Community dynamics of prokaryotic and eukaryotic microbes in an estuary reservoir

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This study demonstrates both prokaryotic and eukaryotic community structures and dominant taxonomies in different positions of the greatest estuary reservoir for drinking water source in the world in four seasons of one year using 454 pyrosequencing method with total of 312,949 16S rRNA and 374,752 18S rRNA gene fragments, including 1,652 bacteria OTUs and 1,182 fungus OTUs. During winter and spring, the community composition at the phylum level showed that microorganisms had similar structures but their quantities were different. Similarly, obvious changes at the genus level were observed among the samples taken in winter and spring between summer and fall. Microorganisms located the reservoir inlet were founded to be different from those in rear at both phylum and genus level. Air temperature had a stronger effect than sampling location on the microbial community structure. Total nitrogen and dissolved oxygen were algae-monitoring indicators during the whole year. Moreover, Bacillus was an efficient indicator during summer and autumn for bacteria OTUs.

Estuary ecosystems are important water sources for numerous cities, particularly world-class, densely populated metropolitan areas located at the junction of river and ocean, such as New York, London, Tokyo, Singapore, Hong Kong, and Shanghai¹. Microbial communities are a key constituent of the microbial food loop of estuary reservoirs, being a decomposer, consumer, primary producer and secondary producer. Water quality is strongly influenced by microbial community dynamics and ecosystem functions such as organic matter content and nutrient recycling. Moreover, these changes in the microbial community trigger algae blooms and off-flavor problems, which negatively affect the water supply. Understanding the community composition and time-site distribution of estuary reservoir microorganisms increases the current understanding of the distribution characteristic and its function in the estuarine ecosystem, and is particularly helpful in judging the environmental safety of the water supply as a source of drinking water.

Microscopic identification and plate cultivation are used to monitor and characterize the microbial diversity of aquatic ecosystems, but these two methods cannot differentiate uncultured species. Some biochemical methods, such as phospholipid fatty acids (PLFA) analysis, are used to characterize the community structure of micro-biota, including both prokaryotes and eukaryotes. However, PLFA analysis has certain limitations, such as overlap in the composition of microorganisms. Both denaturing gradient gel electrophoresis fingerprinting (DGGE) and clone library analyses have made independent cultivation possible. However, these tools usually underestimate the overall diversity of a microbial community and are difficult to use to obtain a comprehensive community profile in complex environments. Fortunately, pyrosequencing, a high-throughput sequencing technology, has gained increasing research interest as a novel tool for examining community structure. Moreover, several researchers showed a general comparison about four basic approaches: miTags, m454Tags, 454Tags and iTags, advocated the technique miTags (metagenomic illumina 16S tags) to acquire a large amount of functional data and avoid a mass of PCR biases. However, many difficulties existed in metagenomic analysis due to variety of genes, variance at evolutionary rates, complex in computations, especially absence of appropriate reference gene sets. Therefore, 454 Pyrosequencing are more fit for RNA diversity analyses.

Similar to prokaryotes, including eukaryotic algae, fungus and macrophytes have also been closely associated with water environment problems, such as algae bloom and off-flavor of freshwater. Principal component analysis (PCA) has shown that phytoplankton could be an indicator of improved water quality at the Golden Horn Estuary. Certain fungi species have been associated with the production of the mycotoxin patulin and the odor secondary metabolite. A study by Kravehuk et al suggested that macrophytes were a positive factor in...
Results

Environmental parameters. Supporting Information (SI) Table S1 lists the environmental parameters, including water temperature, dissolved oxygen (DO), permanganate index (COD Mn), total nitrogen (TN), total phosphorus (TP), transparency, and chlorophyll-a, microcystin-RR (MC_RR) and microcystin-LR (MC_LR). Transparency increased whereas TP decreased from the head to the tail of the reservoir (Fig. 1). COD Mn and chlorophyll-a concentrations were higher in Site 3 than in Site 2 (Fig. 1). The chlorophyll-a of Site 0 was always lower than those of other sites and dramatically influenced by air temperature as well as COD Mn, particularly by air temperatures ranging from 22°C to 30°C, and water quality that meets the current national “Sanitary standard for drinking water” (GB5749-2006). Water temperature varied more significantly in periods. Irregular changes in TN were observed in the six sites. The DO of all samples were relatively stable at approximately 9.11 mg/l. MC_RR and MC_LR were lower than 0.5 μg/l in most samples, with only the MC_LR in sample 0410-3 reaching 0.77 μg/l, which was still lower than the national standard of 1 μg/l.

