Insight into the nitrogen accumulation in urban center river from functional genes and bacterial community

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Abstract

Along with urbanization, the intensified nitrogen pollution in urban rivers and the form of black-odor rivers has become one of the biggest concerns. Better understanding of the nitrogen transformations and microbial mechanisms occurring within urban rivers could help to manage their water quality. In this study, pollution characteristics, potential nitrogen removal rate, composition and function of bacterial community, and abundance of functional genes associated with nitrogen transformation were comparatively investigated in a typical urban river (FC) and a suburban river (LH). Compared with LH, FC was characterized by higher content of nutrients, lower potential nitrogen removal rate and lower abundance of functional genes associated with nitrogen transformation in both overlying water and sediment, especially in summer. Sediment dissolved organic matter characterized by excitation-emission matrix (EEM) showed that FC was more severely polluted by high nitrogen organic matter. Our results revealed that anammox was the main nitrogen removal pathway in both rivers and potential nitrogen removal rates decreased significantly in summer. Bacterial community analysis showed that the benthic communities were more severely influenced by the pollutant than aquatic ones in both rivers. Furthermore, the FC benthic community was dominated by anaerobic respiring, fermentative, sulfate reduction bacteria. Quantitatively, the denitrification rate showed a significant positive correlation with the abundance of denitrification genes, whilst the anammox rate was significantly negatively correlated with bacterial diversity. Meanwhile, NH$_4^+$-N had a significant negative correlation to both denitrification and anammox in sediment. Taken together, the results indicated that the increased nitrogen pollutants in an urban river altered nitrogen removal pathways and bacterial communities, which could in turn exacerbate the nitrogen pollution to this river.

Introduction

Fast growth of population and urbanization cause urban areas to produce more and more pollution. Although improved wastewater collection and treatment can efficiently control point
source pollution, nonpoint source pollution, such as surface runoff caused by rainfall events, as well as endogenous release of the sediment pollutants persist to increase the levels of pollution in urban rivers [1–4]. Usually, urban rivers are characterized by high levels of nutrient inputs associated with human activities [1, 5]. More seriously, many rivers heavily polluted by organic matter, nitrogen, phosphorus, and heavy metals often resulted in excessive oxygen consumption and production of blackening and stinking pollutants, which finally caused rivers to become black and odor smell, especially in summer [6–8]. Nowadays, urban water quality degradation had been observed in developed and developing countries [6, 9]. Stress to local water resources had become one of the biggest concerns associated with urbanization [1, 10]. Therefore, improving urban river water quality, especially elimination of black-odor phenomenon in rivers, is an urgent need for sustainable city development.

High loading of nitrogen (N) is considered as one of the main causes of black-odor phenomenon in rivers [1, 2, 6, 8]. Self-purification function of riverine ecosystems can removal N pollutants through microbial driven processes of ammonification, nitrification, denitrification and anoxic ammonium oxidation (anammox) [11]. However, N removal processes are influenced by variation of environmental factors, contaminants and microbial communities [11–15]. Water quality also deeply influences river ecosystem structure, food-web integrity, as well as ecosystem functions and services [16]. Previous studies have demonstrated that flow velocity and contaminants in urban rivers affected sediment oxygen demand, and the structure and integrity of eukaryotic and bacterial communities [14, 17, 18]. The affected microbial communities in turn disrupted microbial driven self-purification processes, which might aggravate the accumulation of pollution and water quality degradation [8, 19]. For example, effluent inputs and seasonal changes determined the abundance and distribution of nitrifiers, and subsequent transformation of NH$_4^+$-N to nitrate in urban rivers [20]. Till now, many studies about the deterioration of water quality in urban rivers focused on the influent pollution, as well as mechanisms underlying the phenomenon of black and odor in urban rivers [14, 15, 20, 21]. It is pressing to investigate the characteristics of N accumulation and removal in urban rivers and underlying microbial mechanisms to take steps to improve water quality and protect urban aquatic ecosystems.

Wuxi city, adjacent to Lake Taihu, has witnessed degradation of urban river water quality along with fast urbanization. FengChan river (FC) is located in the city center of Wuxi and the FC subwatershed is dominated by urban land cover. LiangHong river (LH) is situated in a wetland protection area within suburb of Wuxi city and is isolated from direct urban impact. FC and LH provide an opportunity to distinguish the influence of the characteristics of urban and suburban pollution into the river, as well as their effects on microbial community structure and self-purification function in rivers. In this study, we compared the biogeochemical properties, bacterial community composition, functional gene abundance, and potential N removal processes in FC and LH. And we discussed the factors that influence the potential N removal processes. The results would help to systematically understand the influence of urbanization on river N pollution and removal at the level of community, functional gene and bacterial denitrification capacity. The results will provide scientific clues to manage and restore urban river water quality.

