was performed to identify differentially abundant taxa, which are highlighted by the phylogenetic tree in cladogram format (A) and the LDA scores (B). Green color indicates an increase taxa in the feces of SLE compared with HC, while the red color indicates an increase taxa in the feces of HC compared with SLE (significant taxa \( p<0.005 \), Kruskal–Wallis test] with LDA score >2 were shown). SLE, Systemic lupus erythematosus patients; HC, healthy controls.

Figure 3. The different microbial diversity between the feces of active SLE patients and remissive SLE patients. A. Principal coordinate analysis illustrating the grouping patterns of the feces of A group and R group based on the unweighted UniFrac distances. Each closed circle represented a sample. Distances between any pair of samples represented their dissimilarities. B. The significantly different phyla in the feces of A group compared with R group. C-D. LEfSe analysis was performed to identify differentially abundant taxa, which are highlighted by by the phylogenetic tree in cladogram format (C) and the LDA scores (D). Significantly discriminative taxa among the active patients (red), remissive patients (blue) and healthy controls (green) were determined using Linear Discriminant Analysis Effect Size (LEfSe) (significant taxa \( p<0.005 \), Kruskal–Wallis test] with LDA score >2 were shown). A, the active SLE patients; R, the remissive SLE patients.
Figure 4. The significantly different predicted metabolic pathways between SLE patients and healthy controls. A. Significantly two predicted metabolic pathways were increased in SLE compared with healthy controls. B. Four predicted metabolic pathways were decreased in SLE patient group compared with healthy controls. It was analyzed by Kruskal–Wallis, and the False Discovery Rate (FDR) q-value was then calculated, and q-values <0.1 was considered significant. SLE, Systemic lupus erythematosus patients; HC, healthy controls.

Figure 5. The significantly different predicted metabolic pathways between active SLE patients and remissive SLE patients. A. Significantly five predicted metabolic pathways were increased in active SLE compared with remissive SLE patients. B. Four predicted metabolic pathways were enriched in remissive SLE patients compared with active SLE patients. A, the active SLE patient group; R, the remissive SLE patient group. It was analyzed by Kruskal–Wallis, and the False Discovery Rate (FDR) q-value was then calculated, and q-values <0.1 was considered significant.

Figure 6. The associations between the abundance of genus Streptococcus and disordered metabolic pathways. A. The predicted metabolic pathway Apoptosis and Pathways in cancer were correlated with SLE-enriched genus Streptococcus. B. The predicted metabolic pathway of Alanine aspartate and glutamate metabolism, Primary bile acid biosynthesis and Secondary bile acid biosynthesis were
negatively associated with active SLE-enriched genus *Streptococcus*. C. Five pathway categories, which were higher in active SLE patients compared to remissive SLE patient group, were positively associated with active SLE-enriched genus *Streptococcus*. They were analyzed by Spearman ranks tests, and the False Discovery Rate (FDR) was calculated for multiple testing. SLE, Systemic lupus erythematosus patients; FDR, the False Discovery Rate.

**Figure 7.** Associations among disease activity, disordered genera and predicted pathways in the gut microbiota of SLE patients. A-C. Three genera, *Streptococcus*, *Campylobacter*, and *Veillonella*, were positively correlated with lupus activity. D-E. Two species, *Streptococcus. anginosus*, and *Veillonella. dispar*, were positively correlated with lupus activity. F. The genus *Bifidobacterium* was negatively related to lupus activity. G-I. The aberrant microbiome-associated pathways, Alanine aspartate and glutamate metabolism, Secondary bile acid biosynthesis and Lipid metabolism, had a positive association with the activity of SLE patients. It was analyzed by Spearman ranks tests, and the False Discovery Rate (FDR) q-value was then calculated for multiple testing. SLEDAI, Systemic Lupus Erythematosus Disease Activity Index; C3, Complement component 3.

**Figure 8.** Receiver operating characteristic (ROC) curves demonstrating the performance of the genus and species in participants. A. Prediction model of the gut microbiota to distinguish the SLE patients
among the healthy controls and RA patients based on the genus/species-level relative abundances using random forests. B. Prediction model of the gut microbiota to differentiate the active SLE patients from remissive SLE patients. AUC, Area under the curves of ROC; CI, Confidence Interval; SLE, Systemic lupus erythematosus patients; RA, Rheumatoid Arthritis patients.

**Supplemental materials**

**Table S1:** The comparison of α-diversity among different groups.

SLE, Systemic lupus erythematosus patients; HC, healthy controls; RA, Rheumatoid Arthritis patients; A, the active SLE patients; R, the remissive SLE patients. The Wilcoxon rank sum test was used to determine significance in α-diversity. *P < 0.05; **P < 0.01; ***P < 0.001.

**Table S2:** The comparisons of β-diversity among different medicine treatments in SLE patients.

SLE, Systemic lupus erythematosus patients; Y= treated with specific drug; N= not treated. The ADONIS analysis was used to determine significance in β-diversity. *P < 0.05; **P < 0.01; ***P < 0.001.

**Table S3:** The comparisons of β-diversity among different groups.

SLE, Systemic lupus erythematosus patients; HC, healthy controls; RA, Rheumatoid Arthritis patients;
A, the active SLE patients; R, the remissive SLE patients. The ADONIS analysis was used to determine significance in β-diversity. *P < 0.05; **P < 0.01; ***P < 0.001.

Table S4: The comparisons of Firmicutes/Bacteroidetes ratio between different groups.

F/B, Firmicutes/Bacteroidetes ratio; SLE, Systemic lupus erythematosus patients; HC, healthy controls; RA, Rheumatoid Arthritis patients; A, the active SLE patients; R, the remissive SLE patients. The Wilcoxon rank sum test was used to determine significance in α-diversity. *P < 0.05; **P < 0.01; ***P < 0.001.

Table S5: Association of disordered genera and aberrant microbiome-associated pathways with activity of SLE.

SLEDAI, Systemic Lupus Erythematosus Disease Activity Index; C3, Complement component 3; CRP, C-reactive protein; ESR, Erythrocyte Sedimentation Rate; Anti-dsDNA, anti-double stranded DNA. They were analyzed by Spearman ranks tests, and False discovery rate (FDR) correction for multiple comparisons was employed. The FDR q-value was then calculated (r coefficient and FDR q-value were indicated for each parameter). *q < 0.05; **q< 0.01; ***q < 0.001.

Table S6: The importance of Genus and species in the random forests model to distinguish the SLE
patients from healthy controls and RA patients.

**Table S7:** The importance of Genus and species in the random forests model to distinguish the active SLE patients from remissive SLE patients.

**Figure S1.** Evaluation of sample preparation and sequencing quality. A. Chao1 dilution curve; B. Shannon dilution curve; C. OTU rank abundance.

**Figure S2.** The associations among α-diversity and Hydroxychloroquine, Glucocorticoid, Cyclophosphamide and Biological agent in SLE patients. SLE, Systemic lupus erythematosus patients; Y= treated with specific drug; N= not treated.

**Figure S3.** The Principal coordinate analysis (PCoA) of different medicine treatments. A-B. PCoA illustrating the grouping patterns of the Hydroxychloroquine-treated group (red) and Hydroxychloroquine-not treated group (blue) based on the weighted UniFrac distances (A) and unweighted UniFrac distances (B). C-D. PCoA illustrating the grouping patterns of the Glucocorticoid-treated group (red) and Glucocorticoid-not treated group (blue) based on the weighted UniFrac distances (C) and unweighted UniFrac distances (D). E-F. PCoA illustrating the grouping patterns of the Cyclophosphamide-treated group (red) and Cyclophosphamide-not treated group (blue)
based on the weighted UniFrac distances (E) and unweighted UniFrac distances (F). G-H. PCoA illustrating the grouping patterns of the Biological agent-treated group (red) and Biological agent-not treated group (blue) based on the weighted UniFrac distances (G) and unweighted UniFrac distances (H). SLE, Systemic lupus erythematosus patients; Y= treated with specific drug; N= not treated.

**Figure S4.** The Principal coordinate analysis (PCoA) of different groups.  A-B. PCoA illustrating the grouping patterns of the SLE patients and healthy controls based on the weighted UniFrac distances (A) and unweighted UniFrac distances (B). C-D. PCoA illustrating the grouping patterns of the SLE patients and RA patients based on the weighted UniFrac distances (C) and unweighted UniFrac distances (D). E-F. PCoA illustrating the grouping patterns of the active SLE patients and remissive SLE patients based on the weighted UniFrac distances (E) and unweighted UniFrac distances (F). SLE, Systemic lupus erythematosus patients; HC, the healthy controls; A, the active SLE patients; R, the remissive SLE patients.
Figure S5. The differentially abundant taxa between SLE patient group and RA patient group. LEfSe analysis was performed to identify differentially abundant taxa, which are highlighted on the phylogenetic tree in cladogram format (A) and for which the LDA scores are shown (B). Green color indicates an increase taxa in SLE compared with RA, while the red color indicates an increase taxa in RA compared with SLE. For cladogram format, from the interior to the exterior, each layer represents the phylum, class, order, family, and genus level (significant taxa [p<0.05, Kruskal–Wallis test] with LDA score >2.5 were shown). SLE, Systemic lupus erythematosus patients; RA, Rheumatoid Arthritis patients.

Figure S6. Receiver operating characteristic (ROC) curves demonstrating the performance of the anti-double stranded DNA and complement C3 to monitor the disease activity.
Table S1: The comparison of $\alpha$-diversity among different groups.

|                | SLE    | HC     | P-value |
|----------------|--------|--------|---------|
| Chao1          | 325.7  | 363.4  | 0.038*  |
| Observed species | 269    | 314.5  | 0.004** |
| Shannon index  | 5.043  | 5.202  | 0.089   |
| Simpson index  | 0.931  | 0.948  | 0.092   |

|                | SLE    | RA     | P-value |
|----------------|--------|--------|---------|
| Chao1          | 325.7  | 288.5  | 0.326   |
| Observed species | 269    | 216    | 0.324   |
| Shannon index  | 5.043  | 4.643  | 0.493   |
| Simpson index  | 0.931  | 0.920  | 0.629   |

|                | A      | R      | P-value |
|----------------|--------|--------|---------|
| Chao1          | 351.7  | 303.1  | 0.123   |
| Observed species | 300    | 251    | 0.092   |
| Shannon index  | 5.244  | 4.871  | 0.359   |
| Simpson index  | 0.931  | 0.924  | 0.663   |

SLE, Systemic lupus erythematosus patients; HC, healthy controls; RA, Rheumatoid Arthritis patients; A, the active SLE patients; R, the remissive SLE patients. The Wilcoxon rank sum test was used to determine significance in $\alpha$-diversity. *$P < 0.05$; **$P < 0.01$; ***$P < 0.001$. 
| Group                        | F. Modle  | R2   | P-value |
|------------------------------|-----------|------|---------|
| Hydroxychloroquine VS no Hydroxychloroquine weighted UniFrac | 1.55      | 0.039| 0.143   |
| Unweighted UniFrac           | 1.081     | 0.028| 0.317   |
| Glucocorticoid VS no Hydroxychloroquine weighted UniFrac | 0.657     | 0.017| 0.647   |
| Unweighted UniFrac           | 1.3       | 0.033| 0.129   |
| Cyclophosphamide VS no Cyclophosphamide weighted UniFrac | 1.315     | 0.033| 0.208   |
| Unweighted UniFrac           | 0.695     | 0.018| 0.908   |
| Biological agent VS no Biological agent weighted UniFrac | 0.155     | 0.004| 0.996   |
| Unweighted UniFrac           | 0.857     | 0.022| 0.672   |