Overall taxonomic richness. A total of 312,949 16S rRNA sequences and 374,752 18S rRNA sequences were selected for classification. Dominant length distributions were approximately 429 and 420 bp, respectively. The taxonomic distributions of the 16S rRNA sequences obtained in the water samples are shown in Fig. 2a. The majority of bacteria sequences (105,673 sequences) belonged to Cyanophyta. The second largest group was Actinobacteria (49,298), followed by Firmicutes (40,013). Betaproteobacteria and Alphaproteobacteria were the most abundant Proteobacteria, making up approximately 56% and 30%, respectively, of the Proteobacteria sequences (43,091). Based on the 97% species similarity, a total of 1,652 OTUs derived from 280,382 sequences were detected in this study. Among these OTUs, 309 were Bacteroidetes OTUs (18.7%), 233 were Cyanophyta (14.1%), and 198 were Alphaproteobacteria (12.0%). The taxonomic distributions of the 18S rRNA sequences obtained in the water samples are shown in Fig. 2b. The majority of fungal sequences (129,819 sequences) belonged to Metazoa. The second largest group was Cryptomonadales (72,845), followed by Chlorophyta (30,194). Eukaryotic algae included Cryptomonadales, Chlorophyta, Diatomea, Dinoflagellata, Kathablepharidae, Chrysophyceae, Glaucophyta, Stramenopiles, and Prymnesiophyceae (126,631 sequences, accounting for 56.4% of total sequences). Similarly, a total of 1,182 OTUs derived from 357,732 fungal sequences were detected in this study. Among these OTUs, 177 were Cerozoa OTUs (15.0%), 129 were Chlorophyta (11.0%), 93 were Metazoa (7.9%), 53 were Cryptomonadales (4.5%), and 27 were Diatomea (2.3%).

Effects of season-site treatments at phylum levels on community. Shifts in the proportions of sequences for different bacteria phyla are shown in Fig. 3a. In general, different samples showed a relative similar trend in terms of richness as well as indicated by Chao1 and Ace indexes during the same season (S2a). The composition and biomass of taxa exhibited a remarkable spatial and temporal heterogeneity with a pronounced longitudinal gradient. Cyanophyta, Proteobacteria, Actinobacteria, and Bacteroidetes were always highest across all water samples, with more than 80% proportion. However, Sites 0 and 2 exhibited greatly decreased relative abundance of Proteobacteria in summer and autumn, as shown in green type. Except for autumn, the proportion of Cyanophyta in Sites 0 and 2 were well below those of other sites. Firmicutes proportions were highest in the summer samples. Changes in the proportion of sequences of different fungal phyla are shown in Fig. 3b, and richness in Spring and Summer significantly higher than others.
(S2b). Interestingly, except for the winter samples, Metazoa had the highest abundance among the samples. Cryptomonadales always existed in all samples, and were highest in number in spring and autumn. Cercozoa were most plentiful during winter, particularly in Sites 3, 8, and 13, accounting for more than 60% in those sites, such a high percentage was associated with low frequency during winter sampling. Chlorophyta had significantly stronger samples in summer than others, similar to Diatomea.

