Analysis of Conjunctival Sac Microbiome in Dry Eye Patients With and Without Sjögren’s Syndrome

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Purpose: To analyze the conjunctival sac microbial communities in patients with Sjögren’s syndrome-associated dry eyes (SSDE) and non-Sjögren’s syndrome-associated dry eyes (NSSDE), compared with normal controls (NC).

Methods: Conjunctival sac swab samples from 23 eyes of SSDE, 36 eyes of NSSDE, and 39 eyes of NC were collected. The V3–V4 region of the 16S ribosomal RNA (rRNA) gene high-throughput sequencing was performed on an Illumina MiSeq platform and analyzed using Quantitative Insights Into Microbial Ecology (QIIME). Alpha diversity was employed to analyze microbiome diversity through Chao1 and Shannon indexes. Beta diversity was demonstrated by the principal coordinates analysis (PCoA) and Partial Least Squares Discrimination Analysis (PLS-DA). The relative abundance was bioinformatically analyzed at the phylum and genus levels.

Results: The alpha diversity was lower in patients with dry eye disease (Shannon index: NC vs. SSDE: \( P = 0.020 \), NC vs. NSSDE: \( P = 0.029 \)). The beta diversity showed divergent microbiome composition in different groups (NC vs. SSDE: \( P = 0.001 \), NC vs. NSSDE: \( P = 0.001 \), NSSDE vs. SSDE: \( P = 0.005 \)). The top 5 abundant phyla were Firmicutes, Proteobacteria, Actinobacteriota, Bacteroidota, and Cyanobacteria in all three groups. The top five abundant genera included Acinetobacter, Staphylococcus, Bacillus, Corynebacterium, and Clostridium_sensu_stricto_1. The relative microbiome abundance was different between groups. The Firmicutes/Bacteroidetes (F/B) ratio was 6.42, 7.31, and 9.71 in the NC, NSSDE, and SSDE groups, respectively (NC vs. SSDE: \( P = 0.038 \), NC vs. NSSDE: \( P = 0.991 \), SSDE vs. NSSDE: \( P = 0.048 \)).

Conclusion: The diversity of conjunctival sac microbiome in patients with NSSDE and SSDE was diminished compared with NC. The main microbiome at the phylum and genus level were similar between groups, but the relative abundance had variations. The Firmicutes/Bacteroidetes ratio was higher in the SSDE group.

Keywords: ocular surface, microbial diversity, microbial composition, dry eyes, Sjögren’s syndrome
INTRODUCTION

Dry eye disease (DED) is a multifactorial inflammatory ocular disease featured by tear film instability and hyperosmolarity, inflammation, and neurosensory abnormalities (1, 2). Symptoms include burning, foreign body sensation, photophobia, and blurred vision, which would adversely affect daily lives (3). Autoimmune disease-associated dry eye is one of the most important types of DED (4). Sjögren’s syndrome (SS) is a chronic autoimmune disease featured by dry mucosal surfaces and other systemic muscular pain, and dry eye is one of the most discomforting symptoms that patients with SS complain about (5, 6). The lacrimal gland becomes infiltrated with activated CD4+ T cells and B cells, and ocular surface diseases develop from reduced lubrication, as well as from cytokines produced by activated epithelial cells and infiltrating inflammatory cells (7, 8).

Recently, mounting evidence shows that microbial imbalance changes the immunity toward an increased inflammatory response, which is closely associated with the occurrence of multiple chronic inflammatory diseases (9, 10). Hence, it is hypothesized that similar mechanisms are also active on the ocular surface and are involved in the pathophysiology of DED including SS. The production of lipases and toxins by different proportions of colonizing bacteria may destabilize the lipid layer of the tear film, interact with the conjunctival mucus (11, 12) and cause ocular surface cellular damage. Thus, dysbiosis might be associated with tear film instability, inflammation, and ocular irritation.

The purpose of the study was to explore the potential changes in the conjunctival sac microbiome in dry eye patients with and without SS, compared with healthy subjects. This may help to better understand its pathophysiology and provide more evidence to develop new treatment strategies.

