



Function of a landscape lake in the reduction of biotoxicity related to trace organic chemicals from reclaimed water



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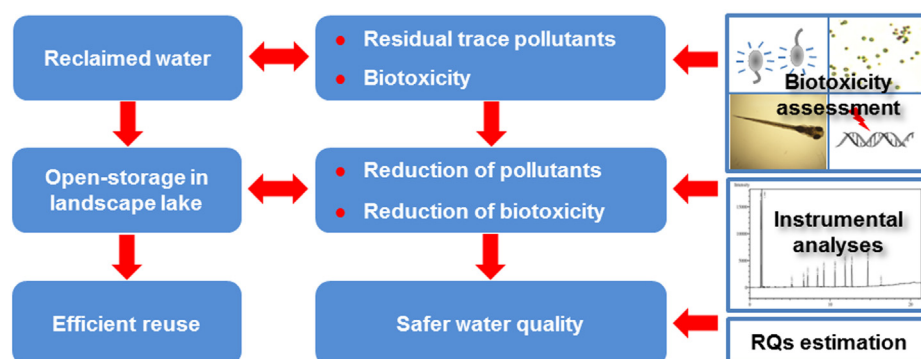
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HIGHLIGHTS

- The lake is a buffer zone in between reclaimed water production and reuse.
- Biotoxicity of reclaimed water decreased during open storage in a landscape lake.
- The dominating chemicals contributing to the biotoxicity were identified.
- The lake provided a favorable semi-natural condition for micropollutants decay.

GRAPHICAL ABSTRACT



ARTICLE INFO

Article history:

Received 18 March 2016
Received in revised form 15 July 2016
Accepted 20 July 2016
Available online 21 July 2016

Keywords:

Biotoxicity
Trace organic chemicals
Reclaimed water
Open storage
Ecological safety

ABSTRACT

The storage of water in a landscape lake can act as a buffer zone between reclaimed water production and reuse, but there is still uncertainty about the variation of water quality and toxic effects during the open-storage process. In this study, long-term sample collection, chemical analyses and biotoxicity assessments were conducted on reclaimed water before and after open storage in a landscape lake. The organic contents, in terms of chemical oxygen demand and total organic carbon, were found to be slightly higher in the lake water than that in the reclaimed water, but substantial reduction of the total concentration of 52 trace organic chemicals was obtained and microorganism toxicity, phytotoxicity, aquatic vertebrate toxicity and genotoxicity, were significantly weakened after open storage. Furthermore, the total risk quotient (RQ_{Total}) decreased from 5.12 (potential ecological risk level) in the reclaimed water to 0.18 (negligible ecological risk level) in the lake water. The removal of chlorpyrifos, dichlorophos and tetracycline was identified as the main reason for biotoxicity reduction after open storage. The semi-natural environment of the landscape lake would have provided a favorable condition for the decay of toxic trace organic chemicals so that the stored water turned to be safer for further reuse.

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1. Introduction

Augmentation of water sources by appropriate reuse of treated wastewater could effectively relieve growing water shortages, but concerns about the safety of reclaimed water often impede its reuse. One direct and effective method to guarantee the safety of water reuse is to monitor the pollutants, particularly trace organic chemicals, using advanced instrumental analysis methods [1]. However, it may not be practically feasible to monitor every potential pollutant because many of them are present only in trace amounts or may have unknown structures that are difficult to identify. Additionally, chemical analyses may not reflect the detrimental effects of the pollutants on aquatic organisms [2]. To complement the limitations of chemical analyses, many *in vitro* and *in vivo* bioassays have been developed to characterize the adverse effects of pollutants in reclaimed water, including their cytotoxicity, genotoxicity, endocrine disrupting effect, photosynthesis inhibition effect, increasing oxidative stress response, and so on [3–6]. Moreover, many websites and organizations, such as the Persistent, Bioaccumulative, and Toxic Profiles, the ECOTOX-ology Knowledgebase, Acute Toxicity Database of the Columbia Environmental Research Center collect experimental and predicted toxicological data of individual chemicals involving species at many trophic levels. That is conducive to the toxicological analyses of pollutants in reclaimed water. A combination of chemical analyses and toxicological analyses can further provide additional information on the ecological safety of reclaimed water reuse.

For water reuse, storage is usually an indispensable intermediate link between reclaimed water production and utilization. Traditionally, the reclaimed water is stored in a nearly enclosed clean-water tank following the treatment process in a reclaimed water treatment plant. An alternative way that is gradually gaining acceptance for effective water reuse is to store the reclaimed water in an open space such as a reservoir, lake, or pond where certain natural processes may promote the stabilization or improvement of the reclaimed water quality [7]. Recharge of the reclaimed water into surface waters is recognized to be a storage approach for indirect potable reuse [8]. On the other hand, water landscaping is usually an important purpose of water reuse in urban area [9]. In this case, the water body receiving the reclaimed water for landscaping can also perform the function of water storage for subsequent environmental uses such as road washing and gardening. During this kind of open storage, the water quality may undergo variations due to the inevitable intrusion of nonpoint sources from the surroundings, and physical, chemical or even biochemical changes may occur in natural surface waters. However, there is little information about the ultimate effects of these actions on the stored water quality, especially regarding the safety of the water for reuse purposes.