SLE, Systemic lupus erythematosus patients. The ADONIS analysis was used to determine significance in β-diversity.
Table S3: The comparison of β-diversity among different groups.

| Group      | F. Modle    | R^2 | P-value     |
|------------|-------------|-----|-------------|
| SLE - HC   | Unweighted UniFrac | 3.406 | 0.054 | 0.000*** |
|            | Weighted UniFrac | 1.705 | 0.028 | 0.132    |
| SLE - RA   | Unweighted UniFrac | 1.271 | 0.021 | 0.143    |
|            | Weighted UniFrac | 0.993 | 0.017 | 0.399    |
| A - R      | Unweighted UniFrac | 1.549 | 0.039 | 0.047*   |
|            | Weighted UniFrac | 2.532 | 0.062 | 0.037*   |

SLE, Systemic lupus erythematosus patients; HC, healthy controls; RA, Rheumatoid Arthritis patients; A, the active SLE patients; R, the remissive SLE patients. The ADONIS analysis was used to determine significance in β-diversity. *P < 0.05; **P < 0.01; ***P < 0.001.
Table S4: The comparison of *Firmicutes/Bacteroidetes* ratio (F/B) between different groups.

| F/B | SLE   | HC    | P-value |
|-----|-------|-------|---------|
| 2.471 | 2.661 | 0.666 |
| A     | HC    | P-value |
| 3.907 | 2.661 | 0.549 |
| R     | HC    | P-value |
| 1.341 | 2.661 | 0.200 |

F/B, *Firmicutes/Bacteroidetes* ratio; SLE, Systemic lupus erythematosus patients; HC, healthy controls; RA, Rheumatoid Arthritis patients; A, the active SLE patients; R, the remissive SLE patients. The Wilcoxon rank sum test was used to determine significance.
### Table S5: Association of disordered genera and aberrant microbiome-associated pathway with activity of SLE.

| Genus and species levels | CRP r | FDR q-value | SLEDAI r | FDR q-value | ESR r | FDR q-value | C3 r | FDR q-value | dsDNA r | FDR q-value |
|-------------------------|-------|-------------|----------|-------------|-------|-------------|------|-------------|---------|-------------|
| *Lactobacillus*         | 0.089 | 0.676       | 0.054    | 0.810       | 0.397 | 0.056       | -0.039 | 0.893       | -0.127  | 0.725       |
| *Streptococcus*         | 0.098 | 0.676       | 0.492    | 0.008**     | 0.370 | 0.056       | -0.502 | 0.008**     | -0.118  | 0.725       |
| *Megasphaera*           | -0.150 | 0.676      | 0.166    | 0.409       | 0.052 | 0.796       | -0.123 | 0.674       | 0.079    | 0.838       |
| *Fusobacterium*         | -0.148 | 0.676      | 0.076    | 0.770       | -0.042 | 0.796       | 0.305  | 0.168       | 0.059    | 0.860       |
| *Veillonella*           | 0.088 | 0.676       | 0.223    | 0.250       | 0.078 | 0.760       | -0.475 | 0.008**     | -0.115  | 0.725       |
| *Oribacterium*          | 0.325 | 0.492       | 0.282    | 0.186       | 0.102 | 0.760       | -0.137 | 0.674       | 0.022    | 0.893       |
| *Campylobacter*         | 0.038 | 0.818       | 0.470    | 0.009**     | 0.423 | 0.056       | -0.201 | 0.427       | -0.114  | 0.725       |
| *Bifidobacterium*       | 0.121 | 0.676       | -0.511   | 0.008**     | -0.217 | 0.358       | 0.009  | 0.954       | -0.038  | 0.889       |
| *mucosae*               | 0.099 | 0.676       | 0.001    | 0.994       | 0.370 | 0.056       | 0.038  | 0.893       | -0.228  | 0.725       |
| *anginosus*             | 0.104 | 0.676       | 0.388    | 0.040*      | 0.244 | 0.311       | -0.253 | 0.275       | -0.130  | 0.725       |
| *dispar*                | 0.081 | 0.676       | 0.237    | 0.242       | 0.082 | 0.760       | -0.473 | 0.008**     | -0.122  | 0.725       |
| *gnavus*                | 0.106 | 0.676       | -0.257   | 0.218       | -0.130 | 0.726       | -0.087 | 0.792       | 0.117    | 0.725       |

| Pathway                  | CRP r | FDR q-value | SLEDAI r | FDR q-value | ESR r | FDR q-value | C3 r | FDR q-value | dsDNA r | FDR q-value |
|--------------------------|-------|-------------|----------|-------------|-------|-------------|------|-------------|---------|-------------|
| Synthesis and degradation of ketone bodies | -0.003 | 0.968 | 0.464 | 0.655 | 0.329 | 0.635 | -0.096 | 0.010* | -0.137 | 0.670       |
| Alanine aspartate and glutamate metabolism | -0.234 | 0.511 | -0.367 | 0.000*** | -0.253 | 0.000*** | 0.195 | 0.154 | 0.126 | 0.670       |
| Apoptosis                | 0.037 | 0.724 | 0.451 | 0.655 | 0.414 | 0.635 | -0.236 | 0.601 | -0.117 | 0.328       |
| Lipid metabolism         | -0.076 | 0.724 | 0.318 | 0.001** | 0.243 | 0.001** | -0.175 | 0.639 | 0.108 | 0.818       |
| Primary bile acid        | -0.172 | 0.850 | -0.371 | 0.655 | -0.196 | 0.635 | 0.137 | 0.927 | 0.144 | 0.670       |
|                      |                   |                   |                   |                   |                   |                   |                   |
|----------------------|-------------------|-------------------|-------------------|-------------------|-------------------|-------------------|-------------------|
| biosynthesis         |                   |                   |                   |                   |                   |                   |                   |
| Secondary bile acid  | -0.196            | 0.511             | -0.382            | 0.000***          | -0.186            | 0.000***          | 0.125             | 0.269             | 0.131             | 0.670             |
| biosynthesis         |                   |                   |                   |                   |                   |                   |                   |
| Secretion system     | 0.034             | 0.450             | 0.375             | 0.012*            | 0.295             | 0.014*            | 0.040             | 0.089             | -0.153            | 0.670             |
| Staphylococcus aureus| 0.179             | 0.724             | 0.420             | 0.232             | 0.360             | 0.225             | -0.462            | 0.002**           | -0.187            | 0.670             |

SLEDAI, Systemic Lupus Erythematosus Disease Activity Index; C3, Complement component 3; CRP, C-reactive protein; ESR, Erythrocyte Sedimentation Rate; Anti-dsDNA, anti-double stranded DNA. They were analyzed by Spearman ranks tests; False discovery rate (FDR) correction for multiple comparisons was employed; the False Discovery Rate q-value was then calculated (r coefficient and FDR q-value were indicated for each parameter). *q < 0.05; **q < 0.01; ***q < 0.001.
Table S6: The importance of genus and species in the random forests model to distinguish the SLE patients from healthy controls and RA patients.