METHODS

Sample Collection

Patients were recruited from the regular outpatient clinic, Department of Ophthalmology, Peking Union Medical College Hospital, between September 1 and October 30, 2021. Patients with SS-associated dry eyes (SSDE), non-SS-associated dry eyes (NSSDE), and normal controls (NC) with healthy ocular surface conditions were recruited. Dry eye was diagnosed based on the Chinese Expert Consensus on dry eyes (97%), and normal controls (NC) with healthy ocular surface conditions were included as the NC group after ruling out dry eyes on a slit-lamp examination. The exclusion criteria included a history of uveitis, glaucoma, retinal disease, ocular trauma/transplantation in the previous 4 weeks, application of antibiotic or immunomodulatory eyedrops in the previous 4 weeks, and eye surgeries within 3 months. A sterile cotton swab was used to collect specimens by rubbing the swab from the medial to the lateral side of the inferior fornix of the conjunctival sac of each right eye without anesthesia. The swabs were then placed in sterile tubes and stored in a refrigerator (at −20°C) before further experiments. The study adhered to the tenets of the Declaration of Helsinki and was approved by the Institutional Review Board of Peking Union Medical College Hospital (ZS-3092). Informed consent forms were obtained from all patients.

DNA Extraction, PCR Amplification, and 16S rRNA Gene Amplicon Sequencing

Deoxynucleobionic acid (DNA) was extracted using the MicroElute Genomic DNA Kit (D3096, Omega, MA, USA) according to the manufacturer’s instructions. The concentration of DNA was measured using a NanoDrop 2000 ultramicrospectrophotometer (Thermo Scientific, Waltham, MA, USA). The V3–V4 region of the 16s ribosomal RNA (rRNA) gene was amplified from each extracted genomic DNA samples with primers (319 F: 5′-ACTCCTACGGGAGGCAGCAG-3′ and 806 R: 5′-GACTACHVGGGTWTCTAAT-3′) manufactured by Sangon Biotech (Shanghai) Co., Ltd. PCR was carried out on a Mastercycler Gradient (Eppendorf, Germany) using 25 µl reaction volumes, containing 12.5 µl 2×Taq PCR MasterMixII (Vazyme Biotech Co., Ltd, China), 3 µl BSA (2 ng/µl), 1 µl Forward Primer (5 µM), 1 µl Reverse Primer (5 µM), 2 µl template DNA, and 5.5 µl ddH2O. The PCR amplification products were purified with AgenCourt AMPure XP magnetic beads (Fisher Scientific, Hampton, NH, USA), dissolved in an elution buffer, and then labeled. The fragment range and the sequence length of the library were detected using the Agilent2100 Bioanalyzer (Agilent, Santa Clara, CA, USA). Qualified libraries were selected for sequencing on the MiSeq PE300 platform based on the size of the inserted fragments.

Bioinformatic Analysis

Samples were sequenced on an Illumina MiSeq platform (Illumina, Inc., San Diego, CA, USA) according to the manufacturer’s instructions. Qualified paired-end reads were separated using the sample-specific barcode sequences and trimmed with the Illumina Analysis Pipeline Version 2.6 (Illumina, Inc., San Diego, CA, USA). Then, the dataset was analyzed using the Quantitative Insights Into Microbial Ecology (QIIME) (Version 1.8.0; Boulder, USA). The sequences were clustered into operational taxonomic units (OTUs) at a similarity ≥97%, to generate rarefaction curves and to calculate the richness and diversity indices. The Ribosomal Database Project (RDP)
A total of 5,368,240 high-quality sequences were generated from the 98 samples of conjunctival sac swabs, with an average of 54,778 sequences per sample. High-quality sequences were clustered into 5,031 OTUs at 97% sequence identity. A modified OTU table was obtained consisting of 4,904 OTUs (ranging from 36 to 693 per sample), corresponding to 51 phyla, 142 classes, 315 orders, 523 families, and 1,028 genera.

The OTUs were compared with a Venn diagram (Supplementary Figure S1) and 811 common OTUs were identified. There were 60 (2,090/3,512), 22 (544/2,442), and 17% (306/1,816) OTUs that were unique in the NC group, NSSDE group, and SSDE group, respectively.

### Alpha and Beta Diversity Analysis

Alpha diversity was employed to analyze the complexity of species diversity for each sample through Chao1, Shannon indexes generated by QIIME. Beta diversity was demonstrated by the principal coordinates analysis (PCoA) and Partial Least Squares Discrimination Analysis (PLS-DA) to evaluate the microbiome complexity between samples. The taxonomy and relative abundance were bioinformatically analyzed at phylum and genus levels.

### Statistical Analysis

The R software (Version 3.2.5; New Zealand) and GraphPad Prism 5.0 (GraphPad Software, San Diego, CA, USA) were used for the statistical analyses. Tukey test was used to identify significant between-group differences for alpha-diversity. The divergence between the two groups was compared by Analysis of Similarity (ANOSIM). The relative abundance of bacteria was compared by one-way ANOVA. The demographic and clinical data are expressed as mean ± SD. A P-value <0.05 was considered statistically significant.

