Bacterial Community Composition of Sapanca Lake During a Cyanobacterial Bloom

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Cite this article as: Özbayram, E. G., Köker, L., Akçaalan, R., İnce, O., Albay, M. (2020). Bacterial community composition of sapanca lake during a cyanobacterial bloom. Aquatic Sciences and Engineering, 35(2), 52–56.

ABSTRACT
Microbial community compositions and functions of freshwater ecosystems vary due to the environmental parameters and water chemistry. Transient bloom events play a crucial role in the community profiles. In this study, a specific focus was set to provide a snapshot of the bacterial community composition in Lake Sapanca, associated with cyanobacterial bloom by high throughput sequencing method. For this purpose, a sample was collected in the shore of Lake Sapanca during a cyanobacterial bloom, and the bacterial community profile was examined by 16S rRNA amplicon sequencing using the Illumina MiSeq platform. Cyanobacteria represented 94% of the all reads. The bacterial community was re-calculated to evaluate the bacterial diversity in detail by filtering cyanobacterial sequences. The community was dominated by Proteobacteria (44%) and Bacteroidetes (33%) species which are abundant in freshwater ecosystems having an ability to degrade complex organics. Among the classified genera, Flavobacterium and Rheinheimera dominated the bacterial community suggesting a strong link between those species and the cyanobacterial bloom. The experimental work presented here provides one of the first investigations of total bacterial communities in Lake Sapanca by the high throughput sequencing method. Further work is needed with more sampling points and time series to fully understand the bacterial diversity and dynamics.

Keywords: Bacterial community, cyanobacterial bloom, illumina miseq, Sapanca lake, 16S rRNA

INTRODUCTION
Freshwater habitats have a vital role in global biogeochemical cycles. However, cyanobacterial blooms are getting more widespread in freshwater lakes due to nutrient runoff and climate change, and have become a serious risk on the sustainability of these ecosystems (Cai et al., 2014; Liu et al., 2019) which cause difficulties to secure and maintain ecosystem health (Woodhouse et al., 2016).

Microorganisms drive crucial functions in the freshwater ecosystems and have a role in the degradation of organic materials, energy conversion and nutrient recycling, overall contribute to the ecosystem balance (Su et al., 2017a; Zhu et al., 2019). Microbial community compositions and functions vary due to the water chemistry, nutrient concentrations, hydrodynamic stability and climate (Steffen et al., 2012), and they respond to environmental alterations quickly being critical indicators (Su et al., 2017b). Furthermore, community structures are highly dependent on transient bloom events which affect the abundance and activity of these communities (Eiler & Bertilsson, 2004). Since microbial communities are key players on biogeochemical cycles, there is still limited information on the characterization and function of bacterial communities inhabiting freshwater
ecosystems, especially during bloom events (Eiler & Bertilsson, 2004; Steffen et al., 2012).

Recent developments in advanced molecular genetic technologies enable us to have deep information about microbial diversity and interactions between the community structure and their environment (Nakatsu, Byappanahalli, & Nevers, 2019). Since the 16S rRNA gene is considered as a molecular marker of prokaryotes and used to investigate microbial communities in a wide variety of habitats, it has also been used to assess microbial community dynamics during cyanobacterial blooms (Nakatsu et al., 2019; Zhu et al., 2019).

Lake Sapanca, a freshwater lake, is located in the northeast of the Marmara Region of Turkey. It is a source for drinking water for the cities of Sakarya and Kocaeli, as well as for industrial usage (Akçaalan et al., 2014; Leroy & Albay, 2010), and has been monitored according to the presence of cyanobacteria and cyanotoxin for years to indicate the ecological status of the lake, and it has now a well-established cyanobacteria diversity (Akçaalan et al., 2007; Akçaalan et al., 2014). Moreover, there are some studies in the literature on Lake Sapanca which evaluated the total bacterial counts, pathogenic bacteria and petroleum-resistant bacteria presence by a culture dependent technique (Altuğ et al. 2006; Çiftçi Türetken et al., 2018), and seasonal dynamics of pathogens by the microarray method (Akçaalan et al., 2018). So far, there have been no attempts to examine the total bacterial community structure in Lake Sapanca, and it has remained unclear. This study, therefore, sets out to assess the bacterial community composition in Lake Sapanca during a cyanobacterial bloom by the high throughput sequencing method. The experimental work presented here provides one of the first investigations into the bacterial community profile of Lake Sapanca by a next generation sequencing platform.