| Genus and species levels | HC and RA (importance) | SLE (importance) | Model | Field |
|--------------------------|-------------------------|------------------|-------|-------|
| Bacteria.Firmicutes.Bacilli.Lactobacillales.Lactobacillaceae.Lactobacillus.mucosae | 100 | 100 | rf | group |
| NA | NA | NA | rf | group |
| Bacteria.Firmicutes.Bacilli.Lactobacillales.Lactobacillaceae.Lactobacillus | 88.8540 | 88.854 | rf | group |
| NA.1 | NA | NA | rf | group |
| Bacteria.Firmicutes.Clostridia.Clostridiales.Veillonellaceae.Acidaminococcus | 88.6828 | 88.682 | rf | group |
| NA.2 | NA | NA | rf | group |
| Bacteria.Firmicutes.Clostridia.Clostridiales.Ruminococcaceae.Faecalibacterium | 86.9773 | 86.977 | rf | group |
| NA.3 | NA | NA | rf | group |
| Bacteria.Firmicutes.Clostridia.Clostridiales.Ruminococcaceae.Fusobacterium | 84.8349 | 84.834 | rf | group |
| NA.4 | NA | NA | rf | group |
| Bacteria.Firmicutes.Clostridia.Clostridiales.Lachnospiraceae.[Ruminococcus].gnavus | 71.9087 | 71.908 | rf | group |
| NA.5 | NA | NA | rf | group |
| Bacteria.Actinobacteria.Actinobacteria.Bifidobacteriales.Bifidobacteriaceae.Scardovia | 68.4609 | 68.460 | rf | group |
| NA.6 | NA | NA | rf | group |
| Bacteria.Firmicutes.Clostridia.Clostridiales.Veillonellaceae.Megasphaera | 68.0529 | 68.052 | rf | group |
| NA.7 | NA | NA | rf | group |
| Bacteria.Firmicutes.Clostridia.Clostridiales.Veillonellaceae.Veillonella | 67.6085 | 67.608 | rf | group |
| NA.8 | NA | NA | rf | group |
| Bacteria.Firmicutes.Bacilli.Lactobacillales.Streptococcaceae.Streptococcus | 65.9776 | 65.977 | rf | group |
| NA.9 | NA | NA | rf | group |
| Bacteria.Firmicutes.Clostridia.Clostridiales.Ruminococcaceae.Oscillospira | 64.0312 | 64.031 | rf | group |
| NA.10 | NA | NA | rf | group |
| Class              | Order                          | Family               | Genus         | Accession Numbers | RF | Group |
|--------------------|--------------------------------|----------------------|---------------|-------------------|----|-------|
| Bacteria           | Actinobacteria                | Coriobacteriia      | Atopobium     | 63.995            | rf | group |
| NA.11              |                                |                      |               | 4483              |    |       |
|                    |                                |                      |               | 63.995            |    |       |
|                    |                                |                      |               | 64483             |    |       |
| Bacteria           | Actinobacteria                | Actinobacteria       | Bifidobacteriales | 59.4554           | rf | group |
| NA.12              |                                |                      | Bifidobacterium.adolescentis | 59.4556 |    |       |
|                    |                                |                      |               | 5463              |    |       |
|                    |                                |                      |               | 45463             |    |       |
| Bacteria           | Firmicutes                    | Bacilli              | Lactobacillales | 57.4657           | rf | group |
| NA.13              |                                |                      | Streptococcaceae | 57.4656           |    |       |
|                    |                                |                      | Streptococcus.anginosus | 57.4656 |    |       |
|                    |                                |                      |               | 1436              |    |       |
|                    |                                |                      |               | 71436             |    |       |
| Bacteria           | Proteobacteria                | Gammaproteobacteria  | Alteromonadales | 57.2296           | rf | group |
| NA.14              |                                |                      | Shewanellaceae | 57.2295           |    |       |
|                    |                                |                      | Shewanella     | 9146              |    |       |
|                    |                                |                      |               | 99146             |    |       |
| Bacteria           | Proteobacteria                | Gammaproteobacteria  | Vibrionales    | 57.1777           | rf | group |
| NA.15              |                                |                      | Vibrio         | 57.1771           |    |       |
|                    |                                |                      |               | 90843             |    |       |
| Bacteria           | Firmicutes                    | Bacilli              | Lactobacillales | 56.0413           | rf | group |
| NA.16              |                                |                      | Lactobacillaceae | 56.0414           |    |       |
|                    |                                |                      | Lactobacillus.ruminis | 56.0414 |    |       |
|                    |                                |                      |               | 187               |    |       |
|                    |                                |                      |               | 3187              |    |       |
| Bacteria           | Firmicutes                    | Clostridia           | Clostridiales  | 55.2231           | rf | group |
| NA.17              |                                |                      | Lachnospiraceae | 55.2229           |    |       |
|                    |                                |                      | Epulopiscium   | 8743              |    |       |
|                    |                                |                      |               | 78743             |    |       |
| Bacteria           | Proteobacteria                | Gammaproteobacteria  | Pasteurellales | 52.8883           | rf | group |
| NA.18              |                                |                      | Pasteurellaceae | 52.8882           |    |       |
|                    |                                |                      | Aggregatibacter | 9954              |    |       |
|                    |                                |                      |               | 39954             |    |       |
| Bacteria           | Firmicutes                    | Clostridia           | Clostridiales  | 52.5463           | rf | group |
| NA.19              |                                |                      | [Mogibacteriaceae] | 52.5464          |    |       |
|                    |                                |                      | Mogibacterium  | 2762              |    |       |
|                    |                                |                      |               | 22762             |    |       |
| Bacteria           | Firmicutes                    | Clostridia           | Clostridiales  | 52.2507           | rf | group |
| NA.20              |                                |                      | Lachnospiraceae | 52.2508           |    |       |
|                    |                                |                      | Coprococcus    | 4908              |    |       |
|                    |                                |                      |               | 74908             |    |       |
| Bacteria           | Firmicutes                    | Bacilli              | Lactobacillales | 50.3798           | rf | group |
| NA.21              |                                |                      | Leuconostocacae.Leuconostocae | 50.3798 |    |       |
|                    |                                |                      | Leuconostocae.Leuconostocae | 50.3793 |    |       |
|                    |                                |                      |               | 187               |    |       |
|                    |                                |                      |               | 8187              |    |       |
| Bacteria           | Actinobacteria                | Actinobacteria       | Bifidobacteriales | 49.8081           | rf | group |
| NA.22              |                                |                      | Bifidobacterium.longum | 49.8080          |    |       |
|                    |                                |                      |               | 9378              |    |       |
|                    |                                |                      |               | 19378             |    |       |
| Bacteria           | Proteobacteria                | Gammaproteobacteria  | Pasteurellales | 49.5807           | rf | group |
|                    |                                |                      | Pasteurellaece.Haemophilus.parainfluenzae | 49.5807 |    |       |
|                    |                                |                      |               | 9343              |    |       |
|                    |                                |                      |               | 79343             |    |       |
| NA.23 | Bacteria.Bacteroidetes.Bacteroidia.Bacteroidales.Bacteroidaceae. | 47.6555 | 47.655 | rf | group |
|       | Bacteroides.coprophilus                        | 8031    | 58031  |    |       |
| NA.24 | Bacteria.Firmicutes.Clostridia.Clostridiales.[Tissierellaceae].Anaerococcus | 46.8295 | 46.829 | rf | group |
|       |                                                | 7796    | 57796  |    |       |
| NA.25 | Bacteria.Firmicutes.Bacilli.Lactobacillales.Streptococcaceae.Lactococcus.garvieae | 45.5228 | 45.522 | rf | group |
|       |                                                | 3009    | 83009  |    |       |
| NA.26 | Bacteria.Firmicutes.Bacilli.Lactobacillales.Enterococcaceae.Enterococcus.cecorum | 44.3246 | 44.324 | rf | group |
|       |                                                | 7961    | 67961  |    |       |
| NA.27 | Bacteria.Firmicutes.Clostridia.Clostridiales.Veillonellaceae.Phascolarctobacterium | 43.7456 | 43.745 | rf | group |
|       |                                                | 4786    | 64786  |    |       |
| NA.28 | Bacteria.Firmicutes.Clostridia.Clostridiales.Veillonellaceae.Phascolarctobacterium | 42.4484 | 42.448 | rf | group |
|       |                                                | 5749    | 45749  |    |       |
| NA.29 | Bacteria.Firmicutes.Clostridia.Clostridiales.Lachnospiraceae.Mor yella            | 42.4231 | 42.423 | rf | group |
|       |                                                | 8053    | 18053  |    |       |
| NA.30 | Bacteria.Proteobacteria.Gammaproteobacteria.Vibrionales.Vibrio naceae.Photosobacterium | 42.2669 | 42.266 | rf | group |
|       |                                                | 8874    | 98874  |    |       |
| NA.31 | Bacteria.Firmicutes.Clostridia.Clostridiales.Veillonellaceae.Veillonella.parvula | 41.8961 | 41.896 | rf | group |
|       |                                                | 9119    | 19119  |    |       |
| NA.32 | Bacteria.Firmicutes.Clostridia.Clostridiales.Lachnospiraceae.Mor yella            | 40.8677 | 40.867 | rf | group |
|       |                                                | 1112    | 71112  |    |       |
| NA.33 | Bacteria.Firmicutes.Clostridia.Clostridiales.Lachnospiraceae.Lach nospira         | 40.7941 | 40.794 | rf | group |
|       |                                                | 2299    | 12299  |    |       |
| NA.34 | Bacteria.Firmicutes.Clostridia.Clostridiales.Lachnospiraceae.Mor yella            | 40.2141 | 40.214 | rf | group |
|       |                                                | 6926    | 16926  |    |       |
| NA.35 | Bacteria.Actinobacteria.Coriobacteria.Coriobacteriales.Coriobacterium            | 39.8194 | 39.819 | rf | group |
| Bacteria, Family, Genus | RF Group | RF Group | RF Group |
|-------------------------|----------|----------|----------|
| **eriaceae.Collinsella** | 3692     | 43692    |          |
| Bacteria.Firmicutes.Clostridia.Clostridiales.Veillonellacea.Megamonas | 39.8118 | 39.811 | rf group |
| | 3832     | 83832    |          |
| **NA.37** | NA       | NA       | rf group |