### RESULTS

#### Demographic Characteristics of Patients

Samples were collected from 23 eyes of SSDE, 36 eyes of NSSDE, and 39 eyes of NC. The mean age was 48.09, 39.89, and 35.61 years old in the SSDE, NSSDE, and NC groups, respectively (Table 1). The average length of Schirmer’s test was 5.72 ± 6.25 (mm/5 min) in the SSDE group and 6.68 ± 8.63 (mm/5 min) in the NSSDE group. The average FBUT was 2.32 ± 1.34 (s) in the SSDE group and 4.71 ± 2.79 (s) in the NSSDE group. The average cornea staining score was 6.67 ± 3.32 in the SSDE group and 2.32 ± 1.03 in the NSSDE group (Table 1).

#### Next Generation Sequencing (NGS) Data

A total of 5,368,240 high-quality sequences were generated from the 98 samples of conjunctival sac swabs, with an average of 54,778 sequences per sample. High-quality sequences were clustered into 5,031 OTUs at 97% sequence identity. A modified OTU table was obtained consisting of 4,904 OTUs (ranging from 36 to 693 per sample), corresponding to 51 phyla, 142 classes, 315 orders, 523 families, and 1,028 genera.

The OTUs were compared with a Venn diagram (Supplementary Figure S1) and 811 common OTUs were identified. There were 60 (2,090/3,512), 22 (544/2,442), and 17% (306/1,816) OTUs that were unique in the NC group, NSSDE group, and SSDE group, respectively.

| Sex      | SSDE | NSSDE | NC     | P-value |
|----------|------|-------|--------|---------|
| Male/female | 0/23 | 9/27  | 12/27  | P = 0.001 |
| Age (years) | 48.09 ± 9.01 | 39.89 ± 13.45 | 35.61 ± 11.03 | P = 0.008 SSDE vs. NSSDE, P = 0.007 NC vs. NSSDE, P < 0.001, NC vs. SSDE |
| Schirmer (mm) | 5.72 ± 6.25 | 6.68 ± 8.63 | ≥10 | P = 0.018 (NSSDE vs. SSDE) |
| FBUT (s) | 2.32 ± 1.34 | 4.71 ± 2.79 | ≥10 | P < 0.001, SSDE vs. NSSDE |
| Cornea staining score | 6.67 ± 3.32 | 2.32 ± 1.03 | 0 | P < 0.001, SSDE vs. NSSDE |

The goods coverage values were all ≥0.99, indicating that the 16S rRNA sequencing results from each library represented the majority of bacterial species present within test samples. The species accumulation curves flattened (Supplementary Figure S2), demonstrating that the sample size is adequate to represent the overall bacterial diversity in the target population.

The alpha diversity was analyzed by the Shannon and chao1 indexes. The Shannon index was significantly lower in the SSDE group (P = 0.020) and NSSDE (P = 0.029) groups compared with the NC group but did not show a statistically significant difference between the NSSDE group and the SSDE group (P = 0.089) (Figure 1A). The chao1 index showed the same tendency as the Shannon index (NC vs. SSDE: P = 0.003, NC vs. NSSDE: P = 0.091, NSSDE vs. SSDE: P = 0.036) (Figure 1B).

There was significant divergence in beta diversity between each group as demonstrated by PCoA (Figure 2A) and PLS-DA (Figure 2B). The samples in the control group were more centralized and resembled each other in the bacterial composition, while the samples in the NSSDE and SSDE groups were more acentric and disperse. This differentiation was further confirmed by ANOSIM analysis, which showed the difference between the groups was greater than the difference within the group (NC vs. SSDE: P = 0.001, NC vs. NSSDE: P = 0.001, NSSDE vs. SSDE: P = 0.005) (Table 2).