**Materials and methods**

**Study sites and sampling**

In this study, water and sediment samples were collected from an urban river (31°36 N, 120’18 E; named FC in this study) located in the city center of Wuxi and a river located in the LiangHong wetland protection area (31°30 N, 120’31 E; named LH in this study). The two rivers
have similar temperature and climate environment. Water and surface sediment samples were collected in April and July, 2018. Three sampling sites were selected for each river with intervals of 500 m. Water samples were collected from the surface layer of the two rivers. The water samples were stored in a sealed polyethylene bucket after large particles of solids and plankton were removed with a net (mesh size of 10 µm). Sediment samples were collected using a beaker-type sampler. Water and sediment samples were stored at 4°C and transferred to the laboratory immediately for subsequent analysis. All water bodies and the riparian areas are public ground. We confirm that no permission was required for the water and sediment sampling.

**Analytical procedures for water and sediments characteristics**

HQ30d Hach portable meter (Hach, USA) was used to measure dissolved oxygen (DO) on sites. Total phosphorus (TP), total nitrogen (TN), ammonium nitrogen (NH₄⁺-N), nitrate nitrogen (NO₃⁻-N), and chemical oxygen demand (COD) were analyzed according to Methods for the Analysis of Water and Wastewater [22]. TN, TP, NH₄⁺-N and NO₃⁻-N were measured by UV-Visible spectrophotometer (UV2450, Shimadzu, Japan). Total organic carbon (TOC) content was determined by TOC analyser (vario TOC, elementar, Germany). Sediment TP, TN, TOC, NH₄⁺-N, and NO₃⁻-N concentrations were measured after extraction with 2 M KCl (1:5 wt/vol). All samples ran in triplicate.

**Spectroscopic characterization of sediment dissolved organic materials**

Dissolved organic materials (DOM) were extracted by mixing ground and sieved freeze-dried sediment (100 mesh sieves) and Milli-Q water in a 1:10 ratio, shaking 24 h at 200 rpm, 25°C, then centrifuged at 8000 rpm for 5 min. The supernatant was filtered through 0.45 µm membrane to get DOM.

Three-dimensional excitation–emission matrix (3D-EEM) fluorescence spectra was analyzed using Horiba F-7000 (Hitachi, Japan) fluorescence spectrophotometer with an excitation (Ex) range from 200 to 450 nm and an emission (Em) range from 250 to 550 nm. The spectra were recorded at a scan rate of 12,000 nm/min, using excitation and emission slit bandwidths of 5 nm. Rayleigh scattering was subtracted from the original EEM data. Fluorescence spectra of Milli-Q water was run under identical conditions to eliminate the effect of Raman scattering. The data of fluorescence spectra were plotted by OrginPro 2017.

Six components were analyzed through 3D-EEM fluorescence spectra. C1 (Ex / Em ≤ 230 (285) / 340 nm) represents amino acid associated tryptophan-like components. C2 (Ex / Em ≤ 240 (350) / 468 nm) represents a typical terrestrial humic-like substance. C3 (Ex / Em ≤ 230 / 420 nm) represents agricultural-soil-derived humic-like or fulvic-like materials. C4 (Ex / Em ≤ 230 (275) / 316 nm) and C6 (Ex / Em ≤ 230 (270) / ≤ 300 nm) represent those of redshifted tyrosine and typical tyrosine substances, respectively [23].

**Potential rates of denitrification and anammox**

The potential rates of denitrification and anammox were measured by incubation experiments using the ¹⁵N isotope pairing technique and the Membrane Inlet Mass Spectrometer (MIMS) (Bay Instruments, Easton, MD) determination of ²⁹N₂ and ³⁰N₂ in the sediment slurry and overlying water [24]. Fresh sediments were mixed with helium (He)-purged water at a ratio (sediment/water) of 1:5 in 12 mL glass vials (Exetainer, Labco, U.K.) to get sediment slurries. Sediment slurry and overlying water were then incubated for 3 days at 20°C, 150 rpm to eliminate nitrate and oxygen. After that, the vials were spiked with 100 µL He-purged solution of ¹⁵NO₃⁻ (99.2% ¹⁵N) to a final concentration of 100 µM ¹⁵N, respectively. Sediment slurry and
overlying water were incubated at 20˚C for 36 h and stopped by adding 200 μL saturated ZnCl₂ solution, then the concentrations of ²⁹N₂ and ³⁰N₂ were determined by MIMS.

Both anammox and denitrification generated ²⁹N₂. Thus, the respective contributions of each process to the total ²⁹N₂ production were quantified by Eq (1):

\[ P_{29} = A_{29} + D_{29} \] (1)

where, \( P_{29} \), \( A_{29} \), and \( D_{29} \) represent the production rate of total ²⁹N₂, ²⁹N₂ from anammox, and ²⁹N₂ from denitrification, respectively. Because the ²⁸N₂, ²⁹N₂, and ³⁰N₂ generated from denitrification follow random isotope pairing, \( D_{29} \) can also be estimated by Eq (2):

\[ D_{29} = P_{30} \times 2 \times (1 - F_n) \times F_n^{-1} \] (2)

where \( P_{30} \) is the total ³⁰N₂ production rate, and \( F_n \) is the mole fraction of ¹⁵N in the nitrate pool. Consequently, the potential rates of anammox and denitrification were estimated by the following equations:

\[ D_{\text{total}} = D_{29} + 2 \times D_{30} \] (3)

\[ A_{29} = P_{29} - D_{29} \] (4)

where \( D_{\text{total}} \) and \( A_{29} \) are the potential rates of denitrification and anammox, respectively.