**Effects of season-site treatments at genus level on abundance and taxonomy.** The hierarchical heatmap was based on the top 50 abundant bacterial communities at the genus level, which were classified into four groups (Fig. 4a). The first group was mainly composed of the summer samples, the second group was composed of the autumn samples, the third group was composed of the spring and winter samples from the middle-rear parts of the reservoir, and the last group was composed of the spring and winter samples from the front-middle parts. The most abundant bacteria at genus level were Chloroplast norank (22.8%), hgcI clade (11.8%), Bacillus (7.6%), Synechococcus (7.0%), and Microcystis (2.4%). Microcystis were most abundant in summer at Sites 3 and 8, and Microcystis aeruginosa accounted for only 3.9% of the bacteria. Moreover, Synechococcus always found during summer and fall. Given the same seasons for all of the samples, such as for the summer and autumn samples that were still clustered together, the features of the top 50 fungus genera were less consistent than those of the bacteria (Fig. 4b). The most abundant fungi were Cryptomonas (16.5%), Maxillopoda (13.5%), Mammalia norank (12.4%), Novel_Clade_2_norank (4.0%), and Rotifera norank (9.7%). Telaulax and Chlamydomonas were the three most abundant eukaryotic algae. Furthermore, algae such as Closterium, Chlorophyceae norank, Hafniomonas, Cyclotella, Actinocyclus, and Ceratium, increased sharply during summer, whereas Thalassiosira were not found in the summer samples.

**Statistical characterization of complete pyrosequencing data obtained from estuary reservoir.** Statistical analysis of 16S rRNA showed that because of the different time-site treatments, environmental variables and bacterial characteristics were different between seasons and stations, particularly among the dominant species (Fig. 5a). The total eigenvalues in the 16S rRNA data set accounted for 2,687,740. The total eigenvalues explained by environmental eigenvalues accounted for 779,902, occupying 29.0% of the total RDA explanatory power. Concerning the variation of species data, the first axis explained 55.1%, the first and second axes together explained 76.7%. The first axis explained 16.5% of the variation of species-environment relation, the first and second axes together explained 26.1%. Biplot scaling of redundancy discriminate analysis (RDA) with original data were shown in Fig. 5a and S3a. Both Temperature and MC_RR got 0.001 Pr with 999 number of permutations based on the 1% level in a permutation test for all constrained eigenvalues for prokaryotes. Corresponding values were 0.002 for TN and 0.01 for DO, and greater than 0.05 for transparency, COD Mn, etc. Therefore, four environmental factors, Temperature, MC_RR, TN and DO, were significant environmental variation for bacteria. Moreover, it made a statistically significant contribution from Bacillus to the species-environment relationship (P = 0.001, F=2.9513, number of permutations=1,000). The higher contents of TN and COD Mn as well as increase in chlorophyll-a and Bacillus and in activity were observed for partial samples from Groups 4 and 5 (0828, 0911, and 0913) when the air temperature during sampling decreased from 30.5°C to 28.8°C. The increase in Synechococcus related to activity...
was observed for partial samples of Groups 3 and 4 (0621, 0712, 0725, and 0807). However, the lower air temperature of samples resulted in higher DO values as well as growth in MC_LR. Compared with the effect of sites, these observations indicated that different air temperatures drastically changed the bacterial community structure. Given the rapid growth of *Microcystis aeruginosa* when air temperature decreased from 30.5°C to 28.8°C, researchers should pay more attention to monitoring indices such as Temperature, MC_RR and TN or the sudden abnormal appearance of *Bacillus*.