### Bacterial Relative Predominance

We summarized the relative abundances of the dominant bacterial community in each group. At the phylum level, 38 phyla were detected from the 23 eyes of the SSDE group, 36 phyla from the 36 eyes of the NSSDE group, and 48 phyla from the 39 eyes of the NC group. The top 5 abundant phyla were Firmicutes, Proteobacteria, Actinobacteriota, Bacteroidota, and Cyanobacteria in all three groups (Figure 3A). The relative percentage of each predominant phylum was demonstrated in Table 3. The Firmicutes/Bacteroidetes (F/B) ratio, which was correlated with inflammation in gut microbiome studies, was calculated. The F/B ratio was 6.42, 7.31, and 9.71 in the NC, NSSDE, and SSDE groups, respectively (P = 0.038, SSDE vs. NC, P = 0.991, NSSDE vs. NC, P = 0.048, SSDE vs. NSSDE).
Among the most abundant phyla, the relative abundance was also different between groups (Figure 4A). The relative abundance of Actinobacteriota was higher in the SSDE group ($P = 0.005$, SSDE vs. SSDE, $P < 0.001$, SSDE vs. NC), followed by the NSSDE group ($P = 0.020$, NC vs. NSSDE) and NC group. Cyanobacteria were more abundant in the NC group ($P < 0.001$, NC vs. NSSDE, $P < 0.001$, NC vs. SSDE), followed by the NSSDE group and SSDE group ($P = 0.680$). Bacteroidota was higher in abundance in the NC group than SSDE ($P = 0.011$), but there was no statistical difference between NC and NSSDE ($P = 0.110$) or NSSDE and SSDE ($P = 0.250$). Proteobacteria was more abundant in the NSSDE group than the SSDE group ($P = 0.014$) and NC group ($P = 0.042$), but its relative abundance was similar between the SSDE and NC group ($P = 0.443$).

At the genus level, 476 genera were detected in the SSDE group, 615 genera in the NSSDE group, and 928 genera.
in the NC group. The top five common abundant genera were *Acinetobacter*, *Staphylococcus*, *Bacillus*, *Corynebacterium*, and *Clostridium_sensu_stricto_1* (Figure 3B). And the relative percentage of each predominant genera was demonstrated in Table 4. *Bacillus* was significantly higher in the NC group compared with the dry eye groups (*P* < 0.001, NC vs. NSSDE, *P* < 0.001, NC vs. SSDE), whereas *Acinetobacter* was significantly higher in the dry eye groups compared with the NC group (*P* < 0.001, NC vs. NSSDE, *P* = 0.001, NC vs. SSDE). The relative abundance of *Acinetobacter* and *Bacillus* was similar between the SSDE and NSSDE groups (*P* = 0.096 for *Acinetobacter*, *P* = 0.727 for *Bacillus*). *Corynebacterium*, *Clostridium_sensu_stricto_1*, and *Geobacillus* were more abundant in the DED groups than the NC (*Corynebacterium*: *P* = 0.003, SSDE vs. NC, *P* < 0.001, NSSDE vs. NC; *Clostridium_sensu_stricto_1*: *P* = 0.039, SSDE vs. NC, *P* < 0.001, NSSDE vs. NC; *Geobacillus*: *P* < 0.001, SSDE vs. NC, *P* < 0.001, NSSDE vs. NC). The top 20 genera with the most significant differences were demonstrated in Figure 4B.

**DISCUSSION**

In recent years, the association of dysbiosis with inflammation and infection has become increasingly recognized (16). On the ocular surface, the antimicrobial activity of tears prevents infection while maintaining a commensal bacterial population (17). In this study, we sampled the conjunctival sac to explore the ocular surface microbiome composition and showed that dry eyes with and without SS have less microbiome diversity than eyes with healthy ocular surface conditions, and have characteristic microbiome composition.

In previous studies, the richness of the conjunctival sac microbiome in patients with dry eyes was inconsistent. While some studies revealed an increased number of bacteria (18, 19), most studies were in accordance with our study that patients with dry eyes have diminished ocular surface microbiome diversity (20–23). In studies regarding SS, bacterial diversity was diminished both for the oral mucosa (24) and the ocular surface (23). The exact mechanism is still unclear but might be an interaction between the mucosa and the commensal bacteria. The microbiome on the ocular surface degrades intracellular mucins, which in turn inhibit bacterial growth, contributing to homeostasis between the ocular microenvironment and commensal bacteria (17). Disruption of either the two ends of homeostasis might cause unbalanced growth of bacteria. In this study, while the major microbiome was stable, the diversity was severely diminished and an aberrant bacterial composition in dry eye patients was demonstrated by the observed clustering and separation demonstrated by the PCoA and PLS-DA analysis.

At the phylum level, *Firmicutes*, *Proteobacteria*, *Actinobacteria*, *Bacteroidota*, and *Cyanobacteria* were the top five phyla in all three groups. These results were consistent with other studies that collect samples from the conjunctival sac (22, 25–27), indicating a relative stable predominant phylum composition of the ocular surface even in different disease status. However, there was a common issue that needs to be addressed in these studies, including ours, that there was no blank control to rule out environmental contamination. *Cyanobacteria* is photosynthetic and was not supposed to be found in the conjunctival sac which has no sun exposure. One possible explanation was that there might be *Cyanobacteria* on the ocular surface and dropped into the conjunctival sac. The 16srRNA method could also identify dead microbiomes as well.