**DNA extraction and quantitative PCR assay**

DNA was extracted from freeze-dried sediment (approximately 0.5 g) or 200 mL overlying water filtered through 0.22 μm filter using a FastDNA SPIN Kit for Soil (MP Biomedicals, Solon, OH, USA) according to the manufacturer’s protocol. The DNA concentration and quantity were determined using a NanoDrop 2000 spectrophotometer (Thermo Fisher Scientific, Schwerte, Germany).

Quantitative PCR (qPCR) assays were conducted using the SYBR-Green approach on ABI 7500 Real-Time PCR System (Applied Biosystems, Foster City, USA). The abundances of anammox bacteria, nitrifier (AOB), denitrifier, and total bacteria were quantified targeting the corresponding specific genes. Details about the primer sets, thermal profiles, and experimental procedures are provided in S1 Table.

**16S rRNA gene high-throughput sequencing and data analysis**

The V3-V4 hypervariable region of sample DNA were amplified using 341F/806R primer pair (341F: 5’-CCTAYGGGRBGCASCAG-3’, 806R: 5’-GGACTACNNGGGTATCTAAT-3’) and sequenced using PE250 (Illumina, CA). The sequence analysis was carried out using QIIME (version 1.9.1). The OTUs were assigned to a set of hierarchical taxa using the ribosomal database project (RDP) classifier. The α-diversity metrics, including Chao 1, OTU richness, Shannon and Simpson index were calculated to compare the bacterial richness and evenness among different treatments. Principal coordinates analysis (PCoA) of Weighted-Unifrac distance and non-metric multidimensional scaling (NMDS) analysis of Bray-Curtis distance were carried out using R (version 3.5.1) package Vegan.

The bacterial taxa (e.g., genera or species) were classified to different functional groups using the functional annotation of prokaryotic taxa (FAPROTAX) database (http://www.zoology.ubc.ca/louca/FAPROTAX/). The Phylogenetic Investigation of Communities by Reconstruction of Unobserved States (PICRUSt) was used to predict the function of bacterial community [25].
The linear discriminant analysis (LDA) effect size algorithm (LEfSe) was applied to explore the statistically significantly features of microbial communities between different samples. An alpha = 0.05 was used in Wilcoxon rank sum test, and threshold for the LDA analysis was set to be 4.0 [26].

Statistical analysis
Statistical analysis of significant difference was analyzed by Graphpad prism 7.0 software with two-way analysis of variance (ANOVA). The level of p < 0.05 was considered statistically significant. Data were expressed as mean ± SD. Spearman correlation analysis was performed to evaluate the correlations among the potential N removal rates, relative abundances of the functional genes, and environmental factors using the SPSS 19.0 package.

Results
Water quality and sediment property in urban river
High concentrations of TN (4.51–8.79 mg L\(^{-1}\)), TP (0.22–0.27 mg L\(^{-1}\)), \(\text{NH}_4^+\)-N (2.47–3.06 mg L\(^{-1}\)) and COD (27.55–29.15 mg L\(^{-1}\)) were observed in FCW (FC overlying water). In LHW (LH overlying water), the concentrations of TN, TP, \(\text{NH}_4^+\)-N and COD were 2.68–7.61 mg L\(^{-1}\), 0.05–0.21 mg L\(^{-1}\), 0.50–1.86 mg L\(^{-1}\) and 23.47–36.09 mg L\(^{-1}\), respectively. In the overlying water, the concentrations of TN, TP, and \(\text{NH}_4^+\)-N in FCW were 1.15–1.68 times, 1.31–4.64 times and 1.33–6.11 times of their counterparts in LHW in spring and summer, respectively, whilst \(\text{NO}_3^-\)-N acted oppositely (Fig 1). In both rivers, the concentrations of TN, TP, and \(\text{NO}_3^-\)-N increased sharply in summer, however, \(\text{NH}_4^+\)-N in FCW decreased in summer (Fig 1).

In sediments, the contents of TN, TP, \(\text{NH}_4^+\)-N, and \(\text{NO}_3^-\)-N in FCS (FC sediment) were also significantly higher than LHS (LH sediment) (p < 0.05) (Fig 2), indicating that more pollutants accumulated in urban center river sediment compared with the suburban one. Despite the significant spatial variability of nutrient contents between urban and suburban rivers, both rivers showed obvious seasonal characteristics. In LHS, the contents of TN, TP and TOC in summer were significantly higher than spring (p < 0.05), which was consistent with the higher load as observed in overlying water in summer. In FCS, the content of TN stayed stable from spring to summer, whereas the contents of \(\text{NH}_4^+\)-N and \(\text{NO}_3^-\)-N in FCS were 3.25 and 1.24 times of their spring level (Fig 2). Furthermore, TP content in FCS decreased in summer.