MATERIALS AND METHODS

Physico-chemical characterization
A surface cyanobacterial bloom occurred in Lake Sapanca on 8 April, 2019 and the sample was collected during the bloom event in the shore. pH, temperature and dissolved oxygen were measured with a portable multiparameter (6600, YSI, USA) on the sampling date.

DNA extraction and amplicon sequencing
First, 10 mL of the sample was filtered with a 0.22 µm filter, and total DNA was extracted from that filter paper using a MoBio PowerWater® DNA Isolation Kit (MoBio Laboratories, Inc., CA, USA) according to the manufacturer’s protocol. The DNA quantification was performed by NanoDrop 1000 (Thermo Fisher Scientific, Inc., DE, USA), and the extracted DNA was stored at -20°C for further analysis.

The bacterial community composition of the sample was analyzed with the ZymoBIOMICS™ Service - Targeted Amplicon Sequencing (Zymo Research, Irvine, CA). 16S ribosomal RNA gene targeted sequencing was performed using the Quick-16S™ NGS Library Preparation Kit (Zymo Research, Irvine, CA). Shortly, the 16S primers used amplification of the V3-V4 region of the 16S rRNA gene (341f-CCTACGGGNGGCWGCAG and 805r-GACTACHVG-GTATCTAATCC). The PCR products were quantified with qPCR fluorescence readings, and pooled together based on equal moi- larity. The final pooled library was cleaned up with Select-a-Size DNA Clean & Concentrator™ (Zymo Research, Irvine, CA), then quantified with TapeStation® and Qubit®. The final library was sequenced on Illumina® MiSeq™ with a v3 reagent kit (600 cycles). The sequencing was performed with >10% PhiX mix and in paired-end mode.

The Dada2 pipeline was used to infer the amplicon sequences from raw reads (Callahan et al., 2016). The raw sequence reads were trimmed with Trimmomatic-0.33 (Bolger, Lohse, & Usadel, 2014). Whereas, SeqPrep were used to assemble the two paired-end reads to have a complete amplicon sequence with (https://github.com/jstjohn/SeqPrep). Usearch (v. 6.1) was used to check and remove chimeric amplicon sequences (Edgar, 2010) in ref mode against a curated database (http://drive5.com/uchime/rdp_gold.fa). Amplicon sequences smaller than 320 bp were removed. For each sample, up to 40,000 sequences were randomly sampled to reduce the potential bias caused by uneven sampling. These amplicon sequences were compiled, clustered and analyzed with Qiime 1.9.1 (Caporaso et al., 2010). OTUs were picked by the workflow of pick_open_reference_otus.py using the GreenGene database (gg_13_8) as reference database. Singleton OTUs were removed. Taxonomy assignment was performed with Qime v.1.9.1 (Caporaso et al., 2010). The microbial community structures were shown by Krona graphs (Ondov, Bergman, & Phillippy, 2011; Ozbayram et al., 2017).