The statistical analysis of 18S rRNA indicated that the summer and fall samples were located at the right side of the PC2 axes, whereas the winter and spring samples were scattered at the left side (Fig. 5b). Correspondingly the total eigenvalues in the 18S rRNA data set accounted for 4,104,485. The total eigenvalues explained by environmental eigenvalues accounted for 924,151, and the combination of these environmental factors provided an additional 22.5% of the total RDA explanatory power. Concerning the variation of species data, the first axis explained 45.1%, the first and second axes together explained 66.8%. The first axis explained 11.0% of the variation of species-environment relation, the first and second axes together explained 15.3%. Biplot scaling of RDA with original data were shown in Fig. 5b and S3b. Temperature and DO got 0.001 Pr with 999 number of permutations based on 1% level in a permutation test for all constrained eigenvalues for eukaryotes. In addition, none of the other factors made a statistically significant contribution to the species-environment relationship (P = 0.173, F = 1.1948, number of permutations = 1000). During the rapid growth of *Microcystis aeruginosa*, TP was related to *Cryptomonas* and partial samples from Groups 4 and 5, which showed a relative important contribution of TP to the activity of eukaryotic microorganisms. Data revealed that *Actinocyclus* and *Cryptomonas* were highly abundant. *Cryptomonas* accounted for 16.5% of the 18S rRNA sequences, indicating that taxonomy was a key factor in the algae community throughout the year. Rapid proliferation of algae during spring and winter were mainly caused by eukaryotic algae, whereas the activity of *Microcystis* was associated with late summer and early fall. Thus,
Figure 4 | (a–b) Heatmap analysis of prokaryote and eukaryote communities based on the Bray-Curtis similarity of 454 pyrosequencing profiles.

Figure 5 | (a–b) Redundancy discriminant analysis (RDA) biplot of the distribution of prokaryote and eukaryote communities with environmental variables in estuary reservoir. The upper right legend shows color-coding indicating six groups in estuary reservoir. group 1: 0226, group 2: 0328, 0410, 0426, group 3: 0529, 0621, group 4: 0712, 0725, 0807, 0828, group 5: 0911, 0913, 1011, group 6: 1029.
more attention should be given to Temperature and DO when monitoring algae in winter and spring.

Discussion

The Changjiang Estuary was an important interface between terrestrial and marine environments, and also had highly complex and dynamic because of the input of freshwater and seawater alternately. The study on temporal and spatial dynamics of prokaryotic and eukaryotic microbes throughout the whole year by pyrosequencing is very important for understanding the structure and function of the ecosystem for estuarine reservoir. Given the supposedly universal primers and database discrepancy, a number of sequences belonging to taxa different from *Cyanophyta* and other unclassified organisms were detected (Fig. 2a,b). Nevertheless, nearly 105,673 *Cyanophyta* sequences were obtained, accounting for 37.7% of the bacterial sequences and forming 233 OTUs. The results were similar to other pyrosequencing study19 on aquatic ecosystems, that is, the dominant bacteria were *Cyanophyta*, accounting for 32.3% of the total sequences in the Guanting Reservoir. This study the eukaryotic algae accounted for 36.4% of fungal sequences, *Cryptomonas* were the most abundant genun (16.5% of fungal). The results are consistent with those of the other methods18,20,29, that is, phytoplankton were the dominant taxa in a reservoir or a lake. In addition, these discussed results conflicted with previous report21 that most of the OTU's belonged to *Actinobacteria*, followed by *Bacteroidetes* and *Betaproteobacteria* by using the restriction fragment length polymorphism (T-RFLP). The discrepancy between the two reservoirs were mainly caused by the higher percentage of uncultured clones by using different methods except for regional difference. Microbial community diversity was traditionally based on morphology or physiology. 16 different algae by observing phytoplankton biovolume in a subtropical estuary22. A total of 142 phytoplankton taxa were identified from eight algae classes during four years in the Golden Horn Estuary23 using quantitative analysis of phytoplankton, which were significantly less than the 233 prokaryotic algae OTUs and the 209 eukaryotic algae OTUs found by pyrosequencing. Since 1993, DGGE has been employed to study microbial community composition24. DGGE isolated 101 sequences and 46 clones, and DNA-based fingerprinting obtained 45 16S rRNA OTUs and 39 18S rRNA OTUs25. However, compared with the traditional molecular biology methods, the pyrosequencing used in this study obtained 312,949 16S rRNA sequences and 374,752 18S rRNA sequences. These results indicate that the numbers of derived sequences were enough to characterize community diversity and richness, and provide more detailed information about uncultured taxa in the estuary reservoir. In addition, the profile of whole community was often reported by only one type26–28 or several types3,17 of species in previous estuary studies. Species compositions known in estuary ecosystems were much less those of terrestrial40 and marine15,27 ecosystems. The use of pyrosequencing27–28 has proven to be one of the most effectively techniques to determine the identity, and to quantify the abundance of microorganisms, as well as to reveal the profile of whole community accurately in natural system. Hence, the long time pyrosequencing results of reservoir had important values for similar reservoir.

