

Application of Environmental DNA Metabarcoding for Predicting Anthropogenic Pollution in Rivers

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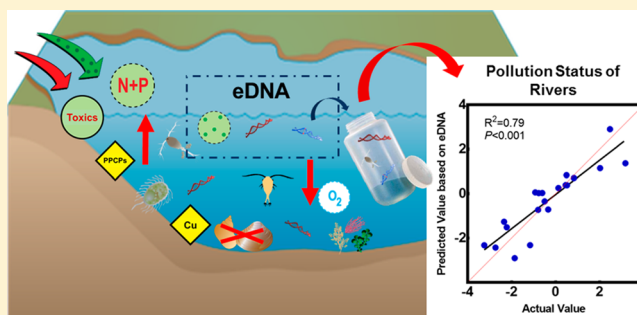
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Supporting Information

ABSTRACT: Rivers are among the most threatened freshwater ecosystems, and anthropogenic activities are affecting both river structures and water quality. While assessing the organisms can provide a comprehensive measure of a river's ecological status, it is limited by the traditional morphotaxonomy-based biomonitoring. Recent advances in environmental DNA (eDNA) metabarcoding allow to identify prokaryotes and eukaryotes in one sequencing run, and could thus allow unprecedented resolution. Whether such eDNA-based data can be used directly to predict the pollution status of rivers as a complementation of environmental data remains unknown. Here we used eDNA metabarcoding to explore the main stressors of rivers along which community structure changes, and to identify the method's potential for predicting pollution status based on eDNA data. We showed that a broad range of taxa in bacterial, protistan, and metazoan communities could be profiled with eDNA. Nutrients were the main driving stressor affecting communities' structure, alpha diversity, and the ecological network. We specifically observed that the relative abundance of indicative OTUs was significantly correlated with nutrient levels. These OTUs data could be used to predict the nutrient status up to 79% accuracy on testing data sets. Thus, our study gives a novel approach to predicting the pollution status of rivers by eDNA data.



INTRODUCTION

Rivers are exposed to multiple stressors, particularly those derived from anthropogenic pollutants, such as excess nutrient, heavy metals, pesticides, or pharmaceuticals.^{1,2} Severe pollution reduces the rivers' provisioning of goods and ecosystem services.³ To alleviate rivers' degradation, and to finally achieve "nontoxic environment" and "good health status" goals, governments implement laws and regulations to manage and improve the water environment.⁴ For example, the European Water Framework Directive (WFD, adopted in 2000) explicitly requires the vast majority of water bodies in member states to reach a "good status" by 2015.⁵ While attempts to monitor chemical contents in waters can directly evaluate the pollution status of rivers, the potential biotoxicity and ecological effects of pollutants can rarely be sufficiently assessed.⁶ Alternatively, biological communities give a comprehensive indication of the physical and chemical properties of rivers, and are both the focus of river protection but can also be used as monitoring targets. Consequently, they

are monitored in the context of applied environmental protection strategies in several countries.⁷ This shift from a focus on chemicals to the focal community to measure quality of waters is widely recognized.^{5,8} However, due to the limitations of the traditional morphology-based species identification approach, river monitoring is extremely time-consuming, labor-intensive, and taxonomic expertise demanding.^{8,9}

Environmental DNA (eDNA) metabarcoding provides a fast and efficient way to uncover biodiversity information, which by now has routinely been used to detect individual species^{10,11} or biological communities in aquatic ecosystem.^{12,13} The eDNA approach has a highly sensitive detection capability and is noninvasive to the organisms themselves,¹⁴ which gives an

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unprecedented opportunity to overcome bottlenecks of traditional morphology-based biomonitoring.¹⁵ Although environmental conditions have been speculated to influence eDNA persistence in aquatic ecosystems,^{16,17} a recent study shows that the decay rate of eDNA can be modeled using first-order constant,¹⁸ and may thus be a rather robust tool. By incorporating eDNA shedding and decay rates, the transport of eDNA can be effectively modeled to estimate species richness in a natural river ecosystem.¹⁹ Comparisons of biodiversity information from eDNA metabarcoding and morphological data sets obtain similar results for freshwater communities.^{9,20} In addition to detecting a set of target taxa, eDNA metabarcoding can also provide access to the broadest set of biodiversity present in the environment.^{12,14} For example, a tree of life metabarcoding or meta-systematics approach has been applied to get a holistic biodiversity perspective at the ecosystem level.¹⁴ This biodiversity revealed by eDNA metabarcoding carries rich information on the local community, but what advantages it carries beyond identifying richness information only is still largely underexplored.

Recently, taxonomy-free molecular indices suggest new measurements for ecosystem assessment using supervised machine learning models.²¹ This method provides a new way to monitor water pollution using high-throughput sequence data. However, the calculation of these taxonomy-free molecular indices still needs information such as taxon-specific ecological weights or categories of tolerance to disturbance.^{21,22} Especially for bacteria or foraminifera that play an important role in ecological processes, these communities with a large proportion of eDNA reads could not be used to calculate indices due to the lack of relevant ecological weights information.²³ Compared with biotic indices, the composition and trophic structure of species in a community may better reflect and capture interactions between the pollution of an ecosystem and the subsequent changes in the ecological network. For example, the abundance of arthropods decreases when pyrethroid is discharged into water, causing algal blooms due to the lack of herbivores.²⁴ Such changes in a river's status can only be understood and predicted by changes in species composition data. Given that eDNA has the advantage of monitoring multiple communities in one sequencing run,¹⁴ it offers a promising tool to assess the species composition of rivers. However, whether such eDNA data can also directly predict the river's pollution status is still insufficiently known.

Here, we used eDNA metabarcoding to profile the species assemblages in rivers from the Yangtze River Delta (YRD), in order to evaluate the method's ability to associate community data with pollution levels. The YRD area is one of the most developed regions in China, and serves as an indispensable water resources for agriculture and industry of 150 million people in eastern China. Large amounts of pollutants discharged into the YRD in recent decades make the study of these rivers a high priority for human welfare. Hence, the main purposes of our study are 3-fold: (1) to profile species assemblages in rivers using eDNA metabarcoding; (2) to explore the main stressors of rivers along which community structure changes, and to rebuild a known stressor gradient based on environmental variables such that it can reveal multiple communities' response under this known stressor gradient; and (3) to predict the pollution status of rivers using eDNA data, and to identify the method's accuracy by comparing testing and training data sets.