This study was conducted by using a water reclamation and reuse system where a lake was constructed to receive the reclaimed water and perform the functions for both landscaping and open storage. To gain insight into the characteristics of water quality variation and its ecological safety after open storage of the reclaimed water under a seminatural condition, comprehensive investigations were conducted by chemical analyses and bioassays with a focus on trace organic chemicals and their bio-toxic effects. The selection of a bioanalytical battery for biotoxicity assessment was based on the notion that the pollutants in reclaimed water and lake water would interact with the species in an aquatic ecosystem by either direct discharge or indirect reuse. Therefore, four bioassays based on the species in three trophic levels of aquatic ecosystem were used here to comprehensively evaluate the non-specific toxicity of waters, and genotoxicity as reactive toxicity was also measured in a complementary assay.

2. Materials and methods

2.1. Description of a water reclamation and reuse system

This study was based on a water recycle system shown in Fig. 1, which starts from groundwater supply for potable purpose, and then enters a circulation of used water collection, water reclamation by anaerobic–anoxic–oxic biological treatment followed by a membrane bioreactor (A²O–MBR), and then reclaimed water distribution for various non-potable purposes such as water landscaping, toilet flushing, and green-belt irrigation. All the collectable used water, including that from toilet flushing, was collected for source enlargement. The A²O–MBR system had a treatment capability of approximately 2000 m³/day and disinfection (by sodium hypochlorite) was performed as the final stage of the treatment. To meet the requirement for different water uses, about half of the reclaimed water was sent directly to a number of buildings for toilet flushing and the other half was transferred to a landscape lake located at the center of the campus. A noticeable feature of the water recycle system was the introduction of the lake which performs both the functions of landscaping and water storage (total storage volume about 5000 m³) for improving the availability of the reclaimed water through multi-step utilization. The depth of the lake was 0.8–1.0 m and the average hydraulic retention time (HRT) was about 5 days which was dynamically controlled by continuous inflow (the reclaimed water) and outflow (pumping for various uses).

2.2. Sample collection and pretreatment

Water samples were collected between December 2014 and June 2015 from the different sampling sites in the water reclamation and reuse system (Fig. 1). During the monitoring period, the sampling frequency was approximately twice a month for normal physicochemical indices analysis, once a month for micropollutants detection and bioassays.

Samples were collected on sunny days to get rid of the influence of rainfall and local runoff. On each sampling day, six liters of the reclaimed water (effluent from the A²O–MBR system) and the same volume of the lake water (near the outlet) were collected into brown glass bottles and taken to the laboratory immediately. Each sample was filtered through a 0.7 μm glass microfiber filter (Φ 150 mm; Whatman™). About 2 L of the filtered water was subjected to chemical analyses, and the remaining volume of the wastewater was directly or indirectly used in bioassays. The raw wastewater and secondary effluent (effluent of A²O) were similarly treated to track the source of pollutants in the generated reclaimed water. Upon collection, the filtered samples were pre-concentrated using solid-phase extraction (SPE) for bioassays and chemical analysis as described in detail in Section SI-1.

2.3. Bioanalytical battery

In total, five different bioassays were conducted to evaluate the biotoxicity effects of pollutants in reclaimed water and lake water. Luminescent bacteria toxicity test (acute toxicity test using *Vibrio fischeri* and *Vibrio qinghaiensis* sp. Q67), algal growth inhibition test using *Chlorella vulgaris*, fish larva mortality test using zebrafish larvae, and genotoxicity assay using the SOS/umu test were performed as described in Section SI-2.

2.4. Chemical analyses

Water quality analyses were firstly conducted by measuring chemical oxygen demand (COD), UV absorbance at 254 nm (UV₂₅₄), total organic carbon (TOC), total phosphorus (TP), NH₄⁺-N, total

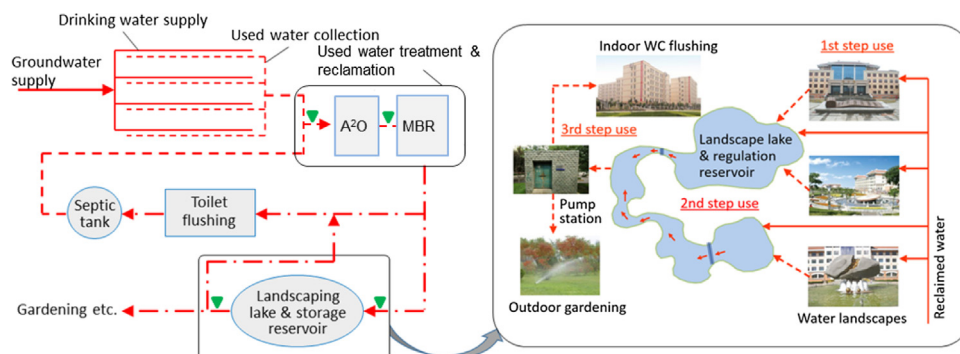


Fig. 1. The water cycle of the wastewater treatment and reuse system [10]. The inverted triangle represents the sampling site.

nitrogen (TN), electric conductivity, turbidity and color according to the standard methods [11]. Specific ultraviolet absorbance (SUVA) was calculated as the ratio of UV_{254} over TOC.