3D-EEM of sediment DOM showed that tryptophan-like (C1) substance in FCS (22.33% in spring, 25.93% in summer) was higher than that in LHS (15.51% in spring, 10.07% in summer). Furthermore, the data also showed that tryptophan-like (C1) substance in FCS significantly increased from spring to summer (Table 1), indicating that more partially-degraded organic matter accumulated in FCS than LHS, especially in summer. Consistent with the increase of tryptophan-like substance, the ratio of tryptophan to tyrosine increased from 5.58 to 6.57 in FCS from spring to summer (S2 Table), suggesting that more fresh protein-like organics accumulated in summer in FCS. In contrast to FCS, LHS had higher proportion of humic- or fluvic-like substance (C3), as well as microbial humic-like substance (C4) (Table 1). Both C3 and C4 components in LHS increased from spring to summer, which demonstrated that more recalcitrant DOM accumulated in LHS, especially in summer.

Functional genes associated with potential nitrogen removal
The abundances of functional genes related to potential N removal, including nitrification (amoA, nxr), denitrification (narG, napA, nirS, nirK, norB, and nosZ), and anammox 16S rRNA genes were analyzed using qPCR. In overlying water, amoA abundance in FCW was
significantly lower than LHW ($p < 0.05$). While the abundance of $amoA$ in LHW significantly decreased from spring to summer, $amoA$ abundance in FCW increased from spring to summer, coinciding with the elevated DO concentration in summer FCW (Fig 1). Abundance of anammox and $nxr$ genes significantly increased from spring to summer in both FCW and LHW, and the highest levels of anammox (7.62–7.75 log gene copies mL$^{-1}$, 0.11% of total bacteria) and $nxr$ (7.30–7.37 log gene copies mL$^{-1}$, 0.05% of total bacteria) were detected in summer FCW (Fig 3). In overlying water, the abundance of all denitrification genes in spring were higher than summer in both rivers, except for $nirK$ in FCW.

Fig 1. The different water quality index between FC and LH in different seasons. (A) DO, (B) TP, (C) TN, (D) NH$_4^+$-N, (E) NO$_3^-$-N, (F) COD.

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Fig 2. The different sediment component content between FC and LH in different seasons, (A) TN, (B) NH$_4^+$-N, (C) TP, (D) NO$_3^-$-N, (E) TOC.

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In sediments, the abundance of amoA, anammox and denitrification genes was higher than overlying water (Figs 3 and 4), demonstrating that sediment was a hotspot of N metabolisms and removal. Compared with LHS, the abundance of amoA, anammox and denitrification genes were significantly lower in FCS (Figs 3 and 4). Just as that observed in overlying water, the abundance of denitrification genes significantly decreased in summer in both rivers (Fig 4).

**Characteristics of bacterial communities in urban river**

Overall, the α-diversity characterized by Chao 1 and Shannon H indices showed that sediment bacterial communities was more diverse than their counterpart in overlying water (Table 2). This result was not surprising since the microenvironment in sediment was more heterogeneous than overlying water and niche selection played an important role in shaping bacterial diversity. In sediment, the α-diversity of LHS in summer was higher than spring, however, FCS had a rather stable α-diversity in both seasons. No matter in overlying water or sediment, the α-diversity of bacterial communities in FC were lower than in LH. In agreement with our results, a previous study showed that the food-web of urban river was low in diversity due to the greater influence of water quality degradation caused by human activities [16].

Beta-diversity (NMDS and PCoA) clearly revealed that bacterial communities in overlying water were separated from the benthic ones (Fig 5). Bacterial communities in overlying water

![Table 1. Six componen ts concentratio n (RU) through 3D-EEM fluorescenc e spectra in FCS and LHS.](https://doi.org/10.1371/journal.pone.0238531.t001)

| Substances | FC-spring | Proportion | FC-summer | Proportion | LH-spring | Proportion | LH-summer | Proportion |
|------------|-----------|------------|------------|------------|------------|------------|------------|------------|
| C1 amino acid associated tryptophan-like components | 1.8925 | 22.33% | 2.623 | 25.93% | 0.929 | 15.51% | 0.767 | 10.07% |
| C2 terrestrial humic-like substance | 4.5253 | 53.39% | 3.814 | 43.71% | 3.470 | 54.22% | 3.388 | 44.48% |
| C3 agricultural-soil-derived humic-like or fulvic-like materials | 0.5071 | 5.98% | 0.789 | 9.05% | 0.525 | 8.21% | 1.072 | 14.08% |
| C4 microbial humic-like substance | 0.3379 | 3.99% | 0.622 | 7.14% | 0.452 | 7.07% | 1.060 | 13.93% |
| C5 redshifted tyrosine | 0.8732 | 10.30% | 0.892 | 10.23% | 0.626 | 9.85% | 0.859 | 11.28% |
| C6 tyrosine substances | 0.3392 | 4.00% | 0.344 | 3.95% | 0.328 | 5.13% | 0.468 | 6.15% |

In sediments, the abundance of amoA, anammox and denitrification genes was higher than overlying water (Figs 3 and 4), demonstrating that sediment was a hotspot of N metabolisms and removal. Compared with LHS, the abundance of amoA, anammox and denitrification genes were significantly lower in FCS (Figs 3 and 4). Just as that observed in overlying water, the abundance of denitrification genes significantly decreased in summer in both rivers (Fig 4).