MATERIALS AND METHODS

Study Area and eDNA Sampling. Twenty-two sites were sampled from the YRD area during April and May 2016 (Figure S1 of the Supporting Information, SI). These sites are located in tributaries of the lower reach of Yangtze River (5 sites, TYR), the Qinhuai River (7 sites, QHR) and the tributaries of Tai Lake (10 sites, TTL), respectively. Qinhuai River is a tributary of Yangtze River flowing through Nanjing City. Tai Lake is the third largest freshwater lake in China as an indispensable water resources for agriculture and industry products.²⁵ These rivers are exposed to various sources of anthropogenic pollutants.^{26,27} At each site, 10 L of surface water were sampled using sterile bottles (Thermo Fisher Scientific, U.S.A.), and immediately transferred on ice. One liter per site was used for eDNA metabarcoding analysis (which has been shown to be sufficient in many settings),^{14,28,29} the remaining 7 L for chemical analyses (see next section). For the eDNA analysis, filtration was done within less than 6 h after sampling. Four independent extractions of 200–250 mL were made from each 1 L water sample by filtering across a Millipore 0.22 μ m hydrophilic nylon membrane (Merck Millipore, U.S.A.). The total volume of water filtered for each membrane disc depended on the turbidity of water. The membrane discs containing captured eDNA were placed in 5.0 mL centrifugal tubes, were immediately frozen and stored at -20°C until DNA extraction.

Analysis of Environmental Variables. Twenty-two environmental variables were measured for each sampling site. Water temperature (WT), pH, and dissolved oxygen (DO) were measured using YSI water quality analyzer in situ (YSI Incorporated, U.S.A.). For each site, the seven 1-L surface water samples were used to measure basic water quality variables, including permanganate index (COD), total phosphorus (TP), total nitrogen (TN), nitrate (NO_3^-), nitrite (NO_2^-), ammonia nitrogen (NH_4^+), and biochemical oxygen demand (BOD) following standard methods (NEPB, 2002), respectively. For heavy metals, 1 L surface water was diluted with 2% HNO_3 and filtered through a 2.5 μ m membrane filter (Whatman, U.K.). We then determined the concentration of Cr, Mn, Ni, Cu, Zn, As, Cd, and Pb using inductively Coupled Plasma Mass Spectrometry (ICP-MS). For organic chemicals, 1 L surface water was analyzed by a Thermo Ultimate 3000 high performance liquid chromatograph (Thermo Fisher, U.S.A.) coupled to a quadrupole-orbitrap instrument (Thermo QExactive Plus) equipped with a heated electrospray ionization (ESI) source (details shown in SI). These organic chemicals were classified into four major classes, including pesticide, medical drug, industrial processing aid (IPA), and personal care (PerC) components. Detailed information on these chemicals is based on Peng et al.²⁵

DNA Extraction, PCR Amplification, and Next Generation Sequencing. eDNA was extracted directly from the filter membrane discs and blank controls (autoclaved tap water) with a DNeasy PowerWater Kit (Qiagen Canada Inc., ON, Canada) following the manufacturer's protocol. Multiple PCR assays were carried out for the target gene following the previously published protocol.¹³ Briefly, a universal eukaryotic primer pair (1380F: TCCCTGCCHTTTGTACACAC; 1510R: CCTTCYGCAGGTTTCACCTAC) was used to amplify the 130 bp fragment of the hypervariable region of 18S rRNA genes.³⁰ In analogy, the 180 bp fragment of

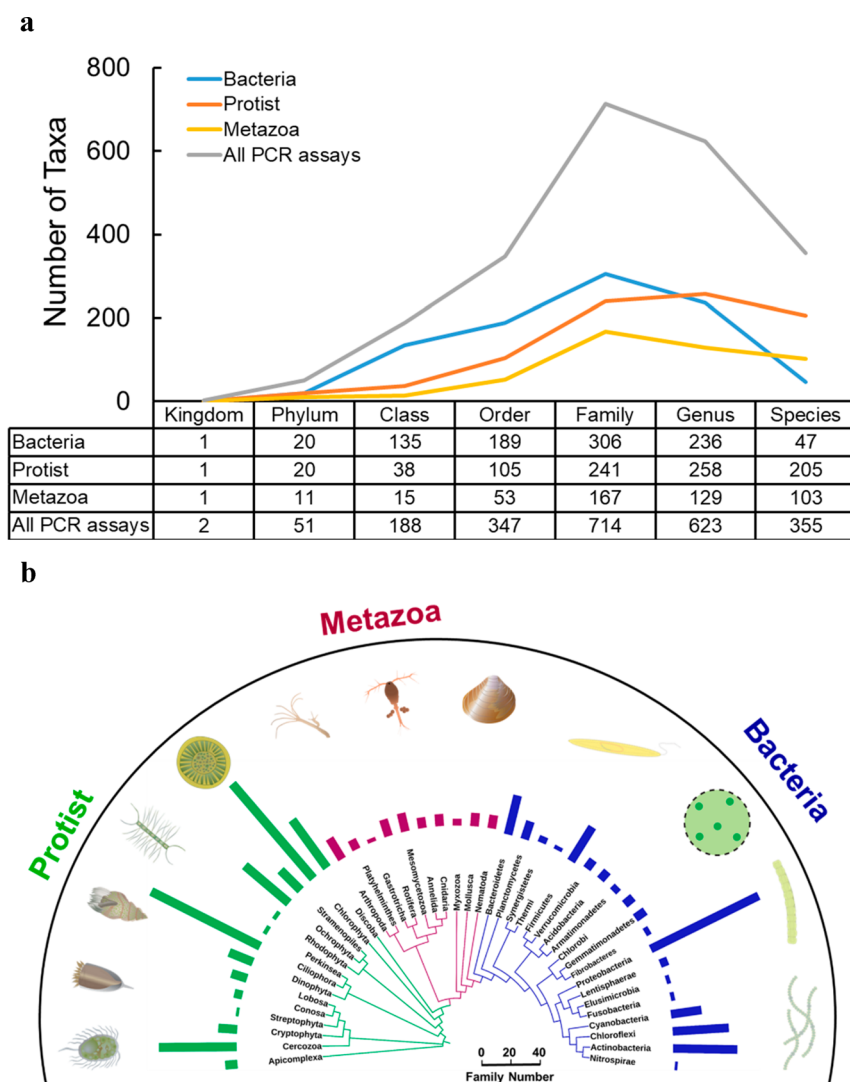


Figure 1. Assignment of eDNA metabarcoding sequences recovered from rivers in this study. (a) Line graph representing the number of each taxonomic rank in different communities, resolved to the highest taxonomic resolution for each OTUs, respectively. (b) Taxonomic phylogenetic tree, built using “tree of life” (ToL) metabarcoding, and bar graphs showing the family numbers per phylum (only families with >3 OTUs are shown).