For trace organic chemicals, instrumental methods were applied targeting 59 chemicals including 13 polycyclic aromatic hydrocarbons (PAHs), 3 pesticides, 16 phenols, and 27 pharmaceuticals. PAHs, pesticides and phenols were analyzed on an Agilent 6890 GC coupled to a 5975 MS (GC/MS), equipped with a DB-5MS (30 m × 0.25 mm × 0.25 μm) capillary column. Detailed instrument conditions for determining PAHs, pesticides and phenols are described in Table SI-1. Pharmaceuticals were analyzed on a UPLC coupled with a Waters Micromass Quattro Premier XE tandem quadruple mass spectrometer (UPLC–MS/MS), equipped with a reversed-phase BEH C18 column (100 m × 2.1 mm × 1.7 μm). Instrument conditions for analyzing different categories of pharmaceuticals were described in detail in previous studies [12].

2.5. Ecological risk quotient estimation

2.5.1. Baseline toxicity prediction for chemicals towards aquatic organisms

Toxicity of chemicals towards luminescent bacteria still does not have a sound available database. The prediction of EC_{50} –15 min values of chemicals towards luminescent bacteria could resort to quantitative structure–activity relationships (QSARs). The typical baseline QSAR based on octanol–water partition coefficient (K_{ow}) was applied to obtain the baseline toxicity of neutral PAHs. The liposome–water distribution coefficient K_{lipw} at a defined pH 7, D_{lipw} (pH 7) was estimated and used in the QSARs to calculate the baseline toxicity of polar organic molecules towards *V. fischeri*, including pesticides, phenols, and pharmaceuticals. The detailed calculating procedures and explanation are shown in Section SI-3.

The predicted EC_{50} –96 h of trace organic chemicals to green algae, LC_{50} –48 h to daphnia and LC_{50} –96 h to fish were obtained from the ECOSARv1.10 database (US Environmental Protection Agency). In consideration of ecological safety, the most conservative effect level is used when predictions are identified from multiple classes. These acute toxicity data from four trophic levels were used to estimate predicted no-effect concentrations (PNECs). The chemicals for which the predicted values exceeded their solubility were excluded from the following risk quotient (RQ) estimation. The solubility values of chemicals were obtained from the experimental data in WSKOWWIN v. 1.42.

2.5.2. Methods for risk quotient estimation

The RQ was quantified using the ratio between the measured environmental concentration (MEC) and the PNEC. The RQ of water samples, RQ_{Total} , was calculated following the concept of concentration addition, where the combined effect of the components

in the water sample is equal to the sum of their individual RQ_i , according to Eq. (1) [13].

$$RQ_{Total} = \sum_{i=1}^n RQ_i = \sum_{i=1}^n \frac{MEC_i}{PNEC_i} \quad (1)$$

The PNEC of each chemical was estimated using the ratio between its acute toxicity $L(E)C_{50}$ value and an assessment factor (AF) (Eq. (2)) [14].

$$PNEC = \frac{L(E)C_{50}}{AF} \quad (2)$$

The lowest $L(E)C_{50}$ value of the four representative species, microorganism, algae, daphnia, and fish, was selected to estimate PNEC. The AF was set as 1000 in accordance with the suggestion of the Technical Guidance Document of the European Commission and Guidance for the Implementation of REACH [15,16]. Other considerations are further discussed in the Results and Discussion section.

It is widely accepted that if $RQ_{mix} \geq 1$, the sample has a potential ecological risk for aquatic organisms, while if $RQ_{mix} < 1$, it has no potential ecological risk for aquatic organisms [17]. In addition, it needed to stress that although genotoxicity has attracted extensive attention, there are no reliable and abundant data for mixture genotoxicity prediction and risk assessment.

2.6. Data analysis

Dose metrics of the water samples were expressed as the enrichment factor (EF) of the SPE, which was calculated as the volume of original water sample ($V_{watersample}$) divided by the volume of its SPE extract ($V_{extract}$), as shown in Eq. (3).

$$EF(\text{times}) = \frac{V_{\text{water sample}}(\text{mL})}{V_{\text{extract}}(\text{mL})} \quad (3)$$

For acute toxicity to luminescent bacteria and genotoxicity, the EC_{50} or $IR_{1.5}$ was derived from concentration (EF) – effect curves, and the toxicity of the water samples was then standardized to a toxic equivalent concentration (TEQ), which was expressed as the EC_{50} or $IR_{1.5}$ of reference compound divided by the EC_{50} or $IR_{1.5}$ of water sample (Eq. (4)).

$$TEQ(\text{mg/L or } \mu\text{g/L}) = \frac{EC_{50 \text{ or } IR_{1.5} \text{ of reference compound}(\text{mg/L or } \mu\text{g/L})}{EC_{50 \text{ or } IR_{1.5} \text{ of water sample}(\text{times})} \quad (4)$$

For phytotoxicity, it was difficult to apply TEQs as quantitative indices because of the low detected effect and the large volume of sample required for testing. So, the growth inhibition of SPE extracts of water samples ($EF=40$ times) was used directly to express the toxicity to *C. vulgaris*. Because of similar considerations, the toxicity of water samples without SPE concentration to fish

Table 1
Bioassays and their quantitative evaluation indices used in this study.