**TABLE 2** | Analysis of similarity (ANOSIM) for diversity between groups.

| Group         | R statistic | P-value |
|---------------|-------------|---------|
| NC-NSSDE      | 0.5485      | 0.001   |
| NC-SSDE       | 0.6064      | 0.001   |
| NSSDE-SSDE    | 0.1195      | 0.005   |
| all           | 0.4428      | 0.001   |

*R* value > 0 means that differences between groups are greater than differences within groups.

**FIGURE 3** | The top five abundant microbiomes at the phylum (A) and genus (B) level showed similar major components of bacteria in each group, but the relative abundance was different.
TABLE 3 | Percentage of the top five phyla in each group.

| Phylum       | NC (%) | NSSDE (%) | SSDE (%) | P-value     |
|--------------|--------|-----------|----------|-------------|
|              |        | NC vs. NSSDE | NC vs. SSDE | NSSDE vs. SSDE |
| Firmicutes   | 43.89  | 37.97     | 40.52    | 0.155  | 0.288 | 0.706 |
| Proteobacteria| 34.78  | 43.61     | 33.04    | 0.014  | 0.396 | 0.014 |
| Actinobacteriota | 6.60   | 8.24      | 15.89    | 0.020  | 0.281 | 0.005 |
| Bacteroidota | 6.84   | 5.19      | 4.17     | 0.110  | 0.027 | 0.250 |
| Cyanobacteria| 4.35   | 1.51      | 0.96     | <0.001 | 0.005 | 0.680 |

FIGURE 4 | The top 20 significantly different microbiomes at the phylum (A) and genus (B) level.

TABLE 4 | Percentage of the top five genus in each group.

| Genus               | NC (%) | NSSDE (%) | SSDE (%) | P-value     |
|---------------------|--------|-----------|----------|-------------|
|                     |        | NC vs. NSSDE | NC vs. SSDE | NSSDE vs. SSDE |
| Acinetobacter       | 5.34   | 20.58     | 13.05    | <0.001  | 0.001 | 0.096 |
| Staphylococcus      | 7.14   | 11.43     | 13.55    | 0.316   | 0.543 | 0.859 |
| Bacillus            | 22.26  | 3.26      | 2.09     | <0.001  | <0.001 | 0.727 |
| Corynebacterium     | 3.82   | 4.15      | 8.62     | 0.076   | 0.003 | 0.050 |
| Clostridium_sensu_stricto_1 | 0.61   | 9.38      | 4.80     | <0.001  | 0.003 | 0.019 |

Thus, Cyanobacteria might also be detected. Supplementary experiments in further studies are needed to give a more pronounced answer.

The relative abundance at the phylum level was different in different groups. Actinobacteriota was more predominant in patients with dry eyes, while Cyanobacteria was less predominant, compared with healthy eyes. These differentiations were further magnified by the etiology of dry eyes. Patients with SSDE had more abundant Actinobacteriota and less Cyanobacteria compared with patients with NSSDE. We assumed that this might be associated with the severity of dry eye, as Schirmer’s test, FBUT test, and cornea staining all showed a more severe degree of dry eye damage to the ocular surface. Proteobacteria was markedly higher in the NSSDE group compared with the SSDE group, and this was in accordance with the other study addressing the different microbiome.
compositions in DED with and without the autoimmune disease (23).

Although the relative abundance of Firmicutes and Bacteroides were similar in these three groups, we calculated the F/B ratio, inspired by the fact that a lower F/B ratio in the gut was associated with the inflammation status in the intestine such as in inflammatory bowel disease (28) and other systemic diseases such as systemic lupus erythematosus (29), while a higher F/B ratio in the gut was associated with obesity (28). Previous studies also showed that the F/B ratio in the gut was lower in patients with SSDE than in NC (30). However, in this study, we showed that the F/B ratio of the conjunctival sac was higher in the SSDE group compared with NC and NSSDE groups, in contrast with the gut F/B ratio. The F/B ratio change in both directions is a sign of dysbiosis (28). It is hypothesized that disrupted microbiome composition might activate distinct immunomodulatory and inflammatory pathways, which result in the instability of tear film and ocular damage. But we have to acknowledge that the gut has a large amount of microbial load while the ocular surface microbiome is paucibacterial. Thus, the effect of microbiome dysbiosis on the ocular surface might be less predominant. Further studies are needed to provide more evidence on the F/B ratio change and its relationship with SSDE and NSSDE.