bacterial 16s rRNA genes was amplified using the modified V3 primer pair (341F: ACCTACGGGSGCWGCAG; 518R: GGTDTTACCGCGGCKGCTG).³¹ To pool and sequence all samples in one sequencing run, unique 12-nt nucleotide codes (also known as tags) were added to the 5'-ends of the forward or reverse primers. All primers were synthesized by Shanghai Generay Biotech Co., Ltd. Each eDNA sample was amplified in three PCR replicates to minimize potential PCR bias, and the products were subsequently combined. PCR negative controls (nuclease-free water as DNA template) were included for all assays. PCRs were carried out in 50 μ L reaction mixture, including 31 μ L of ddH₂O, 10 μ L of 5 \times Phusion Green HF Buffer, 1 μ L of 10 mM dNTPs, 2.5 μ L of each primer (10 μ M), 2.5 μ L of DNA template and 0.5 μ L of Phusion Green Hot Start II High-Fidelity DNA Polymerase (Thermo Fisher Scientific, U.S.A.). The amplification protocol was as follows: initial denaturation at 98 $^{\circ}$ C for 30 s followed by 30 cycles at 98 $^{\circ}$ C for 5 s, 62 $^{\circ}$ C for 30 s, and 72 $^{\circ}$ C for 15 s, with a final extension at 72 $^{\circ}$ C for 7 min, and the PCR was cooled to 4 $^{\circ}$ C until removed. PCR products were visualized on a 2% agarose gel to check the expected size of PCRs yielded

amplicons. Thereafter, the PCR products were purified using the E-Z 96 Cycle Pure Kit (Omega, U.S.A.). All purified PCR products were quantified using Qubit dsDNA HS Assay Kits (Invitrogen, U.S.A.), and were pooled equally for subsequent sequencing. Sequencing adaptors were linked to purified DNA fragments with the Ion Xpress Plus Fragment Library Kit (Thermo Fisher Scientific, U.S.A.) following the manufacturer's protocol. Finally, all samples were diluted to a final concentration of 100 pM. Sequencing templates were prepared with Ion OneTouch 2 and sequenced in the Ion Proton sequencer (Life Technologies, U.S.A.).

Low quality raw sequence (mean quality <20, scanning window = 50, sequences contained ambiguous “N”, homopolymer and sequence length: < 100 bp) were discarded using *split_libraries.py* script in QIIME toolkit.³² The cleaned reads were sorted and distinguished by unique sample tags, all sequences were clustered into OTUs following the UPARSE pipeline at cutoff value of 97% nucleotide similarity. The taxonomy annotation for each OTUs in bacterial, protistan, and metazoan community was assigned against the Greengenes database³³ or the Protist Ribosomal Reference database³⁴

using *align_seqs.py* script, and OTUs number and Shannon entropy index of each community were calculated using *alpha_diversity.py* script in QIIME toolkit.³²

Statistical Analyses. First, eDNA metabarcoding data sets were summarized in separate OTUs table, the taxonomic phylogenetic tree was built using the interactive tree of life (iTOL) online tool.³⁵ Then, to meet the prerequisite of parametric tests, all environmental variables except pH were $\log(x + 1)$ transformed and normalized. To extract the main components explaining the variance of the environmental variables, a principle component analysis (PCA) was performed with the Kaiser-Meyer-Olkin (KMO) and Bartlett's sphericity test, eigenvalues >1 and absolute $r > 0.50$ were taken as criterion for the extraction of the principal components (PC) and the strongly correlation between PC and environmental variables, respectively.³⁶ After, to rebuild a known stressor gradient, all samples were split into three levels (named low, medium and high level) using the one-third of the PC1 distribution as boundaries.³⁷ To detect the difference of environmental variables between each level, nonparametric Kruskal–Wallis (K–W) tests were conducted, followed by post hoc Mann–Whitney–U tests.

To select the significant environmental variables in explaining the variation of bacterial, protistan and metazoan community structure, forward selection distance-based linear models (distLM), based on AIC selection criteria, were used. The significance levels of the variables were assessed by Monte Carlo permutation tests (999 permutations).¹³ To illustrate the variation of communities' structure among three levels, nonmetric multidimensional scaling (nMDS) ordination based on Bray–Curtis (bacteria and protist), and Jaccard (metazoa) dissimilarity matrices were used and the significant differences were assessed by permutational multivariate analyses of variance test (PERMANOVA).³⁸ To identify major OTUs that were responsible for the difference in community structure between each level, a SIMPER analysis was conducted. Using multiple nonlinear regression, we tested the relations between nutrient (surrogated by PC1) and Shannon index of each community. Finally, network analysis was used to explore co-occurrence ecological patterns between OTUs in complex communities, which might be more difficult to detect using either the traditional α - or β -diversity index. Network visualization of the co-occurrence relationships were generated by SparCC with 100 bootstraps to assign P -values.³⁹ Only robust and significant correlations ($|r| > 0.7$ and "two tailed" $P < 0.01$) between nodes were retained in the network.

Indicative OTUs of each level were identified using *multipatt* function in the R package *Indispecies*, the Indicator Values (IndVal) were measured to reflect the conditional probability of the OTUs as an indicator, the significance was tested using a permutation test ($nperm = 999$). To predict the pollution status of rivers based on these indicative OTUs data, predictive models were fitted for three of the four independent subsamples at each site (training data sets) using multivariate linear regression models (MLR) implemented in SPSS 22 software. The reliability and significance of each formula was tested by bootstrap resampling ($n = 1000$). Finally, the remaining subsample at each site were used as testing data sets, to examine the accuracy of predicted values derived from predictive models compared with actual measured values.