Biotoxicity	Assay	Species	Reference compound	Index
Microorganism toxicity	Acute toxicity to luminescent bacteria	<i>V. fischeri</i> ; Q67	Phenol	TEQ _{phenol}
Phytotoxicity	Algal growth inhibition test	<i>C. vulgaris</i>	Potassium dichromate (K ₂ Cr ₂ O ₇)	Growth inhibition
Aquatic vertebrates toxicity	fish larva mortality test	Zebrafish	3,4-Dichloroaniline (3,4-DCA)	Mortality
Genotoxicity	SOS/umu test	<i>Salmonella typhimurium</i> TA1535/pSK1002	4-Nitroquinoline-N-oxide (4-NQO)	TEQ _{4-NQO}

Table 2
Normal physicochemical properties of the reclaimed water before and after storage in the landscape lake.

Parameters	Reclaimed water	Lake water
COD (mg/L)	7.18 ± 4.11	25.89 ± 4.42
UV ₂₅₄ (1/cm)	0.062 ± 0.007	0.072 ± 0.008
TOC (mg/L)	3.59 ± 0.19	5.45 ± 1.05
SUVA (L/mg m)	1.75 ± 0.23	1.35 ± 0.17
TP (mg/L)	0.41 ± 0.13	0.38 ± 0.14
NH ₄ ⁺ -N (mg/L)	0.26 ± 0.23	0.22 ± 0.22
TN (mg/L)	12.81 ± 3.49	6.18 ± 1.27
Electric conductivity (μs/cm)	820.75 ± 27.02	711.63 ± 39.86
Turbidity (NTU)	1.35 ± 0.09	4.35 ± 1.16
Color	7.50 ± 0.96	27.6 ± 5.19

Note: Values are presented as average value ± standard deviation. There were 12 measurements during six months.

larva was directly assayed. The reference compounds and quantitative evaluation indices used in bioassays are summarized in Table 1.

3. Results and discussion

3.1. General water quality before and after storage

As Table 2 shows, the reclaimed water generated by the A²O-MBR system as well as the water stored in the lake achieved the requirements of the reclaimed water standard for toilet flushing, landscape water, and green-belt irrigation [18]. In general, the parameters reflecting inorganic content (such as NH₄⁺-N and electric conductivity) decreased, while the parameters indicating the organic content (COD, TOC, and UV₂₅₄) and sensory indices (turbidity and color) increased after the reclaimed water was stored in the landscape lake. It has been universally acknowledged that the salinity of reclaimed water from urban WWTP is higher than that of source water (normally 1.5–2 times higher than that of tap water) [19]. Salts and nutrients in the reclaimed water may pose potential risks for the normal function of a landscape lake. Unexpectedly, the dissolved inorganic salts and nutrients in the reclaimed water decreased after entering the lake, suggesting that salts and nutrients from reclaimed water did not accumulate in the lake and that those entering the lake from reclaimed water and nonpoint sources did not exceed the environmental capacity of the lake. The natural water purification processes including seasonal uptake by macrophytes, biofouling onto foliage substrates, and feeding by organisms in higher trophic levels may contribute to the removal of salts and nutrients in inland lakes [20]. It is apparent that an open storage system has a high probability of suffering nonpoint pollution, which may cause the increases in turbidity, color, and so on. For example, the increased color of the stored reclaimed water in the lake may result from the transformation of some dissolved chemicals with chromophores that enter the lake water from nonpoint sources. The COD of lake water was more than three times the level of the reclaimed water, and there were slight increases in the UV₂₅₄ and TOC, indicating that the higher COD in the lake water was not due to inputs of organic carbon from nonpoint sources.

3.2. Occurrence of trace organic chemicals before and after storage

3.2.1. Detected chemicals

Of the 59 trace organic chemicals, 52 chemicals including 12 PAHs, 15 phenols, three pesticides, and 22 pharmaceuticals were detected in the reclaimed water, while 42 chemicals consisting 13 PAHs, 12 phenols, one pesticide, and 16 pharmaceuticals were found in lake water (Fig. 2). There was a greater variety of chemicals in the reclaimed water than that in the lake water. For PAHs, 1,12-Benzoperylene was only detected in the lake water, but its detection rate was extremely low. Three phenols (4-chloro-3-methylphenol, 2-chlorophenol, and 2,6-dichlorophenol), two pesticides (chlorpyrifos and dichlorophos), and six pharmaceuticals (acetaminophen, chloramphenicol, diphenhydramine, lincomycin, trimethoprim, and venlafaxine) were founded only in the reclaimed water. The detected chemicals from reclaimed water and lake water were at the average concentrations ranging from 0.18 to 204.08 ng/L, and most were lower than 10 ng/L (Table SI-2). These results are in the previously reported concentration ranges for reclaimed water and surface water [12,21,22].