RESULTS AND DISCUSSION

The physical properties of Lake Sapanca during the bloom are depicted in Table 1. The water was slightly alkaline and the characteristics of the lake matched those observed in early studies (Akçaalan et al., 2007, 2014) showing a typical O₂ saturation level during the bloom event with high dissolved oxygen.

| Parameter          | Lake Sapanca |
|--------------------|--------------|
| Temperature (°C)   | 15.9         |
| pH                 | 8.51         |
| Conductivity (µS/cm) | 215.6     |
| Dissolved Oxygen (mg/L) | 13.57   |
| O₂ Saturation (%)  | 130.3        |

The number of raw reads and after filtration were 78,735 and 65,748, respectively and the rarefaction curve reached a plateau. The microbial community composition of the bloom sample was presented at multiple taxonomic levels by a Krona chart in Figure 1. The microbial community comprised 7 phyla, however, the microbial community was dominated by Cyanobacteria members, representing 94% of all sequences as it was expected. At the genus level, all of the Cyanobacteria reads belonged to Planktothrix. The results are in keeping with previous observational studies,
which showed that the cyanobacterial bloom was mainly caused by *Planktothrix rubescens* in Lake Sapanca (Akçaalan et al., 2007, 2014). Proteobacteria and Bacteroidetes were the following phyla, representing 5% of the microbial community. The results are in agreement with those of previous studies, indicating that the bacterial community was dominated by Proteobacteria, Actinobacteria, and Bacteroidetes during the phytoplankton bloom (Berg et al., 2009; Eiler, Bertilsson, & Centre, 2007; Zhu et al., 2019).

To understand the microbial community diversity better, the bacterial reads were evaluated excluding Cyanobacterial sequences, and re-calculated relative abundances of the bacterial community are depicted in Figure 2. As it is clear from the chart, more than half of the reads were represented by Proteobacteria (44%) and Bacteroidetes (33%). Actinobacteria was the third dominant phylum, representing only 5% of the bacterial community, followed by Planctomycetes (5.3%), Verrucomicrobia (5%) and Gemmatimonadetes. Proteobacteria and Bacteroidetes have the ability to become abundant in the presence of bioavailable organic material, and dominate the community (Eiler et al., 2007). The members are known to be able to decompose complex organic materials, and the peptides of the organic carbon plays a crucial role with their ABC membrane transporters which may support toxin degradation (Lezcano et al., 2017).

Within Proteobacteria, Betaproteobacteria was the dominant class, representing almost half of the total reads in Proteobacteria, followed by Gammaproteobacteria and Alphaproteobacteria. Betaproteobacteria was found as abundant in freshwater ecosystems (Zhu et al., 2019). Flavobacteria (29%) dominated the bacterial community, accounting for 25% of the total sequences (Figure 3). The high abundance suggests that a strong link may exist between the bloom and Flavobacterium species which can degrade various biomacromolecules and carbohydrates. The members react to transient nutrient loads immediately, which is a result of phytoplankton blooms (Buchan, LeCleir, Gulvik, & González, 2014). Whereas some of Flavobacterium species have a potential to degrade cyanotoxin, some of them are reported to have a role in denitrification (Parulekar et al., 2017). Rheinheimera was the second most abundant genus, representing 9% of the bacterial community. Rheinheimera has also higher abundances in the aquatic environments and can hy-
drollyze organic materials. It is speculated that, Rheinheimera species can regulate phosphate exchange in the cyanobacterial mucilage capsule resulting in the enhancement of Microcystis growth (Parulekar et al., 2017).

CONCLUSION

The present research explores, for the first time, the bacterial communities associated with cyanobacterial blooms in Lake Sapanca by 16S rRNA targeted amplicon sequencing. This study has shown that the bacterial community was dominated by bloom-associated phyla, Proteobacteria and Bacteroidetes, having the ability to grow on complex organic materials.

These findings provide a snapshot of the bacterial communities in Lake Sapanca during the cyanobacterial bloom. Further work is needed, with more sampling, to fully understand the bacterial diversity and dynamics.

Conflict of Interest: The author has no conflicts of interest to declare.

Ethics Committee Approval: Ethics committee approval is not required.

Financial Disclosure: The authors declared that this study received no financial support.

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Figure 3. The dominant bacterial genera representing more than 5% of the all sequences are shown in colors. The grey part represented the rest of the bacterial genera which represented <5% of the all sequences.

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