All of the above statistical analyses were performed in the R statistical language (<http://www.r-project.org>), GraphPad Prism 6.01 software, SPSS 22 software and PRIMER7 with

PERMANOVA+ add-on software (PRIMER-E Ltd., Ivybridge, U.K.). The network was analyzed and visualized using Cytoscape V3.⁴⁰

RESULTS AND DISCUSSION

eDNA Metabarcoding Provided a Wide Spectrum of Taxonomic Diversity. We detected a total of 1 640 832 bacterial reads, 3 079 304 protistan reads and 362 672 metazoan reads across all samples after stringent quality filtering (Table S1). These eDNA data were assigned to 5850

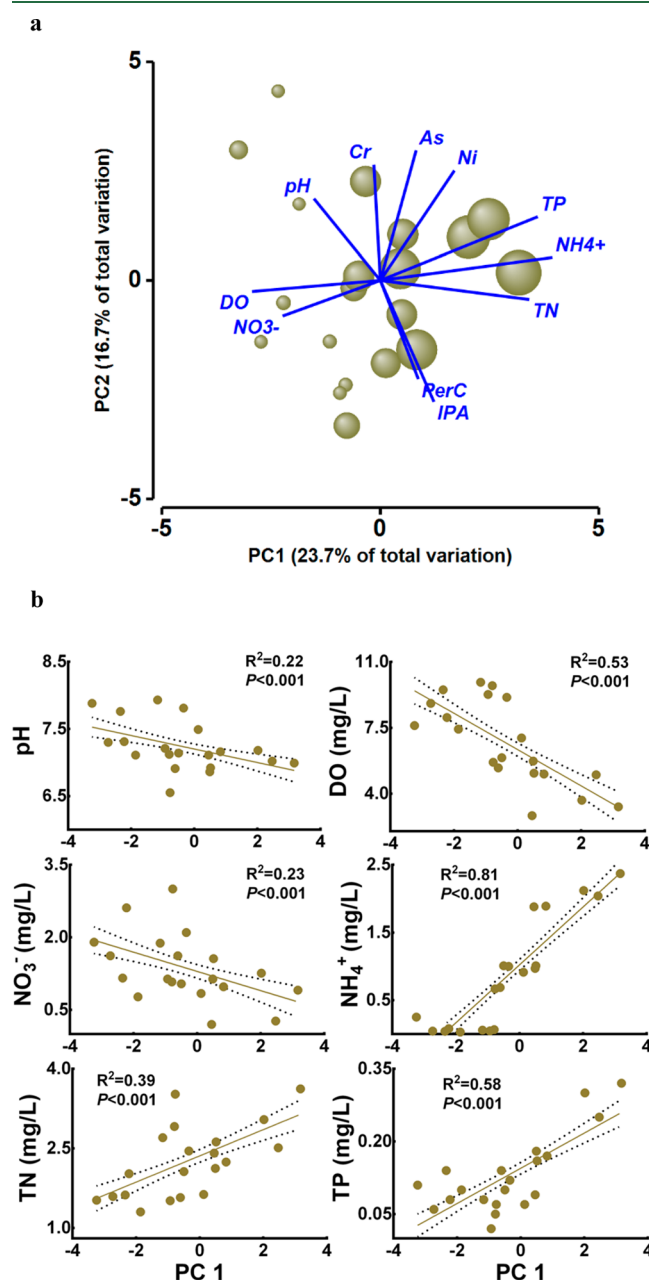


Figure 2. Results of principle component analysis (PCA) on 22 environmental variables (a) and the linear relationship between the first principle components (PC1) and six nutrient-related parameters (b). The bubble size represents the scores of the PC1 in each samples, and the blue line points to the direction of the increase for a given variable, only the strongly correlation (absolute $r > 0.5$) between PC1 and variables are shown (a); the dashed lines are the 95% confidence interval (CI) fitting value (b).

Table 1. Distance-Based Linear Model (distLM) Results of Bacterial, Protistan and Metazoan Community Structures against 22 Environmental Variables in the Full Analysis (9999 Permutations)^a

community	marginal tests			forward selection sequential tests		
	variables	pseudo-F	prop. (%)	pseudo-F	prop. (%)	pumul. (%)
bacteria	pH	6.85	7.54	18.48	18.03	18.03
	NO ₃ ⁻	4.88	5.49	8.42	7.55	25.58
	NH ₄ ⁺	12.78	13.21	8.26	6.81	32.39
	Ni	3.34	3.82	8.59	6.48	38.87
	COD	4.14	4.70	7.40	5.18	44.04
	TN	4.65	5.25	6.28	4.12	48.16
	Cd	9.79	10.44	8.92	5.32	53.48
	TP	4.69	5.29	6.10	3.42	56.90
	Mn	3.67	4.18	6.71	3.50	60.40
protist	NH ₄ ⁺	8.50	9.49	8.50	9.49	9.49
	BOD	7.65	8.63	8.92	9.08	18.58
	NO ₃ ⁻	7.06	8.02	9.90	9.06	27.64
	PerC	6.90	7.85	7.53	6.37	34.01
	TP	5.51	6.37	7.71	6.00	40.01
	Cu	5.52	6.38	6.89	4.99	45.00
	DO	7.23	8.20	8.48	5.59	50.59
	Cd	7.90	8.89	10.11	5.94	56.53
	NO ₂ ⁻	6.08	6.98	11.04	5.71	62.24
metazoa	NH ₄ ⁺	5.07	6.49	5.07	8.49	6.49
	NO ₂ ⁻	3.12	4.10	4.48	5.48	13.97
	Cd	4.69	6.03	6.23	5.11	19.08
	TN	3.98	5.64	4.28	4.66	23.74
	Cr	2.76	3.64	3.37	3.55	27.29
	PerC	2.28	2.72	2.43	2.51	29.80
	TP	3.26	4.27	3.13	3.13	32.93
	drug	2.40	3.18	2.74	2.67	35.61