3.2.2. Possible sources of increased trace organic chemicals

Undoubtedly, trace organic chemicals in reclaimed water are derived from incomplete removal of pollutants in domestic wastewater or by-products of treatment processes. The residuals in the reclaimed water were the major direct source for trace organic chemicals entering the landscape lake, because most of the chemicals were detected in both reclaimed water and lake water (Fig. 2). However, nonpoint source pollution may be another significant pathway for trace organic chemicals to enter the landscape lake. The chemicals found in lake water, but not in reclaimed water, such as 1,12-benzoperylene, might be from nonpoint source pollution. Nevertheless, the new chemicals only from nonpoint source pollution were limited.

In addition, the total PAHs and phenols increased when reclaimed water was stored in the landscape lake (Fig. 3). Because of their lipophilicity and hydrophobicity characteristics, the concentrations of PAHs in water were strongly related with particulate concentration (such as turbidity) [23,24]. The turbidity of the lake water was much higher than that of the reclaimed water, corresponding to the increase in total PAHs (from 21.17 to 36 ng/L). In fact, the concentrations of detected individual PAHs all increased after the reclaimed water entered the landscape lake. PAHs originate from the incomplete combustion of coal, petroleum, timber, and organic polymers, and are widely distributed in the environment [25]. It has been noted that the PAHs content in urban soil around the world is 98–23,250 μg/kg, much higher than that in water [26]. Therefore, the observed increase in PAHs in the lake water likely originated from the ambient environment, such as from contaminated soil or atmospheric deposition [27].

Different changes in concentrations of individual phenols were found in the lake water, although the total concentration of phenols in water increased from 90.52 to 120.07 ng/L during storage in the landscape lake. For example, the concentration of 2,4-dimethylphenol, 3-methylphenol, and 4-methylphenol increased

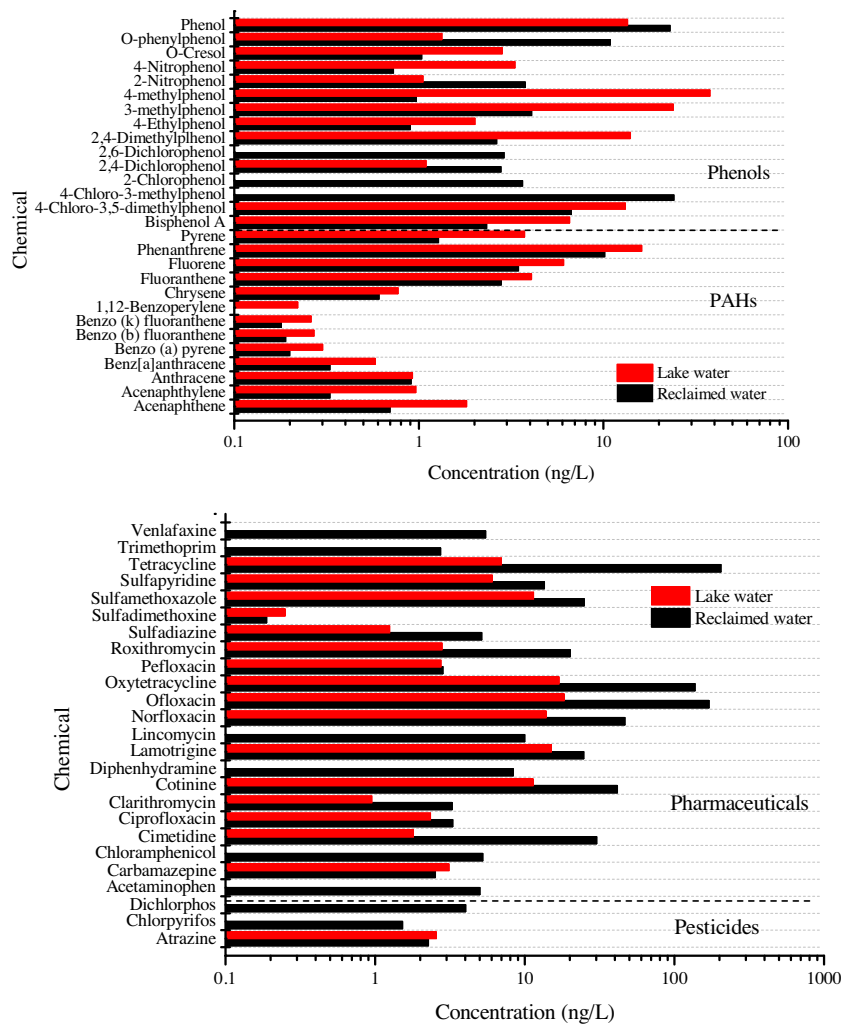


Fig. 2. Average concentrations of chemicals monitored in reclaimed water (n=5) and lake water (n=5). "n" was the number of detected independent samples.