![Fig 3. The abundance of 16S, AOB, anammox bacteria and nxr gene in overlying water (A) and sediment (B) in different seasons of FC and LH.](https://doi.org/10.1371/journal.pone.0238531.g003)
clustered according to seasonal variation, however, benthic communities clustered according to sites (Fig 5). The separation of LH and FC benthic communities might be attributed to the niche differentiation between FC and LH sediment (Fig 2 and Table 1), indicating that pollutants had more legacy effect on benthic communities compared with aquatic communities.

A total of 36 and 58 phyla were identified in aquatic and benthic communities, respectively. For aquatic communities, 7 phyla presented relative abundance higher than 1%, and Oxyphotobacteria, Proteobacteria, Bacteroidetes and Actinobacteria jointly comprised of over 95% of all aquatic sequences in each sample (Fig 6A). Typical water born genera, including Rhodoluna, Polynucleobacter, Fluviicola, and Limnohabitans dominated in aquatic communities of both rivers (Fig 6B). These water born genera are aerobic anoxygenic phototrophs (AAPs), using acetate and other low molecular weight photoproducts, or degradation of high molecular weight compound algae exudates [27–29]. An unidentified Oxyphotobacteria, a bacterial methylotroph (Methylotenera) and organic degraders (Pseudorhodobacter, Flavobacterium, Fluviicola, and Novosphingobium) also dominated in both rivers (Fig 6B). Although heterotrophic processes are considered to prevail in freshwater rivers and lakes with high terrestrial loading and a high content of dissolved organic carbon [30], function analysis of bacterial communities by PICRUSt showed that photosynthesis and methyl-oxidization were the

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**Table 2. The α-diversity analysis metrics of bacterial community in FC and LH.**

|                  | River | Chao 1 | Shannon H | Simpson |
|------------------|-------|--------|-----------|---------|
| **Overlying water** |       |        |           |         |
| Spring           | LH    | 691    | 6.43      | 0.952   |
|                  | FC    | 555    | 3.53      | 0.782   |
| Summer           | LH    | 1477   | 7.28      | 0.983   |
|                  | FC    | 1475   | 7         | 0.973   |
| **Sediment**     |       |        |           |         |
| Spring           | LH    | 2798   | 7.47      | 0.976   |
|                  | FC    | 2600   | 7.05      | 0.968   |
| Summer           | LH    | 4123   | 9.99      | 0.997   |
|                  | FC    | 2667   | 8.11      | 0.972   |

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dominant function in overlying water in both rivers (Fig 7). In the meantime, we found that the relative abundance of some denitrifiers, such as *Malikia*, *Flavobacterium*, and *Rhodocyclaceae*, decreased in summer, especially in FC (Fig 6B), indicating the potential of denitrification decreased in summer.

Benthic communities of LH were dominated by Proteobacteria and Bacteroidetes, which accounted for 88.1% and 77.2% of the total sequences in spring and summer, respectively (Fig 6A). Proteobacteria, Bacteroidetes, and Firmicutes dominated in FCS community (Fig 6A). The high abundance of Firmicutes (40.9% and 20.5% in spring and summer, respectively) in FCS might be attributed to the high level of organic pollution and low DO (Fig 2) as members in Firmicutes have been known for fermenting organic matters anaerobically and producing volatile organic acids [31, 32]. The predominant taxa in FCS community, such as *Novosphingobium*, *Acinetobacter*, *Flavobacterium*, *Geobacter*, and *Dechloromonas* were also identified in other heavily polluted urban river sediments and anaerobic active sludge (Fig 6C) [31, 33].

In benthic communities of FC and LH, LEfSe analysis identified 27 biomarkers at an LDA threshold of 4.0 (Fig 8). The summer FCS community had the greatest abundance of statistically unique taxa. Most of the marker taxa in summer FCS, such as *Smithella*, *Syntrophus*, *Thermomonas*, *Syntrophorhabdus*, *Thiobacillus*, *Christensenellaceae*, and *Rhodobacter* (Fig 6C), could degrade diverse organic matters to acetate [34, 35]. In the spring LHS, a large portion of organic degraders and heterotrophic denitrifiers were identified as the marker phylotype, including *Flavobacterium*, *Massilia*, *Arenimonas*, *Pseudomonas*, and *Lysobacter*, while the summer LHS taxa were dominated by *Dechloromonas*, *Geothermobacter*, *Oxyphotobacteria*, *Geobacter*, *Fluvicola*, and *Methylomonas* (Fig 8). As predicted by PICRUSt, the sediment core communities had higher levels of predicted functions involved in fermentation, anaerobic respiration (such as sulfate, arsenate, fumarate, manganese, and iron), and complex-carbon compound degradation (Fig 7). More fermentation, sulfate reduction and metal oxidation and reduction function were found in FCS community (Fig 7).