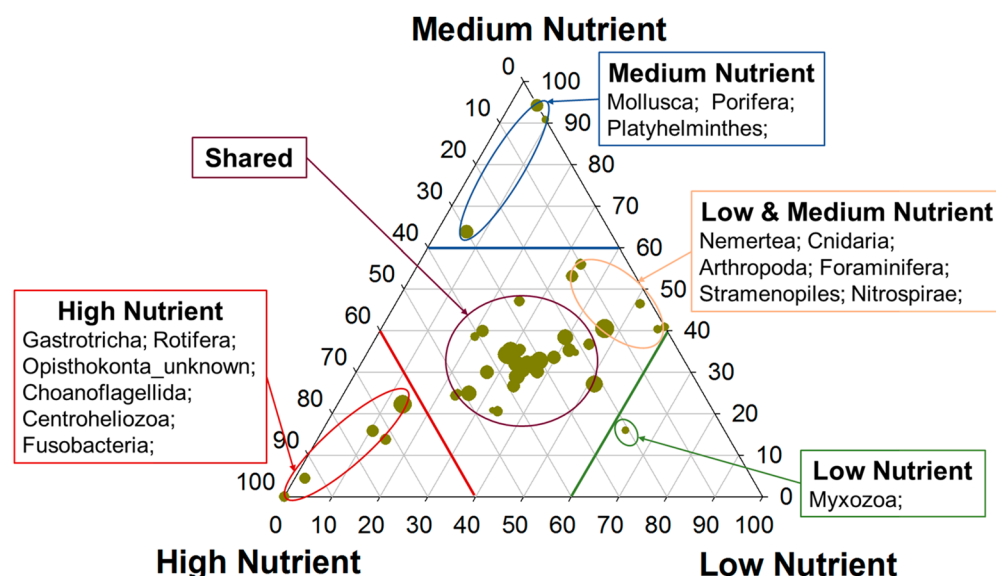
^aProportion of variation explained (Prop. (%)) and cumulative proportion of variation explained (Cumul. (%)) are given. Environmental variables not significantly correlated with community structure ($P < 0.05$) are not shown.

bacterial OTUs, 3475 protistan OTUs and 274 metazoan OTUs (Table S1), annotating (to the highest phylogenetic level resolved) 51 phyla (98.8% of the total OTUs), 188 classes (96.8% OTUs), 347 orders (79.9% OTUs), 714 families (70.9% OTUs), 623 genera (46.9% OTUs), and 355 species (25.9% OTUs) (Figure 1a). The majority of taxonomic lineages at family level belonged to Chlorophyta, Ciliophora, and Proteobacteria (Figure 1b), the relative abundance of these taxa were also disproportionally high (Figure S2). By using different PCR assays, a wide taxonomic lineage including Proteobacteria, Actinobacteria, Chloroflexi, Ciliophora, Chlorophyta, Ochrophyta, Arthropoda, and Mollusca were recovered from the samples. Recent studies have also demonstrated that eDNA methodologies can be used to assess a broad range of prokaryotes and eukaryotes from a variety of environments (e.g., freshwater, seawater, and soil).^{12,14,41} However, there are still some issues (e.g., primer bias, sequencing artifacts and/or contamination) of the eDNA methodologies to be improved,¹⁵ and the taxonomic resolution largely depends on the choice of primer sets and corresponding reference database.⁴² For example, the chloroplastic *rbcL* and nuclear ribosomal 18S genes (e.g., V4 and V9 region) have been used in algae studies,^{42–44} however, there have been controversial in the taxonomic resolution of algae by these primers. These studies suggested that the *rbcL* and V4 region of 18S are more suitable for diatom,^{42,44} yet V9 region of 18S could detect a wide range of taxonomic groups.⁴³ The same issue of primer bias may be found in this study, for example, metazoans have fewer sequences than eukaryotic protists using single V9 region of

18S. Hence, multiple PCR assays using different gene regions are strongly advocated to assess biota in monitoring biodiversity.

Nutrient Identified As a Major Stressor of Rivers. The values of the 22 environmental variables assessed varied largely across samples. Subsequently, all of these variables were reduced to five principal components based on PCA (details were available in SI Table S2). The first two principal components (PC1 and PC2) explained 40.40% of variances of the total variables (Figure 2). Therein, nutrients (including DO, NO₃⁻, NH₄⁺, TN, and TP) were most strongly associated with the first principal component (PC1, Table S2, and Figure 2), which explained 23.70% of the variation in the data. We then used this structuring along the PCA axes as our main environmental descriptors based on which community shifts we wanted to study. Nutrients are relevant in this context, as they have become one of the most severe environmental problems in this region after decades of dense input of nutrients from anthropogenic activities.^{45,46} We used the PC 1 as a new predictor representing nutrient gradients, and then split all samples into three levels (hereafter called “Low nutrient”, “Medium nutrient”, and “High nutrient”) using the 33rd and 66th percentile of the PC1 distribution as boundaries.³⁷ As expected, six nutrient-related parameters were significantly different among these three levels. Specifically, the concentrations of NH₄⁺, TN, and TP in *High nutrient* levels were significantly higher than those in the *Low nutrient* level (Figure S3), while the concentration of NO₃⁻, pH, and DO had an opposing distribution. The other 16 variables (BOD, COD,