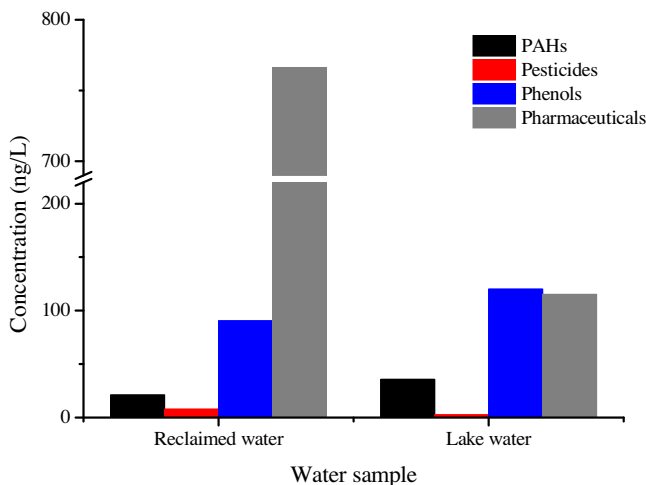


Fig. 3. Total concentration of chemicals monitored in reclaimed water and open-stored water.

dramatically, but 4-chloro-3-methylphenol, 2-chlorophenol, and 2,6-dichlorophenol were not detected in the lake water (Fig. 2). Phenols can derive from degradation products or by-products of natural or artificial compounds (such as lignins and pesticides), or the by-products of the chlorine disinfection

processes [28]. For example, the microbial degradation of chlorinated phenoxyalkanoic-acid pesticides, such as 2,4-dichlorophenoxyacetic acid and 2,4,5-trichlorophenoxyacetic acid, yields plenty of chlorophenols as intermediate metabolites [29,30]. Thus, in addition to nonpoint source pollution, the increase of phenols was likely due to the degradation products or by-products in the seminatural system.

3.2.3. Decay of trace organic chemicals after water storage

Of the four categories of trace organic chemicals, the total concentrations of pesticides and pharmaceuticals decreased after storage in the landscape lake (Fig. 3). Because the single source of raw wastewater was the domestic wastewater of schoolyard where pesticides were rarely used, only three pesticides, including one herbicide and two insecticides, were found in the produced reclaimed water. Of the three detected pesticides, atrazine is one of the most widespread herbicides in the world and its existence and inefficient removal in WWTPs have been reported previously [22,31]. Chlorpyrifos and dichlorophos are also a focus of increasing concern and their occurrence varies with usage in different areas [32,33]. Although these pesticides were not completely removed by the sophisticated treatment processes resulting in residuals in the reclaimed water (Table SI-2), chlorpyrifos and dichlorophos were not detected after open storage (Fig. 2), which suggests that the seminatural system promotes chlorpyrifos and dichlorophos removal.

Table 3

The dominating ecological risk quotients (RQs) of trace organic chemicals based on the mean concentrations in water samples.

Chemicals	Wastewater	Secondary effluent	Reclaimed water	Lake water
Chlorpyrifos	5.26E + 00	4.20E + 00	4.59E + 00	–
Dichlorphos	1.24E + 00	9.55E – 02	1.30E – 01	–
4-Methylphenol	8.34E – 01	4.15E – 04	1.86E – 04	7.19 E – 03
Acetaminophen	4.43 E – 01	–	3.03 E – 03	–
Tetracycline	1.02 E – 01	1.11E – 01	7.12E – 02	2.44E – 03
Total	8.40E + 00	4.68E + 00	5.12E + 00	3.07E – 01

Note: The chemicals with RQ values greater than 0.1 are shown in the table. The chemicals with RQ values greater than 1 are shown in bold. The “Total” was calculated based on all target chemicals.

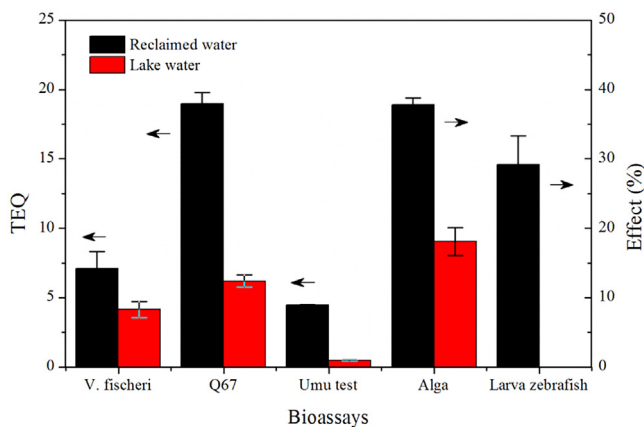


Fig. 4. Results of five bioassays. Luminescent bacteria toxicity test of water sample extracts using *V. fischeri* and Q67 was denoted as TEQ_{phenol} (mg/L). Similarly, the SOS/umu test of water sample extracts was expressed as TEQ_{4-NQO} (μ g/L). Algal growth inhibition tests of water sample extracts (EF = 40 times) were expressed as inhibition at 72 h exposure (%). Fish larva test of water samples without enrichment was denoted as mortality (%) at 96 h exposure. The number of detected independent samples was 5 for every bioassay.