| Bacteria.Actinobacteria.Actinobacteria.Actinomycetales.Micrococaceae.Rothia.mucilaginosa | 39.2398 | 39.239 | rf group |
| | 6869     | 86869    |          |
| **NA.38** | NA       | NA       | rf group |
| Bacteria.Firmicutes.Bacilli.Lactobacillales.Carnobacteriaceae.Coprococcus.eutactus | 38.9446 | 38.944 | rf group |
| | 691      | 6691     |          |
| **NA.39** | NA       | NA       | rf group |
| Bacteria.Proteobacteria.Gammaproteobacteria.Pasteurellales.Pasteurellaceae.Aggregatibacter.pneumotropica | 38.7586 | 38.758 | rf group |
| | 9962     | 69962    |          |
| **NA.40** | NA       | NA       | rf group |
| Bacteria.Firmicutes.Erysipelotrichi.Erysipelotrichales.Erysipelotrichaceae.[Eubacterium] | 38.4991 | 38.499 | rf group |
| | 6132     | 16132    |          |
| **NA.41** | NA       | NA       | rf group |
| Bacteria.Proteobacteria.Gammaproteobacteria.Enterobacteriales.Enterobacteriaceae.Enterobacter | 38.4166 | 38.416 | rf group |
| | 7038     | 67038    |          |
| **NA.42** | NA       | NA       | rf group |
| Bacteria.Firmicutes.Clostridia.Clostridiales.Lachnospiraceae.Coprococcus.eutactus | 38.4081 | 38.408 | rf group |
| | 2124     | 12124    |          |
| **NA.43** | NA       | NA       | rf group |
| Bacteria.Firmicutes.Clostridia.Clostridiales.Veillonellaceae.Pectinatus | 38.0030 | 38.003 | rf group |
| | 853      | 0853     |          |
| **NA.44** | NA       | NA       | rf group |
| Bacteria.Firmicutes.Clostridia.Clostridiales.Ruminococcaceae.Ruminococcus | 37.5858 | 37.585 | rf group |
| | 4095     | 84095    |          |
| **NA.45** | NA       | NA       | rf group |
| Bacteria.Firmicutes.Clostridia.Clostridiales.Lachnospiraceae.[Ruminococcus] | 37.3746 | 37.374 | rf group |
| | 2973     | 62973    |          |
| **NA.46** | NA       | NA       | rf group |
| Bacteria.Firmicutes.Bacilli.Bacillales.Staphylococcaceae.Staphylococcus.sciuri | 37.3398 | 37.339 | rf group |
| | 9497     | 89497    |          |
| **NA.47** | NA       | NA       | rf group |
| Bacteria.Actinobacteria.Actinobacteria.Bifidobacteriales.Bifidobacteriaceae.Bifidobacterium.pseudolongum | 37.0794 | 37.079 | rf group |
| | 9274     | 49274    |          |
| **NA.48** | NA       | NA       | rf group |
| Kingdom                     | Phylum                        | Class                        | Order                              | Genus                      | Sequence Coverage | RF  | Group  |
|-----------------------------|-------------------------------|------------------------------|------------------------------------|----------------------------|-------------------|-----|--------|
| Bacteria                    | Proteobacteria                | Gammaproteobacteria          | Pseudomonadales.                  | Moraxellaceae.Acinetobacter.rhizosphaerae | 36.6336 6509 66509 | rf  | group  |
| NA.49                       |                               |                              |                                    |                            |                   |     |        |
| Bacteria                    | Bacteroidetes.                | Bacteroidia.                 | Bacteroidales.                     | Odoribacteraceae.          | 36.0353 703 36.035 3703 | rf  | group  |
| NA.50                       |                               |                              |                                    |                            |                   |     |        |
| Bacteria                    | Firmicutes.                   | Bacilli.                     | Lactobacillales.                   | Streptococcaceae. Streptococcus.sobrinus | 36.0023 7768 36.002 37768 | rf  | group  |
| NA.51                       |                               |                              |                                    |                            |                   |     |        |
| Bacteria                    | Actinobacteria.               | Actinobacteria.              | Actinomycetales.                   | Microccaceae.              | 35.8898 1106 35.889 81106 | rf  | group  |
| NA.52                       |                               |                              |                                    |                            |                   |     |        |
| Bacteria                    | Firmicutes.                   | Clostridia.                  | Clostridiales.                     | Lachnospiraceae.           | 35.5545 7379 35.554 57379 | rf  | group  |
| NA.53                       |                               |                              |                                    |                            |                   |     |        |
| Bacteria                    | Firmicutes.                   | Clostridia.                  | Clostridiales.                     | Clostridiaceae.            | 35.2804 2355 35.280 42355 | rf  | group  |
| NA.54                       |                               |                              |                                    |                            |                   |     |        |
| Bacteria                    | Firmicutes.                   | Bacilli.                     | Lactobacillales.                   | Lactobacillus. zeae        | 35.2526 0325 35.252 60325 | rf  | group  |
| NA.55                       |                               |                              |                                    |                            |                   |     |        |
| Bacteria                    | Proteobacteria.               | Gammaproteobacteria          | Pasteurellales.                    | Pasteurellaceae.           | 34.9841 941 34.984 1941 | rf  | group  |
| NA.56                       |                               |                              |                                    |                            |                   |     |        |
| Bacteria                    | Firmicutes.                   | Clostridia.                  | Clostridiales.                     | Veillonellaceae.           | 34.8368 1098 34.836 81098 | rf  | group  |
| NA.57                       |                               |                              |                                    |                            |                   |     |        |
| Bacteria                    | Proteobacteria.               | Deltaproteobacteria.         | Desulfovibrionales.                | Desulfovibrionaceae.       | 34.3992 1159 34.399 21159 | rf  | group  |
| NA.58                       |                               |                              |                                    |                            |                   |     |        |
| Bacteria                    | Firmicutes.                   | Erysipelotrichi.             | Erysipelotrichiales.               | Erysipelotrichaceae.       | 34.3124 5602 34.312 45602 | rf  | group  |
| NA.59                       |                               |                              |                                    |                            |                   |     |        |
| Bacteria                    | Bacteroidetes.                | Bacteroidia.                 | Bacteroidales. Porphyromonadacae. | Parabacteroides.           | 33.8313 4522 33.831 34522 | rf  | group  |
| NA.60                       |                               |                              |                                    |                            |                   |     |        |
| Bacteria                    | Proteobacteria.               | Gammaproteobacteria.         | Enterobacteriales.                 | Enterobacteriaceae.        | 33.7498 1453 33.749 81453 | rf  | group  |
| NA.61 | Bacteria.Proteobacteria.Gammaproteobacteria.Enterobacteriales.Enterobacteriaceae.Morganella.morganii |
|-------|---------------------------------------------------------------------------------------------------|
| NA.62 | Bacteria.Firmicutes.Bacilli.Lactobacillales.Aerococcaceae.Abiotrophia |
| NA.63 | Bacteria.Actinobacteria.Actinobacteria.Actinomycetales.Corynebacteriaceae.Corynebacterium.variabile |
| NA.64 | Bacteria.Bacteroidetes.Bacteroidia.Bacteroidales.Prevotellaceae.Prevotella.copri |
| NA.65 | Bacteria.Firmicutes.Clostridia.Clostridiales.[Tissierellaceae].Parvimonas |
| NA.66 | Bacteria.Firmicutes.Clostridia.Clostridiales.Lachnospiraceae.Roseburia.faecis |
| NA.67 | Bacteria.Firmicutes.Erysipelotrichi.Erysipelotrichales.Erysipelotrichaceae.[Eubacterium].cylindroides |
| NA.68 | Bacteria.Proteobacteria.Gammaproteobacteria.Enterobacteriales.Enterobacteriaceae.Serratia |
| NA.69 | Bacteria.Proteobacteria.Betaproteobacteria.Burkholderiales.Alcaligenaceae.Sutterella |
| NA.70 | Bacteria.Proteobacteria.Betaproteobacteria.Burkholderiales.Burkholderiaceae.Lautropia |
| NA.71 | Bacteria.Proteobacteria.Betaproteobacteria.Burkholderiales.Oxalobacteraceae.Oxalobacter.formigenes |
| NA.72 | Bacteria.Actinobacteria.Coriobacteriia.Coriobacterales.Coriobacteriaceae.Adlercreutzia |
| NA.73 | Bacteria.Bacteroidetes.Bacteroidia.Bacteroidales.Porphyromonad |
| Group                                      | Genus | Species | FASN | FLSN | RF | Group       |
|--------------------------------------------|-------|---------|------|------|----|-------------|
| aceae.Porphyromonas                         |       |         | 9366 | 79366|    |             |
| NA.74                                      | NA    | NA      | rf   |      |    | group       |
| Bacteria.Verrucomicrobia.Verrucomicrobiae.Verrucomicrobiaceae.Akkermansia.muciniphila | 29.4384 | 3701 | 29.438 | 43701 | rf | group      |
| NA.75                                      | NA    | NA      | rf   |      |    | group       |
| Bacteria.Firmicutes.Clostridia.Clostridiales.Clostridium. Clostridium                | 29.3992 | 4741 | 29.399 | 24741 | rf | group      |
| NA.76                                      | NA    | NA      | rf   |      |    | group       |
| Bacteria.Firmicutes.Clostridia.Clostridiales.[Tissierellaceae].Peptoniphilus         | 29.3058 | 7477 | 29.305 | 87477 | rf | group      |
| NA.77                                      | NA    | NA      | rf   |      |    | group       |
| Bacteria.Firmicutes.Clostridia.Clostridiales.Clostridium.SMB53                      | 29.2967 | 8578 | 29.296 | 78578 | rf | group      |
| NA.78                                      | NA    | NA      | rf   |      |    | group       |
| Bacteria.Firmicutes.Bacilli.Turicibacterales.Turicibacteraceae.Turicibacter       | 29.0686 | 5286 | 29.068 | 65286 | rf | group      |
| NA.79                                      | NA    | NA      | rf   |      |    | group       |
| Bacteria.Bacteroidetes.Bacteroidia.Bacteroidales.[Paraprevotellaceae].Paraprevotella | 29.0013 | 282  | 29.001 | 3282  | rf | group      |
| NA.80                                      | NA    | NA      | rf   |      |    | group       |
| Bacteria.Proteobacteria.Betaproteobacteria.Burkholderiales.Comamonadaceae.Comamonas | 28.6042 | 1255 | 28.604 | 21255 | rf | group      |
| NA.81                                      | NA    | NA      | rf   |      |    | group       |
| Bacteria.Firmicutes.Erysipelotrichi.Erysipelotrichales.Erysipelotrichaceae.Coprobacillus | 28.4411 | 1353 | 28.441 | 11353 | rf | group      |
| NA.82                                      | NA    | NA      | rf   |      |    | group       |
| Bacteria.Bacteroidetes.Bacteroidia.Bacteroidales.Bacteroidaceae. Bacteroides.acidifaciens | 28.3050 | 7096 | 28.305 | 07096 | rf | group      |
| NA.83                                      | NA    | NA      | rf   |      |    | group       |
| Bacteria.Firmicutes.Clostridia.Clostridiales.Peptostreptococcaceae.Peptostreptococcus | 27.8394 | 6022 | 27.839 | 46022 | rf | group      |
| NA.84                                      | NA    | NA      | rf   |      |    | group       |
| Bacteria.Firmicutes.Clostridia.Clostridiales.Lachnospiraceae.Roseburia              | 26.9286 | 069  | 26.928 | 6069  | rf | group      |
| NA.85                                      | NA    | NA      | rf   |      |    | group       |
| Bacteria.Firmicutes.Clostridia.Clostridiales.Lachnospiraceae.Oribacterium           | 26.8789 | 7591 | 26.878 | 97591 | rf | group      |