a



b

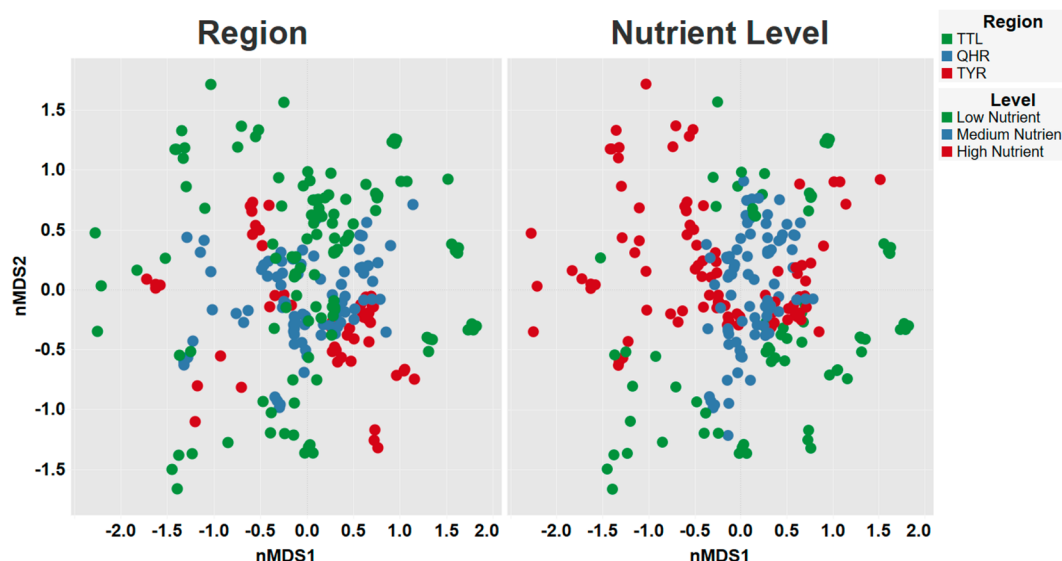


Figure 3. Distribution of dominant taxonomic OTUs at phylum or class level (a) and nonmetric multidimensional scaling (nMDS) analysis of communities' structure (b). The position in the triangle indicates the relative abundance of each phyla or class taxa in bacterial, protistan and metazoan community (a) across three nutrient levels, and the size of the circle represented the relative abundance of each taxon. Significant differences based on Bray–Curtis (bacteria and protist) and Jaccard (metazoa) dissimilarity matrices in the communities' structure are found among the three levels (b). TTL, QHR, and TYR are the abbreviations of the tributaries of Tai Lake, the Qinhuai River, and the tributaries of Yangtze River, respectively.

and heavy metal, and other chemicals) were not significantly different among the three levels.

Changes of Communities' Structure Were Mainly Due to Nutrient. Effects of nutrient on communities' structure were greater than other environment variables. On the basis of distLM analysis, the most parsimonious models explained 60.40%, 62.24%, and 35.61% of the total variation in bacterial, protistan and metazoan communities' structure, respectively (Table 1). In particular, most of the variance in these communities' structure could be explained by nutrient levels (approximately 40%, 30%, and 22%, respectively). These results were coinciding exactly with the PCA of environmental variables indicating that nutrients were the driving stressor of these rivers. Hence, the pollution status of rivers may be

directly revealed by such species information, and may not need to have environmental variables being monitoring.⁸ Importantly, the biological assessment gives an integrated measure of the nutrient exposition of a community over time, while chemical measurements usually only cover one specific time point. To verify this hypothesis, we then analyzed the trend of communities' change under known stressor gradients (nutrient gradients), which is expected to identify some taxa for characterizing the nutrient status in rivers.

First, we found that the dominant taxa in bacterial, protistan, and metazoan communities varied across the nutrient gradients (Figure 3a). Some taxa (Myxozoa, Nitrospirae, Foraminifera and Stramenopiles, Mollusca, and Arthropoda) were primarily identified in *Low* and/or *Medium* nutrient level, and taxa in

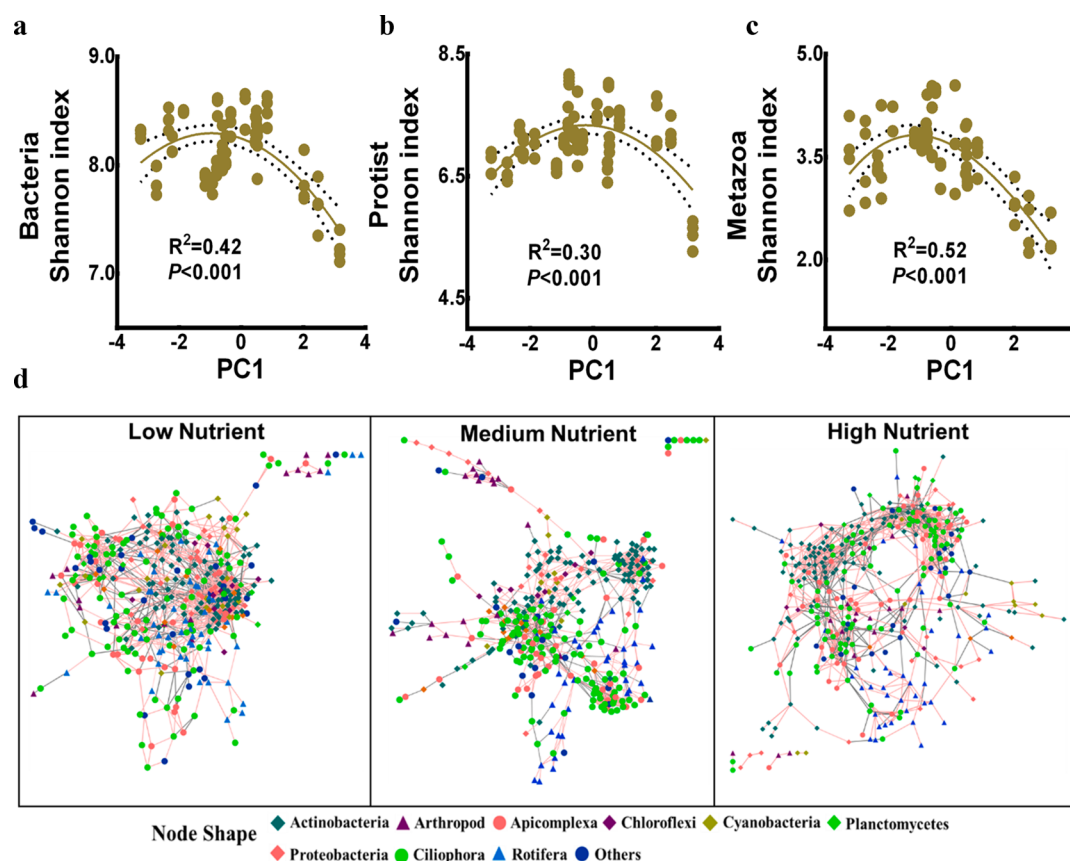


Figure 4. Responses of Shannon index to nutrient (surrogated by PC1, a–c), and ecological interaction network between bacterial, protistan and metazoan communities in each nutrient level (d). Nonlinear polynomial regression included 95% CI (the dash lines) in bacteria (quadratic), protistan (quadratic), and metazoan (cubic) communities. The correlations between each OTUs were generated by SparCC with 100 bootstraps to assign P -values. Only when the absolute $r > 0.7$ and a “two tailed” P value < 0.01 , the nodes and edges in the network remained.