### Potential nitrogen removal rate in urban river and its determining factors

Laboratory $^{15}$N tracer techniques are often used as standard methods to determine the potential rates of denitrification and anammox. This technique has been successfully applied to quantify the in situ N removal rates [24]. Our study found that the rates of
anammox were significantly higher than denitrification in both rivers \( (p < 0.05) \) (Fig 9). The highest anammox rate was detected in the spring FCS \( (3.64 \pm 0.10 \, \mu \text{mol N g}^{-1} \text{h}^{-1}) \), which accounted for 95.87\% of the total N removal rate. In LHS, denitrification rates \( (1.32 \pm 0.065 \, \mu \text{mol N g}^{-1} \text{h}^{-1} \text{ in spring, } 0.30 \pm 0.099 \, \mu \text{mol N g}^{-1} \text{h}^{-1} \text{ in summer}) \) were about 3–5 times higher than LHW \( (0.38 \pm 0.006 \, \mu \text{mol N g}^{-1} \text{h}^{-1} \text{ in spring, } 0.05 \pm 0.004 \, \mu \text{mol N g}^{-1} \text{h}^{-1} \text{ in summer}) \). However, the denitrification rates in FCS \( (0.16 \pm 0.005 \, \mu \text{mol N g}^{-1} \text{h}^{-1}) \) and FCW \( (0.12 \pm 0.002 \, \mu \text{mol N g}^{-1} \text{h}^{-1}) \) were similar in spring and significantly lower denitrification rates were observed in summer FCS \( (0.005 \pm 0.002 \, \mu \text{mol N g}^{-1} \text{h}^{-1}) \), indicating the denitrification in the sediment of FC river was severely inhibited in summer (Fig 9). These data also showed that the rates of denitrification and anammox in both rivers was significantly inhibited in summer regardless of overlying water and sediment \( (p < 0.05) \) compared with their spring counterpart (Fig 9). The potential denitrification rate of LHS was higher than that of FCS in spring and summer, and the potential rate of anammox in LHS was higher than FCS in summer (Fig 9).
We conducted correlation analysis on the factors affecting potential nitrogen removal rate, and the results showed that the denitrification rate in both of overlying water and sediments had a significant positive correlation with the abundance of denitrification genes (S3 and S4 Tables). \(\text{NH}_4^+\cdot\text{N}\) had a significant negative correlation with both of denitrification and ana-mmmox rates in sediment (S4 Table). The diversity of bacterial communities was negatively correlated with anammox rate \((p < 0.01)\) (S3 and S4 Tables). In overlying water, the potential denitrification rate was significantly positively correlated with TOC \((p < 0.01)\) (S3 Table), while the potential denitrification rate was significantly negatively correlated with TOC in sediment \((p < 0.01)\) (S4 Table).

**Discussion**

**Characteristics of water and sediment qualities in an urban center and suburban river**

Water and sediment qualities are integrated representations of multiple physical, chemical, and biological processes. Similar to other polluted urban rivers [2, 6, 16], FC river, which ran through the urban center, was characterized by high contents of TN, TP, \(\text{NH}_4^+\cdot\text{N}\), and low levels of DO (Figs 1 and 2). In the overlying water of both rivers, the concentrations of TN, TP, and \(\text{NO}_3^-\cdot\text{N}\) in summer were higher than spring, which might be attributed to a larger amount of surface runoff containing higher pollutants into rivers caused by rainfall in summer (674.70 mm during May to September, accounting for 64.38% of the annual rainfall in Wuxi) [36]. However, the seasonal variation of water quality in FCW was not as high as LHW (Fig 1),
indicating that pollution flowing into a river in urban area was more intense and stable than in a suburb regardless of seasonal variation.