| NA.86                                      | NA    | NA      | rf   |      |    | group       |
| Taxonomy                                      | RF Value | Group  |
|----------------------------------------------|----------|--------|
| Bacteria. Firmicutes. Clostridia. Clostridiales. Veillonellaceae. Minas. hypermegale | 26.681   | group  |
| NA.87                                        | NA       | rf     |
| Bacteria. Firmicutes. Clostridia. Clostridiales. Lachnospiraceae. [Ruminococcus]. torques | 26.6496  | group  |
| NA.88                                        | NA       | rf     |
| Bacteria. Firmicutes. Clostridia. Clostridiaceae. Clostridium. perfringens          | 25.7679  | group  |
| NA.89                                        | NA       | rf     |
| Bacteria. Bacteroidetes. Bacteroidia. Bacteroidales. Bacteroidaceae. Bacteroides. caccoe | 26.0065  | group  |
| NA.90                                        | NA       | rf     |
| Bacteria. Firmicutes. Clostridia. Clostridiales. Eubacteriaceae. Pseudoramibacter. Eubacterium | 26.0064  | group  |
| NA.91                                        | NA       | rf     |
| Bacteria. Actinobacteria. Coriobacteriia. Coriobacteriales. Coriobacteriaceae. Eggerthella. lentia | 25.9142  | group  |
| NA.92                                        | NA       | rf     |
| Bacteria. Firmicutes. Erysipelotrichi. Erysipelotrichales. Erysipelotrichaceae. Bulleidia | 25.7908  | group  |
| NA.93                                        | NA       | rf     |
| Bacteria. Bacteroidetes. Bacteroidia. Bacteroidales. Bacteroidaceae. Bacteroides. uniformis | 25.7679  | group  |
| NA.94                                        | NA       | rf     |
| Bacteria. Firmicutes. Clostridia. Clostridiales. Lachnospiraceae. Anaerostipes    | 25.3047  | group  |
| NA.95                                        | NA       | rf     |
| Bacteria. Firmicutes. Clostridia. Clostridiales. Lachnospiraceae. Blautia        | 25.2388  | group  |
| NA.96                                        | NA       | rf     |
| Bacteria. Proteobacteria. Gammaproteobacteria. Pseudomonadales. Pseudomonadaceae. Pseudomonas | 25.1646  | group  |
| NA.97                                        | NA       | rf     |
| Bacteria. Bacteroidetes. Bacteroidia. Bacteroidales. Bacteroidaceae. Bacteroides | 24.8466  | group  |
| NA.98                                        | NA       | rf     |
| Bacteria. Actinobacteria. Actinobacteria. Actinomycetales. Corynebacteriaceae. Corynebacterium. durum | 24.8363  | group  |
| NA.99 | Bacteria.Actinobacteria.Coriobacteria.Coriobacteriales.Coriobiaceriaceae.Collinsella.aerofaciens | 24.6407        | 24.640 | rf | group |
| NA.100 | Bacteria.Proteobacteria.Deltaproteobacteria.Desulfovibrioales.Desulfovibrioaceae.Desulfovibrio | 24.3825        | 24.382 | rf | group |
| NA.101 | Bacteria.Firmicutes.Clostridia.Clostridiales.Christensenellaceae.Christensenella | 24.1326        | 24.132 | rf | group |
| NA.102 | Bacteria.Proteobacteria.Epsilonproteobacteria.Campylobacterales.Campylobacteraceae.Campylobacter | 24.1198        | 24.119 | rf | group |
| NA.103 | Bacteria.Firmicutes.Clostridia.Clostridiales.[Tissierellaceae].Wal_L_1855D | 23.8138        | 23.813 | rf | group |
| NA.104 | Bacteria.Firmicutes.Erysipelotrichi.Erysipelotrichales.Erysipelotrichaceae.Holdemania | 23.6500        | 23.650 | rf | group |
| NA.105 | Bacteria.Firmicutes.Erysipelotrichi.Erysipelotrichales.Erysipelotrichaceae.cc_115 | 23.5206        | 23.520 | rf | group |
| NA.106 | Bacteria.Firmicutes.Clostridia.Clostridiales.[Tissierellaceae].Finegoldia | 23.2187        | 23.218 | rf | group |
| NA.107 | Bacteria.Actinobacteria.Actinobacteria.Actinomycetales.Microccaceae.Rothia.aer 
| NA.108 | Bacteria.Firmicutes.Erysipelotrichi.Erysipelotrichales.Erysipelotrichaceae.[Eubacterium].dolichum | 22.4583        | 22.458 | rf | group |
| NA.109 | Bacteroidetes.Bacteroidia.Bacteroidales.Bacteroidaceae.Bacteroides.fragilis | 22.1206        | 22.120 | rf | group |
| NA.110 | Bacteroidetes.Bacteroidia.Bacteroidales.Bacteroidaceae.Bacteroides.ovatus | 22.0084        | 22.008 | rf | group |
| NA.111 | Bacteria.Actinobacteria.Actinobacteria.Actinomycetales.Actinom }
| Group | Species | Phylum | Class | Order | Family | Genus | Species | RF | Group |
|-------|---------|--------|-------|-------|--------|-------|---------|----|-------|
| NA.112 | Actinomyces | Actinobacteria | Actinobacteria | Actinomycetales | Propionibacteriaceae | Propionibacterium | acnes | 21.250 | 19523 | rf | group |
| NA.113 | Pyramidobacter.piscolens | Synergistetes | Synergistia | Synergistales | Dethiosulfovibrio | 1256 | 3295 | 03295 | rf | group |
| NA.114 | Dethiosulfovibrio | Actinobacteria | Actinobacteria | Actinomycetales | Propionibacteriaceae | Propionibacterium | acnes | 20.798 | 41373 | rf | group |
| NA.115 | Dethiosulfovibrio | Actinobacteria | Actinobacteria | Actinomycetales | Propionibacteriaceae | Propionibacterium | acnes | 20.798 | 41373 | rf | group |
| NA.116 | Propionibacterium | Actinobacteria | Actinobacteria | Actinomycetales | Propionibacteriaceae | Propionibacterium | acnes | 19.968 | 70781 | rf | group |
| NA.117 | Propionibacterium | Actinobacteria | Actinobacteria | Actinomycetales | Propionibacteriaceae | Propionibacterium | acnes | 19.968 | 70781 | rf | group |
| NA.118 | Propionibacterium | Actinobacteria | Actinobacteria | Actinomycetales | Propionibacteriaceae | Propionibacterium | acnes | 18.409 | 84322 | rf | group |
| NA.119 | Propionibacterium | Actinobacteria | Actinobacteria | Actinomycetales | Propionibacteriaceae | Propionibacterium | acnes | 18.409 | 84322 | rf | group |
| NA.120 | Propionibacterium | Actinobacteria | Actinobacteria | Actinomycetales | Propionibacteriaceae | Propionibacterium | acnes | 18.346 | 4621 | rf | group |
| NA.121 | Propionibacterium | Actinobacteria | Actinobacteria | Actinomycetales | Propionibacteriaceae | Propionibacterium | acnes | 18.346 | 4621 | rf | group |
| NA.122 | Propionibacterium | Actinobacteria | Actinobacteria | Actinomycetales | Propionibacteriaceae | Propionibacterium | acnes | 18.346 | 4621 | rf | group |
| NA.123 | Propionibacterium | Actinobacteria | Actinobacteria | Actinomycetales | Propionibacteriaceae | Propionibacterium | acnes | 18.346 | 4621 | rf | group |
| NA.124 | Propionibacterium | Actinobacteria | Actinobacteria | Actinomycetales | Propionibacteriaceae | Propionibacterium | acnes | 18.346 | 4621 | rf | group |
| Bacteria | Group | RF | Group | OTU | ID | OTU | ID | Group |
|----------|-------|----|-------|-----|----|-----|----|-------|
| Actinobacteria.Coriobacteria.Coriobacteriales.Coriobacteriaceae.Collinsella.stercoris | 17.362 | 17.362 | rf | group | 4412 | 84412 | 8065 | 18065 | rf | group | 17.362 | 84412 | 8065 | 18065 | rf | group |
| Actinobacteria.Bifidobacteriales.Bifidobacteriaceae.Bombiscardovia | 16.632 | 16.632 | rf | group | 3517 | 43517 | 16.104 | 92837 | rf | group | 15.107 | 18964 | 15.107 | 18964 | rf | group |
| Actinobacteria.Bifidobacteriaceae.Bombiscardovia | 12.959 | 12.959 | rf | group | 0718 | 60718 | 12.511 | 6103 | rf | group | 12.510 | 76103 | 12.510 | 76103 | rf | group |
| Actinobacteria.Coriobacteriaceae.Collinsella.stercoris | 16.704 | 16.704 | rf | group | 8065 | 18065 | 16.632 | 43517 | rf | group | 16.704 | 8065 | 16.704 | 8065 | rf | group |
| Actinobacteria.Bifidobacteriales.Bifidobacteriaceae.Bombiscardovia | 16.704 | 16.704 | rf | group | 8065 | 18065 | 16.632 | 43517 | rf | group | 16.704 | 8065 | 16.704 | 8065 | rf | group |
| Actinobacteria.Coriobacteriaceae.Collinsella.stercoris | 15.107 | 15.107 | rf | group | 81489 | 18964 | 15.107 | 18964 | rf | group | 15.107 | 81489 | 15.107 | 81489 | rf | group |
| Actinobacteria.Coriobacteriaceae.Collinsella.stercoris | 16.104 | 16.104 | rf | group | 92837 | 18964 | 16.704 | 8065 | rf | group | 16.104 | 92837 | 16.104 | 92837 | rf | group |
| Actinobacteria.Coriobacteriaceae.Collinsella.stercoris | 11.838 | 11.838 | rf | group | 76 | 976 | 11.837 | 976 | rf | group | 11.838 | 76 | 11.838 | 76 | rf | group |
| Actinobacteria.Coriobacteriaceae.Collinsella.stercoris | 11.762 | 11.762 | rf | group | 11028 | 11028 | 11.762 | 11028 | rf | group | 11.762 | 11028 | 11.762 | 11028 | rf | group |
| Actinobacteria.Coriobacteriaceae.Collinsella.stercoris | 11.442 | 11.442 | rf | group | 38598 | 38598 | 11.442 | 38598 | rf | group | 11.442 | 38598 | 11.442 | 38598 | rf | group |
| Actinobacteria.Coriobacteriaceae.Collinsella.stercoris | 14.751 | 14.751 | rf | group | 8964 | 18964 | 14.751 | 8964 | rf | group | 14.751 | 8964 | 14.751 | 8964 | rf | group |
| Actinobacteria.Coriobacteriaceae.Collinsella.stercoris | 11.442 | 11.442 | rf | group | 38598 | 38598 | 11.442 | 38598 | rf | group | 11.442 | 38598 | 11.442 | 38598 | rf | group |
| Actinobacteria.Coriobacteriaceae.Collinsella.stercoris | 13.332 | 13.332 | rf | group | 63207 | 18964 | 13.332 | 63207 | rf | group | 13.332 | 63207 | 13.332 | 63207 | rf | group |
| Actinobacteria.Coriobacteriaceae.Collinsella.stercoris | 14.751 | 14.751 | rf | group | 18964 | 18964 | 14.751 | 18964 | rf | group | 14.751 | 18964 | 14.751 | 18964 | rf | group |
| Actinobacteria.Coriobacteriaceae.Collinsella.stercoris | 11.762 | 11.762 | rf | group | 11028 | 11028 | 11.762 | 11028 | rf | group | 11.762 | 11028 | 11.762 | 11028 | rf | group |
| Actinobacteria.Coriobacteriaceae.Collinsella.stercoris | 14.751 | 14.751 | rf | group | 18964 | 18964 | 14.751 | 18964 | rf | group | 14.751 | 18964 | 14.751 | 18964 | rf | group |
| Actinobacteria.Coriobacteriaceae.Collinsella.stercoris | 11.442 | 11.442 | rf | group | 38598 | 38598 | 11.442 | 38598 | rf | group | 11.442 | 38598 | 11.442 | 38598 | rf | group |
| Actinobacteria.Coriobacteriaceae.Collinsella.stercoris | 11.838 | 11.838 | rf | group | 76 | 976 | 11.837 | 976 | rf | group | 11.838 | 76 | 11.838 | 76 | rf | group |
| Actinobacteria.Coriobacteriaceae.Collinsella.stercoris | 11.762 | 11.762 | rf | group | 11028 | 11028 | 11.762 | 11028 | rf | group | 11.762 | 11028 | 11.762 | 11028 | rf | group |
| Actinobacteria.Coriobacteriaceae.Collinsella.stercoris | 11.442 | 11.442 | rf | group | 38598 | 38598 | 11.442 | 38598 | rf | group | 11.442 | 38598 | 11.442 | 38598 | rf | group |
| Actinobacteria.Coriobacteriaceae.Collinsella.stercoris | 11.838 | 11.838 | rf | group | 76 | 976 | 11.837 | 976 | rf | group | 11.838 | 76 | 11.838 | 76 | rf | group |
| Actinobacteria.Coriobacteriaceae.Collinsella.stercoris | 11.762 | 11.762 | rf | group | 11028 | 11028 | 11.762 | 11028 | rf | group | 11.762 | 11028 | 11.762 | 11028 | rf | group |
| Actinobacteria.Coriobacteriaceae.Collinsella.stercoris | 11.442 | 11.442 | rf | group | 38598 | 38598 | 11.442 | 38598 | rf | group | 11.442 | 38598 | 11.442 | 38598 | rf | group |
| Actinobacteria.Coriobacteriaceae.Collinsella.stercoris | 11.838 | 11.838 | rf | group | 76 | 976 | 11.837 | 976 | rf | group | 11.838 | 76 | 11.838 | 76 | rf | group |
| Actinobacteria.Coriobacteriaceae.Collinsella.stercoris | 11.762 | 11.762 | rf | group | 11028 | 11028 | 11.762 | 11028 | rf | group | 11.762 | 11028 | 11.762 | 11028 | rf | group |
| Actinobacteria.Coriobacteriaceae.Collinsella.stercoris | 11.442 | 11.442 | rf | group | 38598 | 38598 | 11.442 | 38598 | rf | group | 11.442 | 38598 | 11.442 | 38598 | rf | group |
|    |    |    |    |    |
|----|----|----|----|----|
| NA | NA | rf | group |
| Bacteria. Proteobacteria. Alphaproteobacteria. Rhodobacterales. Rhodobacteraceae. Paracoccus | 10.9670 | 10.967 | rf | group |
|   | 9042 | 09042 |    |    |
| NA | NA | rf | group |
| Bacteria. Firmicutes. Bacilli. Bacillales. Bacillaceae. Bacillus | 10.9499 | 10.949 | rf | group |
|   | 5981 | 95981 |    |    |
| NA | NA | rf | group |
| Bacteria. Proteobacteria. Gammaproteobacteria. Pasteurellales. Pasteurellaceae. Actinobacillus. parahaemolyticus | 5.84837 | 5.8483 | rf | group |
|   | 1674 | 71674 |    |    |
| NA | NA | rf | group |
| Bacteria. Bacteroidetes. Bacteroidia. Bacteroidales. Porphyromonadaceae. Parabacteroides. gordonii | 0 | 0 | rf | group |