Opisthokonta_unknown, Choanoflagellida, Centroheliozoa, Gastrotricha, and Rotifera were dominant in *High nutrient* level. These results were consistent with other studies showing that eutrophication altered the composition of communities.^{47,48} For example, Mollusca and Foraminifera were sensitive to nutrient,^{49,50} Rotifers increased along a gradient of increasing nutrient levels.⁵¹ Besides, as a chemolithoautotrophic nitrite-oxidizing bacterium,⁵² Nitrospirae was primarily found in *Low* and *Medium nutrient* level, which might explain why the concentration of NO_3^- in these two levels were greater.

Then, the communities' structure also largely varied following the nutrient gradient rather than across regions (Figure 3b), which were further identified as significant by the PERMANOVA tests (pseudo- $F_{\text{bacteria}} = 5.187$, $P < 0.001$; pseudo- $F_{\text{protist}} = 7.854$, $P < 0.001$; pseudo- $F_{\text{metazoa}} = 9.188$, $P < 0.001$). The SIMPER analyses revealed that *Microcystis* sp. (Cyanobacteria, OTU625), *Mycobacterium* sp. (Actinobacteria, OTU368), and ACK.M1 (Actinobacteria, OTU280) in bacterial communities were the major contributors to the dissimilarity across each level (Table S3). *Mediophyceae* sp. (Bacillariophyta, OTU726), *Cryptomonas* sp. (Cryptophyta, OTU544), and *Strobiliidiidae* sp. (Ciliophora, OTU104) in protozoan communities and *Sinocalanus* sp. (Arthropoda, OTU48), *Leiosolenus* sp. (Mollusca, OTU87), and *Brachionus calyciflorus* (Rotifera, OTU103) in metazoan communities were the main contributors to the dissimilarity. As is well-known, *Microcystis* became a dominant taxon during

cyanobacterial bloom periods⁵³ and could produce highly stable and potent polypeptides (microcystins (MCs)) that pose a serious threat to public health.^{46,54} Nutrient enrichment combined with high ambient temperature was regarded as the main stressor that influenced on *Microcystis* blooms.⁵⁵ Besides, some laboratory toxicity studies found that some species in *Brachionus* could be more tolerant of ammonia than cladocerans and copepods.^{56,57}

Alpha Biodiversity and Ecological Interaction Network Varied with Nutrient Status. Alpha biodiversity (Shannon index) indicated a significantly hump-shaped response to nutrient in bacterial, protistan and metazoan communities (Figure 4a–c). Our results contribute to growing evidence of nonlinear responses of aquatic assemblages to nutrient enrichment.^{58,59} Although linear responses of biodiversity to nutrient enrichment were also reported,⁶⁰ some factors might frequently obscure natural nonlinear responses of multiple taxa to stressors. For example, biotic interaction could change structure of the food web within an ecosystem,⁶¹ so that the consumer–resource interactions in communities were often affected by other species.^{62,63} The top predator species could determine how communities' conditions changed across time and space.⁶⁴ The network analysis verifies the above inference, such that the ecological interaction network between OTUs in each community revealed a distinct network topology in each nutrient level (Figure 4d). The number of nodes and edges in network were higher in *Low nutrient*, followed by *High* and *Medium nutrient* (Table S4). In