The wide distribution of pharmaceuticals in the environment has been blamed for their presence in effluent discharge from WWTPs [21]. Miège et al. created a database using the results of 117 scientific publications to summarize the occurrence of pharmaceuticals in WWTPs [34]. Most of the pharmaceuticals detected in the reclaimed water in the present study were identified in the database as the frequently detected pharmaceuticals. However, as shown in Fig. 2, 22 pharmaceuticals were detected in the reclaimed water mostly with concentrations of 10^0 – 10^2 ng/L orders, in contrast to 16 pharmaceuticals detected from the lake water mostly with concentrations of 10^0 – 10^1 ng/L orders, while six pharmaceuticals were undetectable in the lake water. In the reclaimed water, the concentrations of ofloxacin, oxytetracycline, and tetracycline were as high as 100–200 ng/L, but in the lake water, their concentrations were decreased by one order of magnitude or more. As a result, the total concentration of the detected pharmaceuticals decreased from 766.27 ng/L in the reclaimed water to 115.28 ng/L in the lake water (Fig. 3). It has been reported that some antibiotics may undergo cleavage in natural water due to oxidation driven by solar photolysis [35–37]. The activities in the natural environment can significantly stimulate the removal of pharmaceuticals.

3.3. Toxicities before and after storage based on bioassays

For the acute toxicity tests using *V. fischeri* and Q67, the inhibitory effect of reclaimed water expressed as TEQ_{phenol} was reduced from 7.12 to 4.14 mg/L for *V. fischeri* and from 18.96 to 6.20 mg/L for Q67 during open storage (Fig. 4). This indicates that natural water purification processes in the landscape lake decreased the acute toxicity of reclaimed water based on organisms at the lowest trophic level.

In this study, even after many-fold enrichment, the water samples still showed very low phytotoxicity, and the effective concentration corresponding to 50% inhibition could not be experimentally obtained. This might be due to the low concentrations of detected pesticides which were closely related to photosynthesis inhibition [38]. After 72 h exposure with EF = 40 times, the inhibition value decreased from 37.77% for the reclaimed water to 18.11% for the lake water (Fig. 4), indicating that the landscape lake could achieve phytotoxicity removal to certain extent.

For the aquatic vertebrate toxicity test, zebrafish larvae without chorions were used in the test for eliminating the action of chorions of zebrafish embryos as barriers against toxicant uptake, especially for high-molecular-weight chemicals [39]. To avoid the interference of cosolvent (dimethyl sulfoxide solution), actual non-concentrated water samples were used in the zebrafish larvae test. After 96 h exposure to the reclaimed water, 29.17% of larvae death was detected in contrast to non-negative effects for the lake water (Fig. 4). The landscape lake, as a seminatural ecosystem, effectively eliminated the adverse effect of the reclaimed water on zebrafish larvae. Moreover, it was observed that the reclaimed water did not lead to zebrafish larvae death until exposure of 72 h, while the mortality increased suddenly in the remaining exposure period (72–96 h). It was speculated that the action of the pollutants in the reclaimed water on larvae was relatively slow.

Regarding the genotoxicity, in the SOS/umu assay without metabolic activation, the TEQ_{4-NQO} decreased from 4.46 μ g/L in the reclaimed water to 0.48 μ g/L in the lake water, indicating that the open storage in the seminatural ecosystem greatly assisted genotoxicity removal.

In common sense, the A^2O -MBR units employed for water reclamation is an advanced treatment system for effective removal of suspended, colloidal pollutants, and biodegradable organic substances to produce water suitable for reuse. However, as a result of the biotoxicity analysis in this study, the reclaimed water still exhibited apparent toxic effects on aquatic organisms of different trophic levels, as well as significant genotoxicity. Nevertheless, after the reclaimed water was led to the landscape lake, these toxic effects were much significantly reduced or even eliminated during water storage. Although biotoxicity is not a direct measure of any specified contaminant, it is contributed by the integral effects of residual contaminants which pose adverse effects on aquatic organisms. The apparent reduction of the detected trace organic chemicals during the open storage of the reclaimed water in the landscape lake could provide explanations on such kind of relationships based on reported studies which indicated the toxicity of specific chemicals [40]. Taking genotoxicity as an example, PAHs, pesticides (e.g. atrazine), phenols (e.g. 2-chlorophenol) and PPCPs (e.g. ofloxacin) are potential geno-toxins [41–44]. Some disinfection by-products (DBPs), not discussed in this study, may also contribute to genotoxicity [45]. In a seminatural ecosystem, such as the landscape lake in this study, naturally occurring physical, chemical, physicochemical, biological and ecological actions may stimulate the decomposition of the above mentioned trace organic to substantially reduce genotoxicity.