SLE, Systemic lupus erythematosus patients; HC, the healthy controls; RA, Rheumatoid Arthritis patients.
Table S7: The importance of genus and species in the random forests model to distinguish the active SLE patients from remissive SLE patients.

| The species and genus in the model | A    | R    | Model | Field |
|-----------------------------------|------|------|-------|-------|
| Bacteria.Proteobacteria.Epsilonproteobacteria.Campylobacteriales.Campylobacterales.Campylobacteraceae.Campylobacter | 100  | 100  | rf    | group |
| NA                               | NA   | NA   | rf    | group |
| Bacteria.Actinobacteria.Coriobacteria.Coriobacteriales.Coriobacteriaceae.Eggerthella.lenta | 73.529 | 73.529 | rf    | group |
| NA.1                             | NA   | NA   | rf    | group |
| Bacteria.Firmicutes.Bacilli.Lactobacillales.Streptococcaceae.Streptococcus | 67.683 | 67.683 | rf    | group |
| NA.2                             | NA   | NA   | rf    | group |
| Bacteria.Firmicutes.Clostridia.Clostridiales.Lachnospiraceae.[Ruminococcus].gnavus | 63.134 | 63.134 | rf    | group |
| NA.3                             | NA   | NA   | rf    | group |
| Bacteria.Actinobacteria.Actinobacteria.Bifidobacteriales.Bifidobacteriaceae.Bifidobacterium | 60.082 | 60.082 | rf    | group |
| NA.4                             | NA   | NA   | rf    | group |
| Bacteria.Firmicutes.Clostridia.Clostridiales.Lachnospiraceae.[Ruminococcus] | 57.013 | 57.013 | rf    | group |
| NA.5                             | NA   | NA   | rf    | group |
| Bacteria.Actinobacteria.Actinobacteria.Bifidobacteriales.Bifidobacteriaceae.Bifidobacterium.longum | 55.721 | 55.721 | rf    | group |
| NA.6                             | NA   | NA   | rf    | group |
| Bacteria.Firmicutes.Clostridia.Clostridiales.Lachnospiraceae.Oribacterium | 52.499 | 52.499 | rf    | group |
| NA.7                             | NA   | NA   | rf    | group |
| Bacteria.Firmicutes.Erysipelotrichi.Erysipelotrichales.Erysipelotrichaceae.[Eubacterium].dolichum | 51.546 | 51.546 | rf    | group |
| NA.8                             | NA   | NA   | rf    | group |
| Bacteria | Actinobacteria.Acinobacteria.Acinomycetales.Micrococcae.Rothia.aeria | 51.035 | 51.035 | rf | up | group |
|---|---|---|---|---|---|---|
| NA.9 | NA | NA | rf | gro | up |
| Bacteria | Actinobacteria.Acinobacteria.Acinomycetales.Micrococcae.Rothia.mucilaginosa | 49.300 | 49.300 | rf | gro | up |
| NA.10 | NA | NA | rf | gro | up |
| Bacteria | Bacteroidetes.Bacteroidia.Bacteroidales.Bacteroidaceae.Bacteroides.barnesiae | 48.418 | 48.418 | rf | gro | up |
| NA.11 | NA | NA | rf | gro | up |
| Bacteria | Bacteroidetes.Bacteroidia.Bacteroidales.Prevoetella.copri | 46.432 | 46.432 | rf | gro | up |
| NA.12 | NA | NA | rf | gro | up |
| Bacteria | Bacteroidetes.Bacteroidia.Bacteroidales.Bacteroidaceae.Bacteroides.ovatus | 43.175 | 43.175 | rf | gro | up |
| NA.13 | NA | NA | rf | gro | up |
| Bacteria | Bacteroidetes.Bacteroidia.Bacteroidales.Porphyromonadaceae.Parabacteroides.gordonii | 40.198 | 40.198 | rf | gro | up |
| NA.14 | NA | NA | rf | gro | up |
| Bacteria | Bacteroidetes.Bacteroidia.Bacteroidales.Porphyromonadaceae.Parabacteroides.distasonis | 38.728 | 38.728 | rf | gro | up |
| NA.15 | NA | NA | rf | gro | up |
| Bacteria | Proteobacteria.Betaproteobacteria.Neisseriales.Neisseriaceae.Neisseria.subflava | 38.453 | 38.453 | rf | gro | up |
| NA.16 | NA | NA | rf | gro | up |
| Bacteria | Firmicutes.Clostridia.Clostridiales.Lachnospiraceae.Coprococcus.eutactus | 38.060 | 38.060 | rf | gro | up |
| NA.17 | NA | NA | rf | gro | up |
| Bacteria | Firmicutes.Bacilli.Lactobacillales.Carnobacteriaceae.Granulicatella | 36.843 | 36.843 | rf | gro | up |
| NA.18  | NA | NA | rf | group |
|--------|----|----|----|-------|
| Bacteria.Firmicutes.Bacilli.Lactobacillales.Aerococcaceae.Facklamia | 35.791 | 35.791 | rf | group |
| NA.19  | NA | NA | rf | group |
| Bacteria.Firmicutes.Clostridia.Clostridiales.Veillonellaceae.Dialister | 35.669 | 35.669 | rf | group |
| NA.20  | NA | NA | rf | group |
| Bacteria.Proteobacteria.Betaproteobacteria.Neisseriales.Neisseriaceae.e.Neisseria | 35.197 | 35.197 | rf | group |
| NA.21  | NA | NA | rf | group |
| Bacteria.Proteobacteria.Gammaproteobacteria.Enterobacteriales.Ent erobacteriaceae.Citrobacter | 34.496 | 34.496 | rf | group |
| NA.22  | NA | NA | rf | group |
| Bacteria.Actinobacteria.Actinobacteria.Actinomycetales.Actinomyc etaceae.Actinomyces | 34.466 | 34.466 | rf | group |
| NA.23  | NA | NA | rf | group |
| Bacteria.Bacteroidetes.Bacteroidia.Bacteroidales.[Odoribacteraceae] .Butyricimonas | 34.323 | 34.323 | rf | group |
| NA.24  | NA | NA | rf | group |
| Bacteria.Bacteroidetes.Bacteroidia.Bacteroidales.Bacteroidaceae.Bacteroides.coprophilus | 34.323 | 34.323 | rf | group |
| NA.25  | NA | NA | rf | group |
| Bacteria.Firmicutes.Erysipelotrichi.Erysipelotrichales.Erysipelotrich aceae.Clostridium | 34.279 | 34.279 | rf | group |
| NA.26  | NA | NA | rf | group |
| Bacteria.Firmicutes.Clostridia.Clostridiales.Lachnospiraceae.Lachno bacterium | 33.480 | 33.480 | rf | group |
| NA.27  | NA | NA | rf | group |
| Bacteria.Proteobacteria.Betaproteobacteria.Burkholderiales.Comam | 33.117 | 33.117 | rf | group |
| Categories                                                   | ID1  | ID2  | Status |
|--------------------------------------------------------------|------|------|--------|
| Bacteria, Firmicutes, Clostridia, Mogibacteriaceae, Mogibacterium | 32.701 | 68673 | up     |
| Bacteria, Proteobacteria, Burkholderiales, Comamonadaceae, Acidovorax | 31.636 | 28904 | up     |
| Bacteria, Firmicutes, Bacilli, Lactobacillales, Lactobacillaceae, Lactobacillus ruminis | 31.381 | 70548 | up     |
| Bacteria, Firmicutes, Clostridia, Veillonellaceae, Veillonella parvula | 31.235 | 66388 | up     |
| Bacteria, Firmicutes, Bacilli, Lactobacillales, Streptococcaceae, Streptococcus luteciae | 30.932 | 3495 | up     |
| Bacteria, Bacteroidetes, Bacteroidia, Bacteroidales, Paraprevotellaceae, CF231 | 30.560 | 69782 | up     |
| Bacteria, Firmicutes, Clostridia, Lachnospiraceae, Lachnospira | 30.338 | 19745 | up     |
| Bacteria, Actinobacteria, Actinomycetales, Corynebacteriaceae, Corynebacterium | 29.941 | 54974 | up     |
| Bacteria, Proteobacteria, Betaproteobacteria, Burkholderiales, Alcaligenaceae, Sutterella | 29.842 | 49298 | up     |