Although the prokaryotic community was dominated by *Cyanobacteria*, *Chloroplast norank* at the genus level accounted for the majority of *Cyanobacteria* in spring. These groups were not found using techniques in microscopy and were not the study subject in the *Cyanobacteria* bloom. The eukaryotic community was dominated by *Cryptomonas* (accounting for 16.5% of sequences). Although Edward et al29 reported that *Cyanobacteria*, *Diatom*, and *Dinoflagellate* were the dominant algae in an aquatic ecosystem. These results were in accordance with the observation of Pavel et al29 that *Cryptophyte* and *Diatom* prevailed in the reservoir from April to mid-June. These complicated categorical information on microbial communities extend the knowledge obtained from previous studies and reports. At a higher resolution, *Synechococcus* and *Cryptomonas* were the dominant genera among *Cyanobacteria* and *Crytophysa*, respectively. *Synechococcus* survive and propagate more easily than the others because of the main participants in the global carbon cycle and the primary contributors to primary productivity, and are particularly abundant in near shore waters. *Cryptomonas* widely exist because of their ability to adapt to a wide range of irradiance and their high tolerance for eutrophication30. Remarkably, a large number of environmental and unclassified *Chloroplast norank* were found in estuary reservoirs, which is consistent with the finding of Zeng et al31 that most *Cyanobacteria* were related to chloroplasts (93.7%). The developments of both pyrosequencing and microbial taxonomy can provide the basis for estuary reservoirs to be a potential water source for a megalopolis. A more interesting study32 indicated that the relationship of phylogeny and function was sufficiently linked that “predictive metagenomic” approach, which provided useful insights into those uncultivated microbial communities such as “no rank species” by using marker gene data. Aside from *Synechococcus*, *Bacillus*, and *Microcystis*, *hgcI clade* and *Limnophibitans* were also abundant in the bacterial sequences. *hgcI clade* belonged to *Actinomyctetales*, and previous case33 reported that such microbes were affected by the odor and taste of the drinking water. *Limnophibitans* belong to *Betaproteobacteria*, and Kasalický et al34 suggested that *Limnophibitans* had a prominent role in freshwater bacterioplankton communities because of their high rates of substrate uptake and growth, growth on algae-derived substrates, and high mortality rates from bacterivory. The most abundant groups among the eukaryotes were *Maxillopoda*, *Mammalia norank*, *Rotifera norank*, and *Novel_Clade_2 norank* (close relationship with *Cercomonasa*), *Teleaulax*, *Chlamydomonas*, and *Actinocyclos* were also abundant. Such rich algae sequences were due to troubles in these potential water sources largely caused by phytoplankton, and occupied a high biomass in freshwater aquatic systems. This study has identified the key genera involving in abnormal proliferation of algae or off-flavor problems from water source region, also determined their respective roles in aquatic ecological system.