Previous studies have demonstrated that as a city develops along a river, the discharge of rough domestic wastewater results in high concentrations of degradable DOM in urban rivers [37, 38]. Our results of 3D-EEM analysis of sediment DOM indicated that more “fresh” protein-like organics accumulated in FCS, whilst more recalcitrant DOM existed in LHS (Table 1). Thus, the high proportion of protein-like organics might explain the high TN and TP contents in FCW, meanwhile, degradation of protein-like organics would simultaneously release NH$_4^+$-N by ammonification process and exhaust DO in FCW (Fig 1). Our data also showed that the concentration of NO$_3^-$ in FCW was lower than LHW (Fig 1), which might be partly attributed to the low DO in FCW (Fig 1) for O$_2$ is the essential substrate of the nitrification process. The slow transformation of NH$_4^+$-N to NO$_3^-$ subsequently resulted in N accumulation in FC for nitrification was normally considered as the first step of N removal [13].
In sediments, although almost all of the nutrient levels of both rivers increased in summer, we found that TP significantly decreased in FCS (Fig 2). Linear regression analysis showed that contents of sediment TN and TP had a significantly positive relationship with TN and TP levels in overlying water (S1 Fig), indicating the mutual influence of water and sediment qualities. The combination of inflow pollutants and microbial driven biogeochemical processes might explain the difference between FC and LH rivers. Sediments not only act as sink and source of carbon and nutrients, but are also is the hotspots of biogeochemical processes. Firstly, the inflow pollutants into river caused different substance suitable for microbes and changed the environment in sediments. In line with a previous report, protein-like DOM enriched in FC, the river located in an urban center [38], whereas microbially-soil-derived humic-like DOM enriched in LH, the suburban river surrounded by wetlands [38]. The protein-like DOM was mainly mineralized to CO$_2$, which would have consumed the DO in sediments [38]. Thus, the high proportion of biodegradable fresh protein-like organics in FC river sediment might have stimulated microbial activities and caused oxygen depletion in sediment [39], furtherly influencing the biogeochemical processes that control nutrient cycling [40]. Subsequently, internal feedbacks were initiated, including phosphorus release from the sediment [19], reduction in activities of nitrification and denitrification [40, 41].

**Difference of potential nitrogen removal pathway and functional gene abundance in an urban center and suburban river**

In this study, we used $^{15}$NO$_3^-$ to quantify the potential rates of denitrification and anammox in both of overlying water and sediments. The results showed that the potential rates of denitrification were significantly lower than anammox in both rivers and both seasons (Fig 9). The low level of denitrification and high proportion of anammox processes indicated that anammox might be the main potential N removal process in the two rivers we studied, especially in the urban center river. Previous studies have also evidenced the widespread distribution of anammox and the significant contribution of anammox to N removal in many aquatic ecosystems [42–44]. In marine sediments, anammox could account for 24–67% of N loss [45, 46] and in
the Black Sea and Gulfo Dulce, 20–40% of N loss in the suboxic water columns was attributed to anammox [46–48]. In both rivers, the rates of denitrification were lower than that in constructed wetlands [44], whereas the denitrification rates were comparable or even higher than that in riparian sediment (about 0.011 \( \mu \text{mol} \ 15\text{N g}^{-1} \text{ dry soil h}^{-1} \)) [49]. Dalsgaard et al. have demonstrated that anammox tolerates higher oxygen concentrations than denitrification [50]. We speculated that under the same oxygen environment, anammox has a higher rate than denitrification. Meanwhile, high concentration of \( \text{NH}_4^+ \)-N in the two rivers is conducive to the occurrence of anammox.

The rates of anammox and denitrification were severely inhibited in summer in both rivers, which might explain the sharp increase of TN in summer besides the heavier inflow pollutant (Figs 1 and 2). As discussed above, the hypoxia in summer resulted in the reduction of potential N removal activities by inhibiting nitrification processes [41]. Our results showed that the abundance of all functional genes related to potential N removal significantly decreased in summer (Fig 4). These combined results indicated that the propagation of the bacteria harboring these functional genes and activities of nitrification, denitrification and anammox might be inhibited in summer, which subsequently slow down the potential N removal in summer, and exacerbate the accumulation of N in rivers.

**Difference of bacterial communities in an urban center and suburban river**

Our data showed that there was no significant difference in the diversity and bacterial community compositions in the overlying water of FC and LH rivers although the studied characteristics of water quality were significantly different between rivers and seasons. Zhao et al. [16] analyzed 39 stations around Jinan city with different water quality and identified hydraulic parameters, temperature, transparency, electrical conductivity, and dissolved oxygen, instead of water quality, as the main driving factors of aquatic communities. Sánchez-Carrillo [51] studied 10 lakes around the world and found temperature and altitude were the important driving factors of food-web structure. Our results and these previous studies combined indicated that the diversity, composition and functions of aquatic bacterial communities were more driven by physical environmental factors instead of water quality, which explained the similarity of aquatic bacterial communities between FCW and LHW (Fig 5). The results of beta-diversity analysis about the way of the bacteria communities gathered was interesting. We consider that the overlying water was fluid and directly related to human seasonal variation of using water. Therefore, bacterial community in overlying water was aggregated according to seasonal variation. The sediments in the river changed little compared with the overlying water, and the sediments were relatively stable. Therefore, the stable benthic microbial community was formed, showing that the benthic bacterial community gathered according to the location.