At the gene level, patients with dry eyes had less genus variety. The most common five abundant genera were Acinetobacter, Staphylococcus, Bacillus, Corynebacterium, and Clostridium_sensu_stricto_1 (Figure 3B), similar to the results of other studies concerning the ocular surface bacteria (22, 23, 25–27, 31). Compared with the NC group, patients with dry eyes had more abundant Acinetobacter but less Bacillus. Acinetobacter was commonly detected on the ocular surface of critically ill patients, who were always sedated and had unhealthy ocular surface function (32). However, previous studies showed that Bacilli are prominent in patients with dry eye disease, in contrast with the decreased Bacilli in DED groups in our study. Certain Bacillus species can synthesize and secrete lipopeptides, in particular, surfactins and mycosubtilins (33, 34), which might affect tear film stability. Further research needs to be done to explain the Bacillus variation in different studies and its role in dry eye disease. Corynebacterium was significantly higher in abundance in the SSDE group than the NSSDE group, which was in line with the study that compares bacteria composition in patients with dry eyes with and without autoimmune diseases (23). This provided further evidence that Corynebacterium might be associated with autoimmunity, as mycolic acids and the cell wall architecture of Corynebacterium can affect macrophage function (35). However, there was also a study demonstrating that a decreased amount of Corynebacterium in patients with meibomian gland dysfunction (MGD) type of dry eye (36), supported by evidence that Corynebacterium elicited interleukin 17 response from γδ T cells in the ocular mucosa, driving neutrophil recruitment and antimicrobials release into the tears (37). These pieces of evidence were not contradictory, as the former one was a comparison between the etiology of dry eye disease, while the latter focuses on the inflammation status of MGD. Clostridium_sensu_stricto_1 and Geobacillus were also higher in the dry eye groups in our study, but the association with functional microbiology and clinical significance need to be further addressed.

In previous studies regarding MGD, the overgrowth of Staphylococcus and Sphingomonas were detected (36), however, in this study, the amount of Staphylococcus was similar in the three groups; Sphingomonas was more abundant in the NC group. The different findings might be caused by different diagnostic criteria for DED, sampling methods (25), as well as different seasonal regions patients come from (38).

This study has several limitations. First, the F/B ratio at the phylum level was proposed for the first time in our study regarding the field of conjunctival sac microbiome. Although patients in the SSDE group have significantly higher F/B, the intrinsic association with SSDE and its opposite change in the gut of patients with SSDE need to be further investigated. Secondly, this was a single-center study that recruits patients from a certain area with a small sample size. The enrolled population was also homogeneous (mainly Han Chinese) to provide generalized conclusions. Further studies involving more diverse populations are expected. Thirdly, as SSDE more commonly affects mid-aged women, it was hard to have SSDE samples from male patients to match the sex and age balance. The influence of age and sex on microbiome composition was controversial (22, 36, 39). Further studies were expected to address this issue.

CONCLUSIONS

In conclusion, we identified that the diversity of ocular surface microbiome in patients with NSSDE and SSDE was diminished compared with NC. The main microbiome at the phylum and genus level was similar between groups, but the relative abundance of the top five phyla and genera had variations. F/B ratio was significantly higher in the SSDE group compared with NC and NSSDE group.

DATA AVAILABILITY STATEMENT

The datasets presented in this study can be found in online repositories. The names of the repository/repositories and accession number(s) can be found below: https://www.ncbi.nlm.nih.gov/bioproject/, PRJNA802065.

ETHICS STATEMENT

The studies involving human participants were reviewed and approved by the Institutional Review Board of Peking Union Medical College Hospital. The patients/participants provided their written informed consent to participate in this study.

AUTHOR CONTRIBUTIONS

HS and KX wrote the manuscript. HS and QL analyzed the data. ZC applied for the ethical approval of the study. QL designed
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SUPPLEMENTARY MATERIAL

The Supplementary Material for this article can be found online at: https://www.frontiersin.org/articles/10.3389/fmed.2022.841112/full#supplementary-material

Supplementary Figure S1 | Venn diagram showing the number of operational taxonomic units (OTUs) that each group consisted of and its relationship with other groups.

Supplementary Figure S2 | Species accumulation analysis showing the increase in OTUs detected with the addition of each sample.
Conflict of Interest: The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

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