Microorganism community compositions in the estuary23,24 were reported with little knowledge about the effect of environmental factors on the diversity and distribution of microorganism. RDA was powerful27,28 in determining the relationship between microbiological community composition and environmental factors. Previous studies23,24,35 demonstrated that temperature and nutrients were the key factors in the composition of aquatic communities, and that DO, CODMn, and chlorophyll-a were the main direction indices for algae growth. Thus, this study selected six abiotic environmental indices (water temperature, transparency, DO, CODMn, TP, and TN) as well as biotic environmental indices (Chlorophyll-a, MC-RR, and MC-LR concentrations, five predominant sequences). Pyrosequencing studies not only provide an insight into the microbial assemblages in estuary reservoirs but also determine their ecological function through the shifts in their compositions under various time-site treatments. Compared with the estuary reservoirs used in other studies29–31, the one used in this study had acceptable levels of chlorophyll-a and microcystin risk during the entire summer (average 6.41 µg/l, 0.12 of MC-RR, and 0.05 MC-LR µg/l). Environmental conditions were varied among the time-site treatments so that the communities varied as well, thereby increasing the heterogeneity of microbial communities, in accordance with the findings of Yan et al35. Samples from 0 and 2 denoted the inlet and start site of the reservoir, respectively (Fig. 2). Heterogeneity of environmental conditions resulted in the minimum number of *Cyanophyta* and maximum number of *Proteobacteria* in different monitored sampling sites. High abundance of *Proteobacteria* was observed in a previous study32. Heterogeneity was also reported from the similar estuary inflow to the dam37. Microclimate condition was the primary factor
resulting in change of aquatic microorganism diversity for similar water quality between reservoir head region and Changjiang estuary. Therefore, the study was important significance, and researchers should pay more attention to focusing on microbial communities of estuary reservoir. Furthermore, compared with the effect of sampling positions, air temperature changed the microbial community structures more violently for samples were clustered into three parts (Fig. 5).

The pyrosequencing analysis showed that Temperature, MC_RR, TN and DO were key environmental parameters for prokaryotic communities, and Temperature and DO were also the critical factors for eukaryotic communities (Fig. 5). While compared to prokaryotes, majority of the environmental factors for eukaryotes did not explained perfectly for community structure changes. Therefore, roots of changes were derived from the species themselves, really not constrained variables. Transparency, COD_Mn, and TN were shown to be significant factors for prokaryote from late August to mid-October. A strong relationship was also observed among transparency, TN, DO, and the eukaryotic community during winter and spring. Moreover, chlorophyll-a had a close affiliation to both prokaryotic algae in summer and autumn and eukaryotic algae in winter and spring. Transparency also had a strong relationship with algae caused by the increase in algae biomass. Nutrients, such as TN and TP, were shown to be weakly related, and TN had a close relation to both prokaryotic algae in summer and autumn and eukaryotic algae in winter and spring. Data revealed that the TN and TP concentrations had significant discrepancy among the locations (S1). Thus, TN was related to the algae community throughout the year, whereas TP had a lower effect, contrary to the findings of another study conducted on Lake Taihu. TN and TP were also considered to be the major environmental factors at the hypertrophic sampling site35. COD_Mn represented the status of water pollution by organic compounds. This study demonstrated that COD_Mn was associated with the rapid proliferation of Microcystis. The concentration of microcystin (MC-RR and MC-LR) dropped to the lowest level during Microcystis rapid proliferation, which was attributed to the intracellular microcystin that was not released into the water. Low water temperature was accompanied by high DO value, and this study showed that low DO had a weaker relation with Cyanophyta bloom36. Shu et al reported some similar results that the combined shared features of temperature and nutrition conditions led to relatively high abundances of total Microcystis, and other results with minor difference that the concentrations of chlorophyll-a and microcystin were higher than those found in this study. This knowledge is valuable for in-depth understanding of microbial community biodiversity in estuary reservoirs, particularly those that are important potential water sources.

Methods

Description of sampling sites. A total of 80 samples were collected monthly from February to October 2013. The sampling region was located in the Changjiang Estuary and samples were collected according to climate and the four seasons. Spring started on March 4 and the temperature increased from 10°C to 22°C. Summer started on May 5 and the temperature was above 22°C. Autumn started on October 15 and the temperature decreased from 22°C to 10°C. Winter started on November 28 and the temperature was below 10°C. Phytoplankton reproduction rapidly in summer. Six sampling locations were selected for environmental heterogeneity, as shown in Fig. 1. The sampling sites were as follows: (1) Site 0 was the water inlet of the reservoir; (2) Site 2 was situated south of the reclamation area; (3) Site 3 was located north of the reclamation area; (4) Site 8 was at the center of the reservoir; (5) Site 13 was near the water outlet and water intake of the reservoir; (6) Site 35 was the tail of the reservoir. The samples were evaluated by sampling time and sampling station and were marked from 0226-0 (month date-point) to 1029-13 in sequence.