Differently structured bacterial communities differed in their metabolic substance, degradation pattern and production, and community composition often directly influenced ecosystem function [52]. In contrast to aquatic microbial communities, sediment microbial communities differed between the FC and LH river (Fig 5). Compared with LHS, fermentation and sulfate reduction bacteria existed in high abundance in FCS (Fig 7), which might be explained by the niche selection and adaptation of such functional bacteria to the environment of high carbon and low DO content (Figs 1 and 2). The fermentative process is not effective for mineralization of organics, and the production of fermentation, such as short-chain fatty acids would contribute to the stench odor of urban river [31]. The production of FeS, H\(_2\)S, organic sulfides, \( \text{NH}_3 \), amines, and short chain fatty acids through sulfate reduction and fermentation was considered as the main cause of the black and stench of urban rivers [8]. In addition, the
reduced sulfur might inhibit nitrification [53] and subsequent N removal process [54]. In addition, it was interesting to find that autotrophic denitrifiers, *Thermomonas* and *Thiobacillus*, instead of heterotrophic denitrifiers, dominated in FCS (Fig 6C). The autotrophic denitrifiers, *Thermomonas* and *Thiobacillus*, could reduce nitrate by coupling Fe(II) and reduced sulfur oxidation, respectively [55, 56]. However, denitrification efficiency of autotrophic denitrifiers was lower than heterotrophic denitrifiers due to their slow growth and the inadequate supply of electrons to reduce NO$_3^-$N. Therefore, we proposed that the biochemical processes in FCS might be less effective in removal of organics and N compared with LHS, resulting in the accumulation of organics and N in the urban river (Fig 2), which would then aggravate the accumulation of pollution in the river.

**Factors influencing potential nitrogen removal activities**

We further analyzed the factors influencing potential N removal activities. The denitrification rate in both of overlying water and sediments showed a significant positive correlation with the abundance of denitrification genes (S3 and S4 Tables), so the low level of denitrification genes in summer could explain the corresponding sharply decreased denitrification rate. DO was not correlated with most of the functional genes and potential N removal rates, probably because N was primarily removed through anammox (S3 Table). Although NH$_4^+$-N is one of the substrates of anammox, NH$_4^+$-N had a significant negative correlation with both denitrification and anammox rates in sediment (S4 Table). The high protein-like substance content of FC led to high ammonium formation from organic matter mineralization, resulting in NH$_4^+$-N concentration of (192.4 ± 0.66) mg kg$^{-1}$ dry sediment in FC. A previous study had reported that free un-ionized ammonia inhibited anammox at concentrations exceeding 2 mg N L$^{-1}$ [57]. High NH$_4^+$-N concentration could stimulate the growth and activity of nitrifiers [58], however, nitrifying bacteria have been found to be more sensitive than heterotrophic bacteria to free un-ionized ammonia [59]. High concentrations of NH$_4^+$-N such as those found in FCS, have resulted in the inhibition of nitrification and anammox activity due to presence of free un-ionized ammonia [57, 60, 61]. Organic carbon and nitrate can serve as electron donors and substrates for denitrifying bacteria. The samples were amended by N when potential denitrification rate was determined in our study, the availability of C sources affected denitrification. Our results showed that the potential denitrification rate in water was significantly positively correlated with TOC, which is consistent with previous studies [24]. However, the potential denitrification rate in sediments was significantly negatively correlated with TOC, which may be related to differences in bacterial communities in sediments as above discussion.

Sometimes, higher species diversity of bacterial communities results in greater rates of many functions in ecosystems [62], but we found that diversity was negatively correlated with anammox rate ($p < 0.01$) (S3 and S4 Tables). This finding was probably due to the competition for NH$_4^+$-N with other bacteria affected the growth and gene expression of these functional bacteria related with nitrogen, which dampened the function of potential N removal. Previous studies with synthesized microcosms had demonstrated that high richness increases the inhibitory effects of competitors and dampens ecosystem functions [63, 64]. Both the overlying water and sediment of the rivers had extremely high diversity of bacterial communities in summer, where competitive or antagonistic stress associated with substrate and inter-specific interactions may contribute to the suppression of potential N removal rates in summer. Reduced removal rate of N in the urban river would have intensified N accumulation and would contribute to the continued decline in water quality.
Conclusion

The characteristics of N transformation and underlying bacterial mechanisms was comparatively investigated in a typical urban river and a suburban river, in Wuxi, China. Our results indicated that the concentration of total nitrogen, ammonia and nitrate accumulated faster in the urban river. Consistently, the abundance of functional genes associated with N transformation in the urban river was lower. Bacterial community analysis showed that acidogenic, fermentative and sulfate reductive bacteria dominated in benthic community of the urban river. Isotopic pairing experiments further revealed that anammox was the main potential N removal pathway in both rivers and potential N removal was inhibited in the urban river. Both gene abundance and isotopic pairing experiments showed that potential N removal was inhibited in summer in both rivers. Taken together, the altered N biotransformation and bacterial communities in an urban river could further promote pollution in urban rivers.

Supporting information

S1 Fig. Linear regression analysis of the sediment TN (A) and TP (B) with the water.
(DOCX)

S1 Table. Primers and programs of the target genes in the quantitative PCR analysis.
(DOCX)

S2 Table. The DOM component tryptophan to tyrosine ratio in FCS and LHS.
(DOCX)

S3 Table. Spearman’s correlation coefficient of potential N removal rate with relative abundances of functional genes and environmental factors in overlying water.
(DOCX)

S4 Table. Spearman’s correlation coefficient of the potential N removal rate with relative abundances of functional genes and environmental factors in sediment.
(DOCX)

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