**Legend**
- ID1: Identification Number 1
- ID2: Identification Number 2
- Status: "up" indicates an upregulated status.
| Genus/Species | 29.634 | 29.634 | rf | group  |
|---------------|--------|--------|----|--------|
| Bacteria.Firmicutes.Bacilli.Lactobacillales.Lactobacillaceae.Lactobacillus.zeae | 42159 | 42159 |    |        |
| NA.38         | NA     | NA     | rf | group  |
| Bacteria.Actinobacteria.Coriobacteria.Coriobacteriales.Coriobacteriaceae.Collinsella.stercoris | 29.022 | 29.022 | rf | group  |
| NA.39         | NA     | NA     | rf | group  |
| Bacteria.Firmicutes.Clostridia.Clostridiales.[Tissierellaceae].Parvimonas | 28.887 | 28.887 | rf | group  |
| NA.40         | NA     | NA     | rf | group  |
| Bacteria.Firmicutes.Clostridia.Clostridiales.Eubacteriaceae.Anaerostipes | 28.805 | 28.805 | rf | group  |
| NA.41         | NA     | NA     | rf | group  |
| Bacteria.Bacteroidetes.Bacteroidia.Bacteroidales.Bacteroidaceae.Bacteroides.plebeius | 28.792 | 28.792 | rf | group  |
| NA.42         | NA     | NA     | rf | group  |
| Bacteria.Firmicutes.Clostridia.Clostridiales.Lachnospiraceae.Anaerostipes | 28.788 | 28.788 | rf | group  |
| NA.43         | NA     | NA     | rf | group  |
| Bacteria.Bacteroidetes.Bacteroidia.Bacteroidales.Prevotellaceae.Prevotella.stercorea | 28.392 | 28.392 | rf | group  |
| NA.44         | NA     | NA     | rf | group  |
| Bacteria.Actinobacteria.Actinobacteria.Bifidobacteriales.Bifidobacteriaceae.Bifidobacterium.adolescentis | 28.126 | 28.126 | rf | group  |
| NA.45         | NA     | NA     | rf | group  |
| Bacteria.Firmicutes.Clostridia.Clostridiales.Ruminococcaceae.Faeclibacterium | 28.1115 | 28.1115 | rf | group  |
| NA.46         | NA     | NA     | rf | group  |
| Bacteria.Firmicutes.Clostridia.Clostridiales.Lachnospiraceae.Roseburia.faecis | 27.783 | 27.783 | rf | group  |
| NA.47         | NA     | NA     | rf | group  |
| Bacteria | Firmicutes.Clostridia.Clostridiales.Clostridiaceae.SMB53 | 27.180 | 27.180 | rf |  | group | up |
|----------|--------------------------------------------------------|--------|--------|----|----|--------|----|
|          | NA.48                                                  | NA     | NA     | rf | gro | group | up |
|          | Bacteria.Proteobacteria.Alphaproteobacteria.Rhizobiales.Bradyrhizobiaceae.Bradyrhizobium | 26.832 | 26.832 | rf | gro | group | up |
|          | Bacteria.Proteobacteria.Gammaproteobacteria.Enterobacteriales.Enterobacteriaceae.Enterobacter | 26.832 | 26.832 | rf | gro | group | up |
|          | NA.49                                                  | NA     | NA     | rf | gro | group | up |
|          | NA.50                                                  | NA     | NA     | rf | gro | group | up |
|          | Bacteria.Bacteroidetes.Bacteroidia.Bacteroidiales.Bacteroidaceae.Bacteroides | 26.652 | 26.652 | rf | gro | group | up |
|          | NA.51                                                  | NA     | NA     | rf | gro | group | up |
|          | Bacteria.Firmicutes.Clostridia.Clostridiales.Lachnospiraceae.Blautia | 26.473 | 26.473 | rf | gro | group | up |
|          | NA.52                                                  | NA     | NA     | rf | gro | group | up |
|          | Bacteria.Proteobacteria.Alphaproteobacteria.Sphingomonadales.Sphingomonadaceae.Sphingomonas | 25.600 | 25.600 | rf | gro | group | up |
|          | NA.53                                                  | NA     | NA     | rf | gro | group | up |
|          | Bacteria.Firmicutes.Clostridia.Clostridiales.Lachnospiraceae.Blautia.producta | 25.583 | 25.583 | rf | gro | group | up |
|          | NA.54                                                  | NA     | NA     | rf | gro | group | up |
|          | Bacteria.Proteobacteria.Gammaproteobacteria.Pasteurellales.Pasteurellaceae.Aggregatibacter.segnis | 25.385 | 25.385 | rf | gro | group | up |
|          | NA.55                                                  | NA     | NA     | rf | gro | group | up |
|          | Bacteria.Actinobacteria.Actinobacteria.Actinomycetales.Propionibacteriaceae.Propionibacterium.acnes | 25.278 | 25.278 | rf | gro | group | up |
|          | NA.56                                                  | NA     | NA     | rf | gro | group | up |
|          | Bacteria.Proteobacteria.Gammaproteobacteria.Enterobacteriales.Enterobacteriaceae.Erwinia | 25.217 | 25.217 | rf | gro | group | up |
| NA.57 | Bacteria.Firmicutes.Erysipelotrichi.Erysipelotrichales.Erysipelotrichaceae.Allobaculum | 25.014 | 62599 | rf | group |
| NA.58 | NA | NA | rf | group |
| NA.59 | Bacteria.Proteobacteria.Betaproteobacteria.Burkholderiales.Comamonadaceae.Delftia | 24.862 | 47222 | rf | group |
| NA.60 | Bacteria.Firmicutes.Bacilli.Lactobacillales.Enterococcaceae.Enterococcus | 24.453 | 21906 | rf | group |
| NA.61 | Bacteria.Firmicutes.Clostridia.Clostridiales.Christensenellaceae.Christensenella | 24.237 | 42137 | rf | group |
| NA.62 | Bacteria.Proteobacteria.Deltaproteobacteria.Desulfovibrionales.Desulfovibrionaceae.Desulfovibrio | 24.049 | 06378 | rf | group |
| NA.63 | Bacteria.Firmicutes.Clostridia.Clostridiales.Peptococcaceae.Peptococcus | 23.584 | 04599 | rf | group |
| NA.64 | Bacteria.Firmicutes.Bacilli.Lactobacillales.Lactobacillaceae.Lactobacillus | 23.461 | 52666 | rf | group |
| NA.65 | Bacteria.Firmicutes.Clostridia.Clostridiales.Lachnospiraceae.Dorea.formicigenerans | 23.102 | 78563 | rf | group |
| NA.66 | Bacteria.Proteobacteria.Gammaproteobacteria.Cardiobacteriales.Car | 22.832 | 22.832 | rf | group |
| diobacteriaceae.Cardiobacterium | 39661 | 39661 | up |
|--------------------------------|-------|-------|----|
| NA.67                          | NA    | NA    | rf |
| Bacteria.Bacteroidetes.Bacteroidia.Bacteroidales.Porphyromonadaceae.Porphyromonas | 22.748 | 22.748 | rf |
| NA.68                          | NA    | NA    | rf |
| Bacteria.Actinobacteria.Coriobacteriia.Coriobacteriales.Coriobacteriaceae.Slackia | 22.606 | 22.606 | rf |
| NA.69                          | NA    | NA    | rf |
| Bacteria.Firmicutes.Erysipelotrichi.Erysipelotrichales.Erysipelotrichaceae.Coprobacillus | 22.474 | 22.474 | rf |
| NA.70                          | NA    | NA    | rf |
| Bacteria.Firmicutes.Clostridia.Clostridiales.Ruminococcaceae.Oscillospira | 22.222 | 22.222 | rf |
| NA.71                          | NA    | NA    | rf |
| Bacteria.Bacteroidetes.Bacteroidia.Bacteroidales.[Odoribacteraceae].Odoribacter | 22.168 | 22.168 | rf |
| NA.72                          | NA    | NA    | rf |
| Bacteria.Actinobacteria.Actinobacteria.Actinomycetales.Corynebacteriaceae.Corynebacterium.durum | 21.983 | 21.983 | rf |
| NA.73                          | NA    | NA    | rf |
| Bacteria.Firmicutes.Clostridia.Clostridiales.Clostridiaceae.Clostridium.m.neonatale | 21.982 | 21.982 | rf |
| NA.74                          | NA    | NA    | rf |
| Bacteria.Bacteroidetes.Bacteroidia.Bacteroidales.Bacteroidaceae.Bacteroides.eggerthii | 21.791 | 21.791 | rf |
| NA.75                          | NA    | NA    | rf |
| Archaea.Euryarchaeota.Methanobacteria.Methanobacteriales.Methanobacteriaceae.Methanobrevibacter | 21.529 | 21.529 | rf |
| Bacteria.Firmicutes.Clostridia.Clostridiales.[Tissierellaceae].Finegolia | 21.529 | 21.529 | rf |
| NA.76 | NA | NA | rf | group |
|-------|----|----|----|-------|
| NA.77 | NA | NA | rf | group |
| Bacteria.Firmicutes.Clostridia.Clostridiales.Dehalobacteriaceae.Dehalobacterium | 21.086 | 32381 | 21.086 | 32381 | rf | group |
| NA.78 | NA | NA | rf | group |
| Bacteria.Actinobacteria.Coriobacteriia.Coriobacteriales.Coriobacteriaceae.Atopobium | 21.077 | 00539 | 21.077 | 00539 | rf | group |
| NA.79 | NA | NA | rf | group |
| Bacteria.Firmicutes.Clostridia.Clostridiales.[Tissierellaceae].Peptoniphilus | 21.051 | 92188 | 21.051 | 92188 | rf | group |
| NA.80 | NA | NA | rf | group |
| Bacteria.Proteobacteria.Gammaproteobacteria.Pseudomonadales.Moraxellaceae.Enhydrobacter | 20.806 | 61893 | 20.806 | 61893 | rf | group |
| NA.81 | NA | NA | rf | group |
| Bacteria.Bacteroidetes.Bacteroidia.Bacteroidales.Bacteroidaceae.Bacteroides.caccae | 20.670 | 882 | 20.670 | 882 | rf | group |
| NA.82 | NA | NA | rf | group |
| Bacteria.Firmicutes.Clostridia.Clostridiales.Veillonellaceae.Acidaminococcus | 20.655 | 32082 | 20.655 | 32082 | rf | group |
| NA.83 | NA | NA | rf | group |
| Bacteria.Firmicutes.Bacilli.Lactobacillales.Streptococcaceae.Lactococcus.garvieae | 19.643 | 70853 | 19.643 | 70853 | rf | group |
| NA.84 | NA | NA | rf | group |
| Bacteria.Firmicutes.Clostridia.Clostridiales.Lachnospiraceae.Dorea | 19.554 | 29461 | 19.554 | 29461 | rf | group |
| NA.85 | NA | NA | rf | group |
| Bacteria.Firmicutes.Clostridia.Clostridiales.Peptostreptococcaceae.Peptostreptococcus | 19.490 | 68014 | 19.490 | 68014 | rf | group |
| NA.86 | NA | NA | rf | group |
| Bacteria: Firmicutes, Bacilli, Lactobacillales, Aerococcaceae, Abiotrophia | 19.449 | 19.449 | rf | group up |
| NA.87 | NA | NA | rf | group up |
| Bacteria: Proteobacteria, Gammaproteobacteria, Pasteurellales, Pasteurellaceae, Actinobacillus, parahaemolyticus | 19.237 | 19.237 | rf | group up |
| NA.88 | NA | NA | rf | group up |
| Bacteria: Firmicutes, Bacilli, Bacillales, Staphylococcaceae, Staphylococcus | 19.185 | 19.185 | rf | group up |
| NA.89 | NA | NA | rf | group up |
| Bacteria: Firmicutes, Clostridia, Clostridiales, Veillonellaceae, Phascolarctobacterium | 18.896 | 18.896 | rf | group up |
| NA.90 | NA | NA | rf | group up |
| Bacteria: Firmicutes, Clostridia, Clostridiales, Ruminococcaceae, Anaerotruncus | 18.833 | 18.833 | rf | group up |
| NA.91 | NA | NA | rf | group up |
| Bacteria: Firmicutes, Clostridia, Clostridiales, Veillonellaceae, Veillonella | 18.253 | 18.253 | rf | group up |
| NA.92 | NA | NA | rf | group up |
| Bacteria: Proteobacteria, Gammaproteobacteria, Pseudomonadales, Moraxellaceae, Acinetobacter | 17.856 | 17.856 | rf | group up |
| NA.93 | NA | NA | rf | group up |
| Bacteria: Proteobacteria, Gammaproteobacteria, Vibrionales, Vibrionaceae, Vibrio | 17.662 | 17.662 | rf | group up |
| NA.94 | NA | NA | rf | group up |
| Bacteria: Firmicutes, Clostridia, Clostridiales, Ruminococcaceae, Ruminococcus | 17.600 | 17.600 | rf | group up |
| NA.95 | NA | NA | rf | group up |
| Bacteria: Proteobacteria, Gammaproteobacteria, Pasteurellales, Pasteurellaceae, Aggregatibacter | 17.395 | 17.395 | rf | group up |
| NA.96                      | NA | NA | rf | group |
|---------------------------|----|----|----|-------|
| Bacteria.Synergistetes.Synergistia.Synergistales.Dethiosulfovibrionaceae.Pyramidobacter.piscolens | 17.295 | 17.295 | rf | group |
|                           | 99524 | 99524 |    |       |
| NA.97                     | NA | NA | rf | group |
| Bacteria.Firmicutes.Clostridia.Clostridiales.Eubacteriaceae.Pseudoramibacter_Eubacterium | 17.229 | 17.229 | rf | group |
|                           | 92569 | 92569 |    |       |
| NA.98                     | NA | NA | rf | group |
| Bacteria.Proteobacteria.Gammaproteobacteria.Enterobacteriales.Enterobacteriaceae.Morganella.morganii | 16.806 | 16.806 | rf | group |
|                           | 09917 | 09917 |    |       |
| NA.99                     | NA | NA | rf | group |
| Bacteria.Firmicutes.Clostridia.Clostridiales.Clostridiaceae.Clostridium.perfringens | 16.801 | 16.801 | rf | group |
|                           | 47762 | 47762 |    |       |
| NA.100                    | NA | NA | rf | group |
| Bacteria.Bacteroidetes.Bacteroidia.Bacteroidales.Rikenellaceae.AF12 | 16.471 | 16.471 | rf | group |
|                           | 67993 | 67993 |    |       |
| NA.101                    | NA | NA | rf | group |
| Bacteria.Actinobacteria.Coriobacteriia.Coriobacterales.Coriobacteriaceae.Adlercreutzia | 16.453 | 16.453 | rf | group |
|                           | 71933 | 71933 |    |       |
| NA.102                    | NA | NA | rf | group |
| Bacteria.Firmicutes.Erysipelotrichi.Erysipelotrichales.Erysipelotrichiaceae.Holdemania | 16.356 | 16.356 | rf | group |
|                           | 76276 | 76276 |    |       |
| NA.103                    | NA | NA | rf | group |
| Bacteria.Actinobacteria.Actinobacteria.Actinomycetales.Micrococaceae.Rothia.dentocariosa | 16.226 | 16.226 | rf | group |
|                           | 55915 | 55915 |    |       |
| NA.104                    | NA | NA | rf | group |
| Bacteria.Proteobacteria.Gammaproteobacteria.Pasteurellales.Pasteurellaceae.Haemophilus | 16.207 | 16.207 | rf | group |
|                           | 60864 | 60864 |    |       |
| NA.105                    | NA | NA | rf | group |
| Bacteria.Actinobacteria.Coriobacteriia.Coriobacterales.Coriobacteriaceae | 15.439 | 15.439 | rf | group |
| Genome | x | y | z | w |
|--------|---|---|---|---|
| aceae.Collinsella.aerofaciens | 21086 | 21086 | up |
| NA.106 | NA | NA | rf | group |
| Bacteria.Tenericutes.Mollicutes.Mycoplasmales.Mycoplasmae.Mycoplasma | 14.998 | 14.998 | rf | group |
| NA.107 | NA | NA | rf | group |
| Bacteria.Proteobacteria.Gammaproteobacteria.Alteromonadales.Shewanellaceae.Shewanella.algae | 14.912 | 14.912 | rf | group |
| NA.108 | NA | NA | rf | group |
| Bacteria.Bacteroidetes.Bacteroidia.Bacteroidales.[Paraprevotellaceae].[Prevotella].tannerae | 14.858 | 14.858 | rf | group |
| NA.109 | NA | NA | rf | group |
| Bacteria.Proteobacteria.Alphaproteobacteria.Rhizobiales.Methylobacteriaceae.Methylobacterium.organophilum | 14.541 | 14.541 | rf | group |
| NA.110 | NA | NA | rf | group |
| Bacteria.Bacteroidetes.Bacteroidia.Bacteroidales.[Paraprevotellaceae].Paraprevotella | 14.417 | 14.417 | rf | group |
| NA.111 | NA | NA | rf | group |
| Bacteria.Proteobacteria.Gammaproteobacteria.Enterobacteriales.Entercardiaceae.Serratia | 13.760 | 13.760 | rf | group |
| NA.112 | NA | NA | rf | group |
| Bacteria.Actinobacteria.Actinobacteria.Bifidobacteriales.Bifidobacteriaceae.Scardovia | 13.372 | 13.372 | rf | group |
| NA.113 | NA | NA | rf | group |
| Bacteria.Firmicutes.Clostridia.Clostridiales.Clostridiaceae.Clostridium | 13.196 | 13.196 | rf | group |
| NA.114 | NA | NA | rf | group |
| Bacteria.Bacteroidetes.Flavobacteriia.Flavobacteriales.Flavobacteriaceae.Flavobacterium | 13.025 | 13.025 | rf | group |
| NA.115 | NA | NA | rf | group |
| Bacteria | Actinobacteria | Actinobacteria | Actinomycetales | Nocardiaceae | Rhodococcus | 12.411 | 12.411 | rf | group | 73113 | 73113 | 73113 |
|----------|----------------|----------------|-----------------|-------------|-------------|--------|--------|----|-------|--------|--------|--------|
| NA.116 | | | | | | NA | NA | rf | group | | | |
| Bacteria | Proteobacteria | Alphaproteobacteria | Sphingomonadales | Sphingobium | | 12.411 | 12.411 | rf | group | 63161 | 63161 | 63161 |
| NA.117 | | | | | | NA | NA | rf | group | | | |
| Bacteria | Bacteroidetes | Bacteroidia | Bacteroidales | Bacteroidaceae | Bacteroides | fragilis | 12.134 | 12.134 | rf | group | 45195 | 45195 | 45195 |
| NA.118 | | | | | | NA | NA | rf | group | | | |
| Bacteria | Proteobacteria | Gammaproteobacteria | Vibrionales | Vibrionaceae | Vibrio | rumoiensis | 12.076 | 12.076 | rf | group | 04882 | 04882 | 04882 |
| NA.119 | | | | | | NA | NA | rf | group | | | |
| Bacteria | Firmicutes | Bacilli | Lactobacillales | Streptococcaceae | Streptococcus | sobrinus | 12.012 | 12.012 | rf | group | 89651 | 89651 | 89651 |
| NA.120 | | | | | | NA | NA | rf | group | | | |
| Bacteria | Firmicutes | Clostridia | Clostridiales | Veillonellaceae | Megamonas | | 11.868 | 11.868 | rf | group | 28997 | 28997 | 28997 |
| NA.121 | | | | | | NA | NA | rf | group | | | |
| Bacteria | Bacteroidetes | Bacteroidia | Bacteroidales | Bacteroidaceae | Bacteroides | uniformis | 11.032 | 11.032 | rf | group | 3025 | 3025 | 3025 |
| NA.122 | | | | | | NA | NA | rf | group | | | |
| Bacteria | Firmicutes | Clostridia | Clostridiales | Ruminococcaceae | Ruminococcus | bromii | 10.923 | 10.923 | rf | group | 45167 | 45167 | 45167 |
| NA.123 | | | | | | NA | NA | rf | group | | | |
| Bacteria | Firmicutes | Bacilli | Bacillales | Staphylococcaceae | Staphylococcus | sciuri | 10.516 | 10.516 | rf | group | 56122 | 56122 | 56122 |
| NA.124 | | | | | | NA | NA | rf | group | | | |
| Bacteria | Proteobacteria | Gammaproteobacteria | Pseudomonadales | Pseudomonadaceae | Pseudomonas | | 10.258 | 10.258 | rf | group | 18194 | 18194 | 18194 |
| NA.125 | | | | | | NA | NA | rf | group | | | |
| Kingdom | Phylum | Class | Order | Family | Genus | OTU | Genus | Taxonomy | OTU | Genus | Taxonomy | OTU | Genus | Taxonomy | OTU | Genus | Taxonomy | OTU | Genus | Taxonomy | OTU | Genus | Taxonomy | OTU | Genus | Taxonomy | OTU | Genus | Taxonomy | OTU | Genus | Taxonomy | OTU | Genus | Taxonomy | OTU | Genus | Taxonomy | OTU | Genus | Taxonomy | OTU | Genus | Taxonomy | OTU | Genus | Taxonomy | OTU | Genus | Taxonomy | OTU | Genus | Taxonomy | OTU | Genus | Taxonomy | OTU | Genus | Taxonomy | OTU | Genus | Taxonomy | OTU | Genus | Taxonomy | OTU | Genus | Taxonomy | OTU | Genus | Taxonomy | OTU | Genus | Taxonomy | OTU | Genus | Taxonomy | OTU | Genus | Taxonomy | OTU | Genus | Taxonomy | OTU | Genus | Taxonomy | OTU | Genus | Taxonomy | OTU | Genus | Taxonomy | OTU | Genus | Taxonomy | OTU | Genus | Taxonomy | OTU | Genus | Taxonomy | OTU | Genus | Taxonomy | OTU | Genus | Taxonomy | OTU | Genus | Taxonomy | OTU | Genus | Taxonomy | OTU | Genus | Taxonomy | OTU | Genus | Taxonomy | OTU | Genus | Taxonomy | OTU | Genus | Taxonomy | OTU | Genus | Taxonomy | OTU | Genus | Taxonomy | OTU | Genus | Taxonomy | OTU | Genus | Taxonomy | OTU | Genus | Taxonomy | OTU | Genus | Taxonomy | OTU | Genus | Taxonomy | OTU | Genus | Taxonomy | OTU | Genus | Taxonomy | OTU | Genus | Taxonomy | OTU | Genus | Taxonomy | OTU | Genus | Taxonomy | OTU | Genus | Taxonomy | OTU | Genus | Taxonomy | OTU | Genus | Taxonomy | OTU | Genus | Taxonomy | OTU | Genus | Taxonomy | OTU | Genus | Taxonomy | OTU | Genus | Taxonomy | OTU | Genus | Taxonomy | OTU | Genus | Taxonomy | OTU | Genus | Taxonomy | OTU | Genus | Taxonomy | OTU | Genus | Taxonomy | OTU | Genus | Taxonomy | OTU | Genus | Taxonomy | OTU | Genus | Taxonomy | OTU | Genus | Taxonomy | OTU | Genus | Taxonomy | OTU | Genus | Taxonomy | OTU | Genus | Taxonomy | OTU | Genus | Taxonomy | OTU | Genus | Taxonomy | OTU | Genus | Taxonomy | OTU | Genus | Taxonomy | OTU | Genus | Taxonomy | OTU | Genus | Taxonomy | OTU | Genus | Taxonomy | OTU | Genus | Taxonomy | OTU | Genus | Taxonomy | OTU | Genus | Taxonomy | OTU | Genus | Taxonomy | OTU | Genus | Taxonomy | OTU | Genus | Taxology of the Bacteria domain.