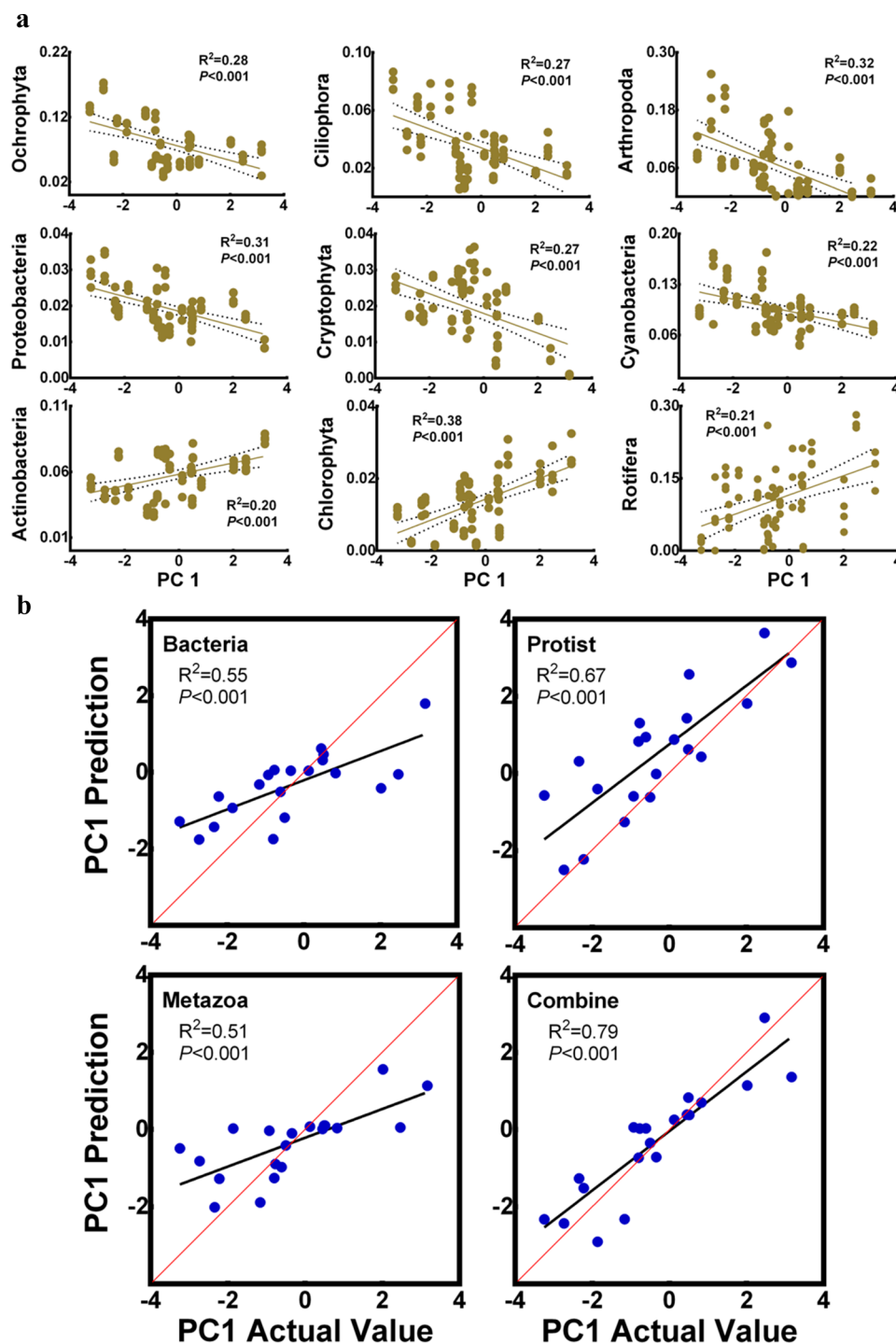


Figure 5. Relationship between nutrient (surrogated by PC1) and the relative abundance of indicative OTUs, the dashed lines are the 95% CI fitting value, only significant correlation ($P < 0.05$) are shown (a). Comparison between the PC1 predicted value given by the indicative OTUs in test samples and the PC1 actual value derived from environmental variables, the red diagonal lines represent the ratio (1:1) between the predicted and measured values (b).

addition, more complex ecological interactions in *Low* and *High nutrient* level according to the betweenness centrality and average closeness centrality parameters.⁶⁵

Novel OTUs Based Indicator Could Rapidly Predict Nutrient Status of Rivers. Indicator analysis identified 960 OTUs in different taxa (i.e., Ochrophyta, Ciliophora, Actinobacteria, Proteobacteria and Chlorophyta) that were characteristic of each nutrient level (Figure S4). These OTUs included their trophic positioning and responsiveness to a range of

nutrient gradients.^{42,66} In addition, the relative abundance of indicative OTUs in Ochrophyta, Ciliophora, Arthropod, Proteobacteria, Cryptophyta and Cyanobacteria were significantly negatively correlated with nutrient, indicative OTUs in Actinobacteria, Chlorophyta, and Rotifera could increase with nutrient gradient (Figure 5a). Here it is noteworthy that the relationship between the relative abundance of indicative OTUs in cyanobacteria and nutrient was contradictory to previous study.⁵⁵ One possible explanation is that more than

Table 2. On the Basis of Multivariate Linear Regression Models (MLR) to Predict Nutrient Status (as Dependent Variable, y) of Rivers Using the Relative Abundance of Indicative OTUs Data (% Indicative OTUs, as Independent Variable, x) in Training Data Sets^a

predictor variables	predictor formula	Adj- R^2	F
% indicative OTUs (bacteria)	$y = 1.61 - 98.33 \times \% \text{ Proteobacteria}$	0.41	16.41
% indicative OTUs (protist)	$y = 3.91 + 27.86 \times \% \text{ Stramenopiles} - 30.93 \times \% \text{ Ciliophora} - 28.36 \times \% \text{ Chlorophyta} - 68.52 \times \% \text{ Cryptophyta}$	0.60	12.94
% indicative OTUs (metazoa)	$y = 0.12 - 6.68 \times \% \text{ Arthropoda}$	0.43	7.13
% indicative OTUs (combine)	$y = 3.45 - 48.46 \times \% \text{ Proteobacteria} + 39.33 \times \% \text{ Stramenopiles} - 41.02 \times \% \text{ Ciliophora} - 7.84 \times \% \text{ Arthropoda} - 358.39 \times \% \text{ Bacteroidetes} - 10.24 \times \% \text{ Ochrophyta}$	0.75	12.63

^aCombine, integration of bacteria, protistan, and metazoan communities; Adj- R^2 , the adjust R^2 .

half of sites had reached eutrophication or severe eutrophication status (Table S2). A previous study within this study area confirmed this phenomenon that high cyanobacteria cell concentrations were negatively correlated with ammonia, especially in appropriate external water temperature.⁶⁷ Recently, palaeo-limnological views based on subfossil DNA also supported strong correlations between trophic status changes and microbial eukaryotes community succession.⁶⁸ By contrast, the largest proportions of these indicators were Ochrophyta, particularly more than 50% of OTUs in Ochrophyta belonging to *Cyclotella* sp., *Nitzschia* sp., *Melosira* sp., or others. (Bacillariophyceae). Diatoms inhabit a variety of waterbodies and different species could respond differently and characteristically to environmental status.⁶⁹ In particular, the *Nitzschia* sp. has been used to characterize the nutrient status in waters.⁷⁰ Besides, diatoms were considered as the most sensitive group to TP, and the occurrence of OTUs declined with increasing concentrations of TP.³⁸

Using identified indicative OTUs, we could identify the nutrient status with 41%–75% accuracy on training data sets, but the predictive ability of single communities was lower than combined communities' data (Table 2). When comparing the nutrient (surrogated by PC1) predicted value with the actual value using test data sets, there was a good consistency between each value ($R^2 = 0.51$ – 0.79 , Figure 5b). However, we also found underprediction or overprediction of the nutrient status using single community data. For example, the predicted value in *Low nutrient* level was higher than actual ones using bacterial or metazoan data, but the lower predicted value was occurred in *High nutrient* level. Besides, almost all predictions were higher than the actual ones in protistan data. Corresponding to single community data, we achieved up to 79% accuracy to predict the nutrient status via combining multiple communities' data. A recent study has also demonstrated that multitrophic metabarcoding biotic index has higher predictive potential for pollutant status.²³ In another study, it has also been found that sequence data was a better predictor than environmental variables to predict cyanobacterial blooms.²⁸ In short, eDNA metabarcoding offers novel and promising tools to monitor and predict anthropogenic contamination of aquatic ecosystems (e.g., rivers, lakes, marine ecosystems) by DNA sequence-based data.

■ ASSOCIATED CONTENT

Supporting Information

The Supporting Information is available free of charge on the ACS Publications website at DOI: 10.1021/acs.est.8b03869.

Detailed information regarding the method for the detection of organic chemical and principal component

analysis of environmental variables; figures showing location of sampling sites, the relative abundance of taxa at phylum level, comparison of environmental variables, and the number of indicative OTUs; tables showing sequencing counts and shannon entropy index, summary of environmental variables, and the results of PCA, SIMPER analysis, and parameters of network topology (PDF)

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Notes

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