3.4. Identification of dominating toxic chemicals based on risk quotient estimation

The acute toxicity data based on four species including bacteria, algae, daphnia, and fish were used to estimate the RQ. The value assigned to AF seriously affected the PNEC estimation that was used to calculate RQ. The Technical Guidance Document of the European Commission suggested that AF was set as 1000 if acute toxicity data are available in at least three test systems on three trophic levels: algae, daphnia, and fish [15]. In this study, acute toxicity data regarding bacteria (*V. fischeri*) was also introduced to estimate PNEC. There were no guidance documents or studies to confirm AF value in this case. Guidance for the Implementation of REACH stated that *V. fischeri* (MICROTOX® test) has limited relevance with WWTP function [16]. So, there was no recommended AF for the effect assessment of microorganisms in WWTP. However, it was different with the objective of calculating PNEC in this study. That was because acute toxicity of chemicals in water towards *V. fischeri* has a close relationship with its ecological risk. Comparison of the collected L(E)C₅₀ values of target trace organic chemicals for the four species revealed that daphnia was the most sensitive species to most target trace organic chemicals except PAHs, to which *V. fischeri* was the most sensitive species. Because of the low concentration of PAHs in water sample, the PNECs of PAHs showed little influence on the estimated result of RQ_{Total}. The PNECs of target chemicals were mainly affected by one species (daphnia). Hence, the AF was set as 1000 in this study.

The RQ_{Total}s of reclaimed water and lake water are shown in Table 3, as well as those of raw wastewater and the secondary effluent for biotoxicity tracking. The RQ_{Total}s of wastewater, secondary effluent, and reclaimed water exceeded one and only the RQ_{Total} of lake water was less than one. This indicated that all the water samples except lake water had potential ecological risk for aquatic organisms. The ecological safety of reclaimed water improved after reserving into the landscape lake, although non-point source pollution introduced some contaminants. This further verified the above biotoxicity test results for reclaimed water and lake water. The lake as a part of campus landscape could not only adjust the amount of water required as a buffer zone, but also improve the ecological safety of reclaimed water for subsequent reuse.

The RQs of target chemicals are shown in Table SI-3. The RQs of the dominating trace organic chemicals in the water samples are shown in Table 3. Chlorpyrifos and dichlorphos, which had RQs exceeding 1, made the greatest contribution to the biotoxicity of wastewaters from reclaimed water production processes (as shown in bold in Table 3). This is because the chlorpyrifos and dichlorphos are extremely toxic to daphnia even in very low concentrations (Table SI-3). Therefore, these two insecticides should receive more attention than other trace organic chemicals in the ecosystem, even though pesticides were not the primary trace organic chemicals based on chemical analysis. Furthermore, 4-methylphenol, acetaminophen, and tetracycline had high concentrations in the water samples and contributed greatly to RQ_{Total}, as shown in Table 3. It is clear that ecological risk reduction of reclaimed water stored in the lake occurred mainly because of the decreased risk from chlorpyrifos, dichlorphos, and tetracycline, which can be tracked to raw wastewater. The concentration reduction of other chemicals in lake water had a certain contribution to reduce its adverse impact on ecological system and then the trace organic chemicals introduced into the lake water by non-point source or treatment processes could not increase the ecological risk observably.

4. Conclusion

Water reuse through a water cycle is an effective way to realize multi-step utilization of the reclaimed water as shown by the system investigated in this study. Within the water cycle, the landscape lake acts as a quantitative and qualitative buffer zone between reclaimed water production and reuse. In addition to its primary function of water landscaping, the lake acts as a reservoir for open-storage of the reclaimed water and inevitably receives non-point pollutant sources. Therefore, it has been uncertain whether the water after storage still meets the quality requirements for subsequent reuse. Through this study, it was identified that although the lake water was slightly deteriorated comparing with the reclaimed water regarding certain conventional parameters such as COD, TOC, turbidity and color, many of the trace organic chemicals detected from the reclaimed water were substantially reduced during water storage. This eventually resulted in a remarkable reduction of the biotoxicity as indicated by the toxicity on aquatic organisms of various trophic levels and genotoxicity, and the RQ_{Total} as a comprehensive index for quantifying the ecological risk level. The method of RQs estimation for each detected trace organic chemical has effectively assisted identifying the dominating toxic organic chemicals contributing to the biotoxicity and helped us to clarify the main reasons for toxicity reduction in the lake water. It can thus be concluded that the seminatural environment of the landscape lake has provided a favorable condition for the decay of toxic trace organic chemicals to improve the ecological safety of the stored water. Further studies are still necessary for investigating how certain trace organic chemicals residual in the reclaimed water are reduced during open storage in such a system.

Acknowledgements

This research was supported by the National Natural Science Foundation of China (Grant No. 51508449), the National Program of Water Pollution Control in China (Grant No. 2013ZX07310-001), Fund for Postdoctoral Scientific Research Project, China (2015M572531) and the Program for Innovative Research Team in Shaanxi (Grant No. IRT2013KCT-13). The authors are grateful to Mr. Oda Yoshimitsu for the provision of *Salmonella typhimurium* TA1535/pSK1002.

Appendix A. Supplementary data

Supplementary data associated with this article can be found, in the online version, at <http://dx.doi.org/10.1016/j.jhazmat.2016.07.050>.

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