- **Bacteria.Bacteroidetes/[Saprospirae].[Saprospirales].Chitinophagaceae.Sediminibacterium**
  - OTU: 10.137
  - Genus: NA
  - Numerical data:
    - 2439

- **Bacteria.Firmicutes.Erysipelotrichi.Erysipelotrichales.Erysipelotrichaceae.Bulleidia**
  - OTU: 9.6591
  - Genus: NA
  - Numerical data:
    - 89488

- **Bacteria.Actinobacteria.Actinobacteria.Actinomycetales.Pseudonocardia.Bulleidia**
  - OTU: 9.4716
  - Genus: NA
  - Numerical data:
    - 62675

- **Bacteria.Actinobacteria.Actinobacteria.Bifidobacteriales.Bifidobacteriaceae.Bombiscardovia**
  - OTU: 9.2896
  - Genus: NA
  - Numerical data:
    - 09429

- **Bacteria.Proteobacteria.Gammaproteobacteria.Xanthomonadales.Xanthomonadaceae.Stenotrophomonas**
  - OTU: 9.2626
  - Genus: NA
  - Numerical data:
    - 01412

- **Bacteria.Verrucomicrobia.Verrucomicrobiae.Verrucomicrobiales.Verrucomicrobiaceae.Akkermansia.muciniphila**
  - OTU: 9.1251
  - Genus: NA
  - Numerical data:
    - 89649

- **Bacteria.Proteobacteria.Betaproteobacteria.Burkholderiales.Burkholderiaceae.Burkholderia**
  - OTU: 9.0364
  - Genus: NA
  - Numerical data:
    - 12749

- **Bacteria.Proteobacteria.Gammaproteobacteria.Vibrionales.Vibrionaceae.Photobacterium**
  - OTU: 8.9156
  - Genus: NA
  - Numerical data:
    - 01809

- **Bacteria.Firmicutes.Clostridia.Clostridiales.Lachnospiraceae.Moryella**
  - OTU: 8.7452
  - Genus: NA
  - Numerical data:
    - 2191

- **Bacteria.Proteobacteria.Betaproteobacteria.Burkholderiales.Oxalobacteraceae.Oxalobacter.formigenes**
  - OTU: 8.6865
  - Genus: NA
  - Numerical data:
    - 73361

- **Bacteria.Firmicutes.Clostridia.Clostridiales.Lachnospiraceae.Moryella**
  - OTU: 8.7452
  - Genus: NA
  - Numerical data:
    - 2191
| ID  | Group               | Bacteria                                                                 | Phylum                        | Class                          | Order                     | Family                  | Genus           | Species         | Log_{10} DNA reads | RF  | Group |
|-----|---------------------|--------------------------------------------------------------------------|-------------------------------|--------------------------------|----------------------------|-------------------------|-------------------|------------------|-------------------|------|--------|
| NA.135 |                     | Bacteria.Firmicutes.Clostridia.Clostridiales.Veillonellaceae.Megasphaera | NA                            | NA                             | rf                         | Group                   |                   |                  | 8.6761           | 9462 | Group |
| NA.136 |                     | Bacteria.Proteobacteria.Deltaproteobacteria.Desulfovibrionales.Desulfovibrionaceae.Bilophila | NA                            | NA                             | rf                         | Group                   |                   |                  | 8.5509           | 60344| Group |
| NA.137 |                     | Bacteria.Firmicutes.Bacilli.Lactobacillales.Lactobacillaceae.Lactobacillus.mucosae | NA                            | NA                             | rf                         | Group                   |                   |                  | 8.4448           | 65919| Group |
| NA.138 |                     | Bacteria.Firmicutes.Clostridia.Clostridiales.Veillonellaceae.Selenomas    | NA                            | NA                             | rf                         | Group                   |                   |                  | 8.3328           | 26301| Group |
| NA.139 |                     | Bacteria.Actinobacteria.Actinobacteria.Actinomycetales.Micrococcaceae.Micrococcus | NA                            | NA                             | rf                         | Group                   |                   |                  | 7.7464           | 41536| Group |
| NA.140 |                     | Bacteria.Firmicutes.Clostridia.Clostridiales.Lachnospiraceae.Coprococcus | NA                            | NA                             | rf                         | Group                   |                   |                  | 7.6351           | 27255| Group |
| NA.141 |                     | Bacteria.Proteobacteria.Gammaproteobacteria.Enterobacteriales.Entrobacteriaceae.Klebsiella | NA                            | NA                             | rf                         | Group                   |                   |                  | 7.3767           | 70496| Group |

A, the active SLE patients; R, the remissive SLE patients.