Reservoir characteristics analysis. Water samples from the estuary reservoir were collected using a 500 ml water sampler (WB-PM, Beijing Pujite Instrument Co., Ltd., China). Water temperature was instantaneous value measured using a multi-parameter water quality analyzer (Multi3410, WTW Company, Germany), and air temperature was average value from local meteorological bureau. Water quality indices, including DO, COD_Mn, TN, TP, transparency, and chlorophyll-a, were analyzed using the standard methods37. Microcystin was determined using an Agilent 1200 High Performance Liquid Chromatography (HPLC)38. Microcystin standard substances (MC-LR and MC-RR) were purchased from Sigma, guaranteed reagents were bought from Fisher, other laboratory reagents were obtained from Sinopharm. Freshwater was sampled at a depth of approximately 0.5 m from the top water layer. Microorganisms were collected by filtering 300 ml of the collected water through hydrophilic nucleopore filters (0.22 μm, Jingteng Laboratory Equipment Co., Ltd., China) and then stored at −80°C.

DNA extraction, amplification, and sequencing. Genomic DNA was extracted from the same amount of membrane (0.1 g) on the day after sampling by using the E.Z.N.A. Stool DNA Kit (Omega, USA) according to the manufacturer protocol. As previously described by Bossard et al.39 The V1-V3 region of the 16S rRNA gene was amplified using the forward primer 5’-AGAGTTTGATCCT-GGCTCAG-3’ and reverse primer 533R (5’-TACGGCGGTGTGCTGGCAGC-3’). The V4 region of the 18S rRNA gene was amplified using 3ND1 (5’-GGCAGGTGTG-CCGTCAAG-3’) and reverse primer V4_euk_R2 (5’-AGGTATCTTCTTCTGCG-3’). Polymerase chain reaction was conducted in 20 μl reaction mixtures containing 1 μl of deoxynucleoside triphosphate at a concentration of 2.5 mM, and 0.8 μl of forward and reverse primers, respectively (5 μm each), 0.4 μl FastPlu Polymerase, 5 × FastPlu Buffer 4 μl, and 10 ng of DNA. Each reaction mix received 1 μl of genomic community DNA as a template. The following cycling parameters were used for prokaryotes: retain 2 min at 95°C, 25 cycles (95°C for 30 s, 55°C for 30 s, and 72°C for 30 s), followed by a final extension at 72°C for 5 min, until halted at 10°C by the user. Similar reaction parameters were used for eukaryotes for 35 cycles (95°C for 30 s, 55°C for 30 s, and 72°C for 45 s). Triplicate reaction mixtures were pooled per sample, purified using an AxyPrep DNA gel extraction kit (Axygen, USA) and quantified using the QuantiFluor-ST Fluorescence quantitative system (Promega, USA). The mixtures were then pyrosequenced using a Roche 454 Genome Sequencer FLX Titanium platform (Majorbio Bio-Pharm Technology Co. Ltd., Shanghai, China) as previously described40,41.

Processing of pyrosequencing data. Data were processed using the Quantitative Insights Into Microbial Ecology (Qime) pipeline for 16S rRNA and 18S rRNA data sets (http://qiime.org/tutorials/index.html)42. In brief, the sequences with a quality score below 25 and 200 bp in length were trimmed and then assigned to water samples based on unique 10-bp barcodes. Sequences were binned into operational taxonomic units (OTUs) using a 97% identity threshold, and the most abundant sequence from each OTU was selected as a representative sequence for that OTU. Taxonomy was aligned and compared with the SILVA database (SILVA version 115)43. To compare richness between different treatments, Qime randomly chose a series of subsets of each library in different sizes (100, 200, 300, 400, 500, 600,... with a step of 100) to calculate respective richness indices. Bar plots were done by Microsoft Excel 2010, and heatmap analysis were conducted using the R-gplots for Linux with species-environment correlations44.

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